Stat 615 Final_Project

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ACKNOWLEDGEMENTS

"I am not what happened to me, I am what I choose to become" by Christopher Gardner, The Pursuit of Happiness.

It is always a pleasure to remember the fine people who guided me in the Regression program. I received to uphold my practical and theoretical skills during the respective session. Firstly, I would like to thank **Pro. James C. Dickens** and secondly, I want to thank my family & friends for their love, motivation, and support during this semester in American university. Thanks for all the ideas, opinions, knowledge, and suggestions given to me to help me to complete this report. We are very thankful to American University for giving us the opportunity to pursue this project.

Title Page with Executive Summary

Title: Estimating Medical Cost via random forest.

Type of analysis: Application analysis

Table 1:

Name	course
Dhruv Jain	STAT -615
Mekdim Ashebo	STAT -615

```
# calling all the libraries used in the code book
library(olsrr)
library(tidyverse)
library(dbplyr)
library(dplyr)
library(Matrix)
library(MASS)
library(ggplot2)
library(tibble)
library(data.table)
library(ggmosaic)
library(ggforce)
library(ggmap)
library(ggthemes)
library(purrr)
library(keep)
library(readr)
library(gridExtra)
library(randomForest)
library(corrplot)
library(PerformanceAnalytics)
```

1 All About Data Set

1.1 Data Set Source

Summary: The type of analysis is **application based**. This data set is collected from different open-source databases manually. offer a preliminary description of the data set. For example, indicate the size of the data source, describe the variables, and include any other data profile information that would be of interest.

Data source: website name:Kaggle. Website link

Overall approach is Cleaning, analyzing, again cleaning, answering the five questions, playing with data to deliver more better output and finally graphical representation. **Defining** the issues and trying to resolve that by presenting the medical cost data set . **Measure** overall what can be done. **Analyze** the data to use for future capability. **Improving** You will use information gathered in the previous phases to design and implement improvements in processing with consistency. Overall Approach to this question is using various tools and coding sets with providing the statistical data with convincing evidence.

Clean -> Design -> Plot --> Types Regression model -> hypothesis -> statistical findings -> Random forest -> conclusion

Recommendations: Using R-studio markdown one can achieve high graphical results with better quality and one can also conclude that big data can be easily handled on this platform. Using this platform, we will continue to evaluate the medical cost for personal data using this tool. The issue is that what can be done with the data can we provided some explanation to justify the results. We would like to address those five questions with different visual ideas.

IMPORTANT KEYWORDS:

- charges
- Smoker
- sex
- BMI (body mass index)
- P-value
- Random forest

Columns Description

age: age of primary beneficiary sex: insurance contractor gender, female, male bmi Body mass index, providing an understanding of body, weights that are relatively high or low relative to height, objective index of body weight (kg / m ^ 2) using the ratio of height to weight, ideally 18.5 to 24.9. children: Number of children covered by health insurance / Number of dependents smoker: Smoking region: the beneficiary's residential area in the US, northeast, southeast, southwest, northwest. charges: Individual medical costs billed by health insurance

We had to randomly sample 300 rows from our original data. We then saved these 300 rows into csv file so that we can import them later. going forward We will take that csv file. (which has only be run once)

```
# The preliminary steps we did
#insurance <- read_csv('Downloads/insurance.csv')
#insurance_300 <- sample_n(insurance, 300)
#write.csv(insurance_300 , file = "Desktop/insurance_300.csv")</pre>
```

```
insurance_new <- read_csv("insurance_300.csv",
col_types = cols(
   age = col_double(),
   sex = col_character(),
   bmi = col_double(),
   children = col_double(),
   smoker = col_character(),
   region = col_character(),
   charges = col_double()
))</pre>
```

- The data contains 300 rows and 7 columns. This project is about determining the factors that affect medical costs billed by health insurance.
- The independent variables include three categorical variables and three quantitative variables. Sex (male/female), region(Northeast, northwest etc), and smoker(whether a person smokes or not) are the categorical variables. While the quantitative variables include the BMI index, the age and the number of children the person have.

```
head(insurance_new,8)
```

```
## # A tibble: 8 x 7
##
                     bmi children smoker region
       age sex
                                                     charges
                  <dbl>
                            <dbl> <chr>
                                          <chr>>
##
     <dbl> <chr>
                                                       <dbl>
## 1
        63 female 25.1
                                                      14255.
                                0 no
                                          northwest
## 2
        18 male
                    38.2
                                0 yes
                                          southeast
                                                      36308.
## 3
        48 male
                    29.6
                                0 no
                                          southwest
                                                      21232.
## 4
        46 female
                   33.4
                                          southeast
                                                       8241.
                                1 no
## 5
        52 male
                    30.2
                                1 no
                                          southwest
                                                       9725.
## 6
        36 female
                   19.9
                                0 no
                                          northeast
                                                       5458.
## 7
                    20.9
        19 male
                                1 no
                                          southwest
                                                       1832.
## 8
        48 male
                    36.7
                                 1 no
                                          northwest
                                                      28469.
nrow(insurance_new)
## [1] 300
ncol(insurance_new)
## [1] 7
colnames(insurance_new)
```

[1] "age" "sex" "bmi" "children" "smoker" "region" "charges"

• Let us quickly investigate the summary of our dependent variable. The median insurance charge is around 10097 and the mean of 13283. The standard deviation is 11399.

summary(insurance_new)

```
##
                         sex
                                               bmi
                                                              children
         age
##
           :18.00
                     Length:300
                                                 :17.29
                                                           Min.
                                                                  :0.00
    Min.
                                         Min.
##
    1st Qu.:27.00
                     Class : character
                                          1st Qu.:25.25
                                                           1st Qu.:0.00
    Median :40.50
                                         Median :30.01
                                                           Median:1.00
##
                     Mode :character
##
    Mean
            :39.88
                                          Mean
                                                 :30.02
                                                           Mean
                                                                  :1.02
##
    3rd Qu.:53.00
                                          3rd Qu.:34.20
                                                           3rd Qu.:2.00
##
            :64.00
                                                 :46.75
                                                                  :5.00
    Max.
                                          Max.
                                                           Max.
##
                                                charges
       smoker
                           region
    Length:300
                        Length: 300
##
                                             Min.
                                                    : 1136
##
    Class : character
                        Class : character
                                             1st Qu.: 5134
##
    Mode :character
                        Mode :character
                                             Median :10097
##
                                             Mean
                                                    :13283
##
                                             3rd Qu.:17154
##
                                             Max.
                                                     :51195
```

• Type of columns used in data frame (double, charterer)

str(insurance_new)

```
## spc_tbl_ [300 x 7] (S3: spec_tbl_df/tbl_df/tbl/data.frame)
## $ age : num [1:300] 63 18 48 46 52 36 19 48 19 19 ...
             : chr [1:300] "female" "male" "male" "female" ...
## $ sex
## $ bmi
             : num [1:300] 25.1 38.2 29.6 33.4 30.2 ...
## $ children: num [1:300] 0 0 0 1 1 0 1 1 0 0 ...
## $ smoker : chr [1:300] "no" "yes" "no" "no" ...
## $ region : chr [1:300] "northwest" "southeast" "southwest" "southeast" ...
   $ charges : num [1:300] 14255 36308 21232 8241 9725 ...
##
##
   - attr(*, "spec")=
##
    .. cols(
         age = col_double(),
##
         sex = col_character(),
##
    .. bmi = col_double(),
    .. children = col_double(),
##
##
       smoker = col_character(),
##
    .. region = col_character(),
##
       charges = col_double()
##
    ..)
## - attr(*, "problems")=<externalptr>
```

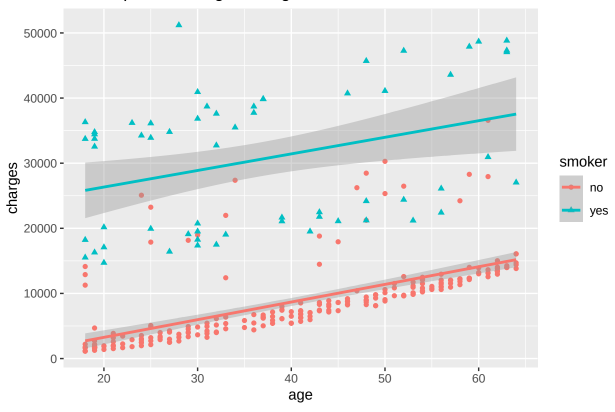
1.2 Cleaning the data and type of columns

[1] "northwest" "southeast" "southwest" "northeast"

```
# calculating NA/missing data in columns
colSums(is.na(insurance_new))
                                                  region charges
##
        age
                 sex
                          bmi children
                                         smoker
##
          0
                   0
# converting to factor variable
insurance_new$sex = as.factor(insurance_new$sex)
insurance_new$smoker = as.factor(insurance_new$smoker)
# how many unique values
unique(insurance_new$sex)
## [1] female male
## Levels: female male
unique(insurance_new$children)
## [1] 0 1 2 5 3 4
unique(insurance_new$smoker)
## [1] no yes
## Levels: no yes
unique(insurance_new$region)
```

```
# Check levels of smoker variable
table(insurance_new$smoker)
##
## no yes
## 239 61
# Check levels of region variable
table(insurance_new$region)
##
## northeast northwest southeast southwest
         67
             77
# Check levels of sex variable
table(insurance_new$sex)
##
## female male
##
     134
          166
1.3 Visualization
1.3.a Does age affect medical charges for smoker?
## 'geom_smooth()' using formula = 'y ~ x'
```

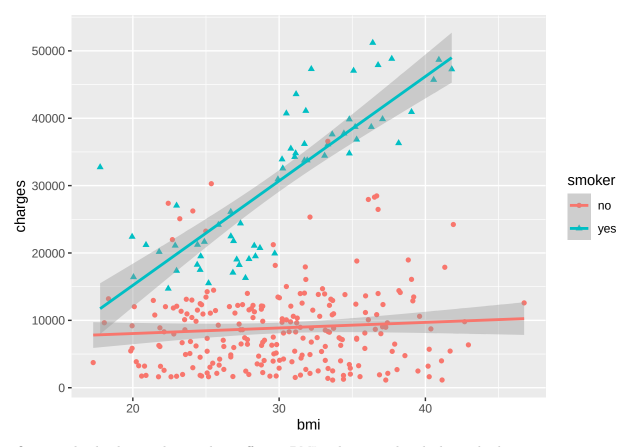
Scatter plot for charges vs Age



- Yes the charges are increased as we increase the number of age. Now the fun part is if a person smokes he/she is paying more charges on medical then the person not smoking.

1.3.b Does Body mass index (BMI) affect medical charges for smoker?

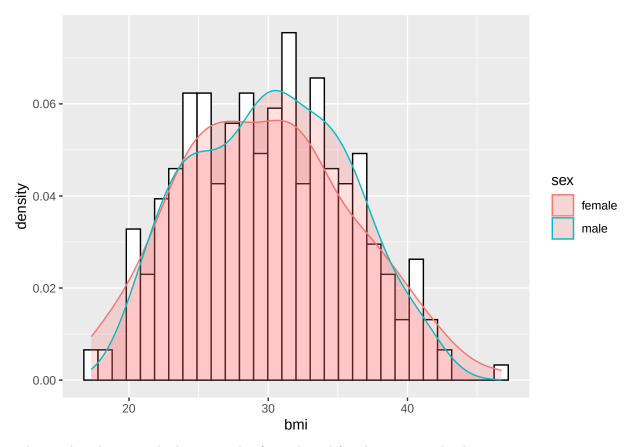
'geom_smooth()' using formula = 'y ~ x'



- One can clearly observe that smoking affect in BMI and increased with the medical expenses.

1.3.c Histogram for density graph for Body mass index (BMI)

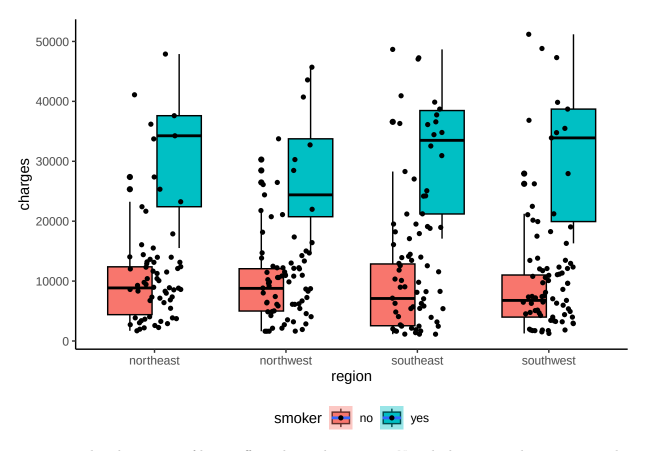
'stat_bin()' using 'bins = 30'. Pick better value with 'binwidth'.



- The graph is density vs body mass index for male and female counts in the data set.

1.3.d High no. of smokers from which region ?

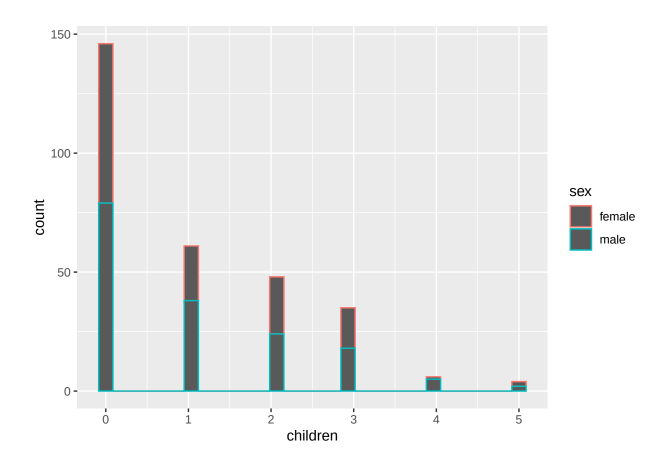
'geom_smooth()' using formula = 'y ~ x'



- comparing that does region of living affects the smokers or not. Now, looking at graph one cans say that the region does affect the charges on medical insurance.

1.3.e

'stat_bin()' using 'bins = 30'. Pick better value with 'binwidth'.



1.4 No. of counts

```
# Female
insurance_new%>%
  filter(sex == "female")%>%
  count(sex,children,smoker,region)%>%
  arrange(sex, smoker) -> counts
head(counts, n=5)
## # A tibble: 5 x 5
            children smoker region
     sex
                                           n
     <fct>
               <dbl> <fct> <chr>
##
                                       <int>
## 1 female
                   0 no
                            northeast
## 2 female
                   0 no
                            northwest
                                          17
## 3 female
                   0 no
                            southeast
                                          14
## 4 female
                   0 no
                            southwest
                                          13
## 5 female
                   1 no
                            northeast
                                           5
# Male
insurance_new%>%
  filter(sex == "male")%>%
  count(sex,children,smoker,region)%>%
```

```
arrange(sex, smoker) -> counts_male
head(counts_male, n=5)
## # A tibble: 5 x 5
##
           children smoker region
     sex
                                           n
##
     <fct>
              <dbl> <fct>
                            <chr>
                                       <int>
## 1 male
                  0 no
                            northeast
                                          17
## 2 male
                   0 no
                            northwest
                                          15
## 3 male
                   0 no
                            southeast
                                          14
## 4 male
                   0 no
                            southwest
                                          14
## 5 male
                            northeast
                                           6
                   1 no
```

2 QQ & Multicollinearity plot description

2.1 Multicollinearity plot

- The response variable is not dependent on explanatory variable interns of multicollinearity.
- The highest correlation is between charges and age with only 0.24. But if we exclude charges since charges is dependent variable, the highest correlation among the independent variables.
- The age with bmi with only 0.04 which is nearly 0. So there exists no colinearity among the independent variables. This suggests that each of the variables might be useful if they are included in the regression model as they don't have any correlation with each other.

```
numeric_insurance <- cor(insurance_new[,c("bmi", "children", "age", "charges")])
numeric_insurance</pre>
```

```
## bmi children age charges

## bmi 1.00000000 -0.01371482 0.04733455 0.1785191

## children -0.01371482 1.00000000 0.03529611 0.0781793

## age 0.04733455 0.03529611 1.00000000 0.2461625

## charges 0.17851911 0.07817930 0.24616249 1.0000000
```

- we can also the scatter plots between the independent variables clearly there is no pattern that we can see verifying our output from the correlation matrix.
- One can say from the graph that the points are independently plotted and one cannot find any kind of pattern on left side of graph. On the other hand one can identify the

2.2 Normality plot

```
# Light tailed at the end
qqnorm(insurance_new$bmi)
```

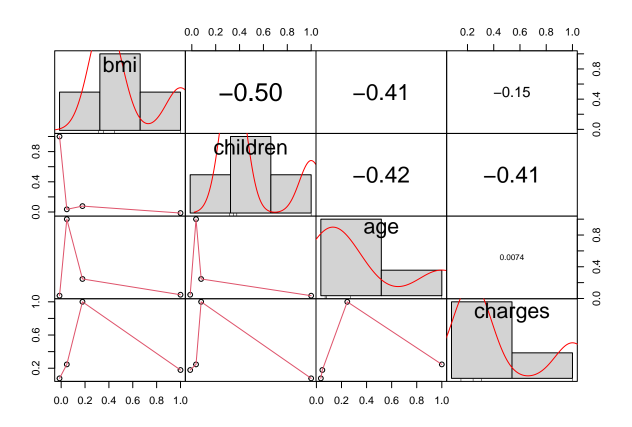
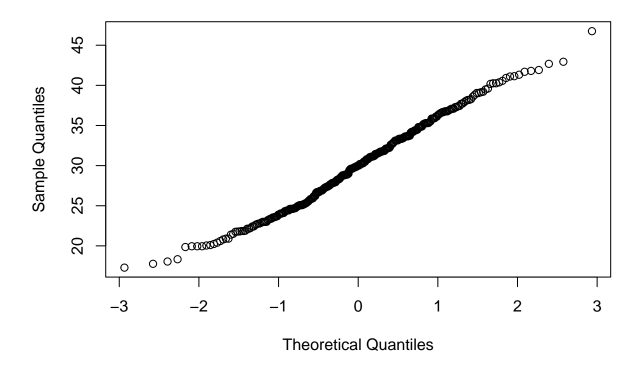


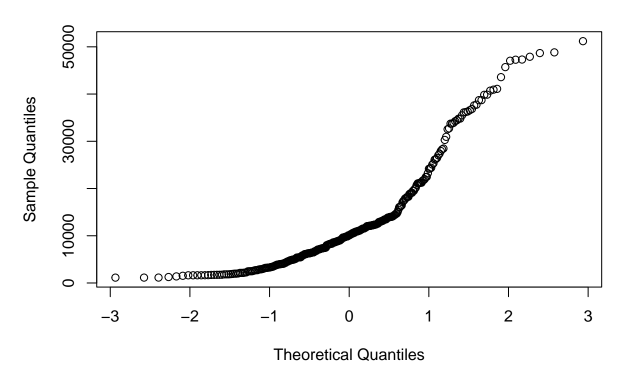
Figure 1: multicollinearity plot

Normal Q-Q Plot



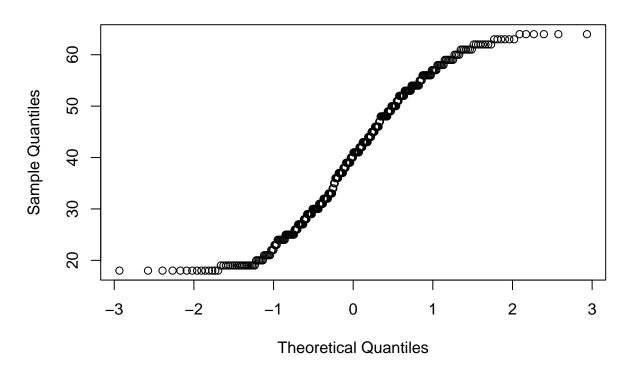
right skewed
qqnorm(insurance_new\$charges)

Normal Q-Q Plot



Heavy tailed at the end
qqnorm(insurance_new\$age)

Normal Q-Q Plot



3 Full Regression Model

• full regression model including both categorical and quantitative variables

```
lm(charges ~ age + children + bmi + region + sex + smoker, data = insurance_new) -> x
##
## Call:
## lm(formula = charges ~ age + children + bmi + region + sex +
       smoker, data = insurance_new)
##
##
##
   Coefficients:
##
       (Intercept)
                                              children
                                                                     bmi
                                 age
          -12033.2
                               261.1
                                                 532.8
                                                                  353.3
##
                    regionsoutheast
                                     regionsouthwest
## regionnorthwest
                                                                sexmale
           -1545.5
                             -1505.4
                                                                  607.8
##
                                               -1719.9
##
         smokeryes
##
           22876.4
```

4 Matrix

4.1 Fitted & Residual values using matrix for quantitative variables

```
Xm <- model.matrix(~age + children + bmi , data=insurance_new )</pre>
Ym <- as.matrix(insurance_new%>%dplyr::select(charges))
# Let's use R code to establish matrix X :
# A = (X^T*X)^{-1}*X^TY
(solve(t(Xm)%*%Xm))%*%(t(Xm)%*%Ym)
##
                charges
## (Intercept) -4825.6927
          185.6491
669.1986
## age
## children
## bmi
               333.8682
# fitted values
Xm%*%((solve(t(Xm)%*%Xm))%*%(t(Xm)%*%Ym)) \rightarrow fitted_values
#-----
# residual values
# producing the table of residula and fitted plot for the model.
matrix = data.frame(fitted_values, residual_values)
names(matrix)[1] <- "fitted_values"</pre>
names(matrix)[2] <- "residual_values"</pre>
head(matrix, 6)
    fitted_values residual_values
##
       15243.617 -989.0085
## 1
## 2 11259.742 25048.0563
## 3 13967.964 7264.2178
## 4 15547.919 -7307.3293
## 5 15580.081 -5855.5505
## 6 8486.629 -3028.5827
#-----
# Both fitted values and residual values match with matrix model
lm(charges ~ age + children + bmi, data= insurance_new) -> AB
#summary(AB)
#residuals(AB)
#fitted(AB)
```

4.2 Matrix method with both quantitative and qualitative variables (dummy variables included automatically)

```
# The results are the same using both the matrix method and lm method.
Xm <- model.matrix(~age + children + bmi + region + sex+ smoker , data=insurance_new )</pre>
Ym <- as.matrix(insurance_new%>%dplyr::select(charges))
# Let's use R code to establish matrix X :
# A = (X^T*X)^{-1}X^TY
(solve(t(Xm)%*%Xm))%*%(t(Xm)%*%Ym)
##
                     charges
## (Intercept) -12033.2491
## age
                   261.1443
## children
                   532.7552
## bmi
                    353.3493
## regionnorthwest -1545.5302
## regionsoutheast -1505.3521
## regionsouthwest -1719.9011
## sexmale
                   607.7807
              22876.3789
## smokeryes
# fitted values
Xm%*%((solve(t(Xm)%*%Xm))%*%(t(Xm)%*%Ym)) \rightarrow fitted_values
# residual values
Ym-Xm%*%((solve(t(Xm)%*%Xm))%*%(t(Xm)%*%Ym)) \rightarrow residual_values
#-----
# producing the table of residula and fitted plot for the model.
matrix = data.frame(fitted_values, residual_values)
names(matrix)[1] <- "fitted_values"</pre>
names(matrix)[2] <- "residual_values"</pre>
head(matrix,6)
## fitted_values residual_values
## 1 11735.311 2519.298
## 2
       28133.497
                       8174.301
## 3
        9848.695
                      11383.488
     10822.791
## 4
                      -2582.201
## 5 11638.037
## 6
                      -1913.507
## 6
       4383.695
                       1074.351
# Both fitted values and residual values match with matrix model
lm(charges ~ age + children + bmi+ region + sex + smoker, data= insurance_new) -> AB
#summary(AB)
#residuals(AB)
#fitted(AB)
```

5 Analyze and Evaluate the full model

```
lm(charges ~ age + children + bmi+ region + sex + smoker, data= insurance_new) -> AB
summary(AB)
```

```
##
## Call:
## lm(formula = charges ~ age + children + bmi + region + sex +
       smoker, data = insurance new)
##
##
## Residuals:
     Min
              1Q Median
                            30
                                  Max
## -10712 -3120 -1095
                          1496
                                24152
##
## Coefficients:
##
                    Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                   -12033.25
                                2155.91
                                        -5.582 5.46e-08 ***
## age
                      261.14
                                  24.05 10.858
                                                < 2e-16 ***
## children
                      532.76
                                 279.27
                                         1.908
                                                  0.0574 .
## bmi
                      353.35
                                  62.03
                                         5.696 2.99e-08 ***
                   -1545.53
## regionnorthwest
                                 992.47
                                        -1.557
                                                  0.1205
## regionsoutheast
                   -1505.35
                                1036.22 -1.453
                                                  0.1474
## regionsouthwest
                   -1719.90
                                 990.21
                                         -1.737
                                                  0.0835
## sexmale
                      607.78
                                                  0.3823
                                 694.59
                                         0.875
## smokeryes
                    22876.38
                                 865.37 26.435
                                                < 2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 5915 on 291 degrees of freedom
## Multiple R-squared: 0.7379, Adjusted R-squared: 0.7307
## F-statistic: 102.4 on 8 and 291 DF, p-value: < 2.2e-16
```

5.1 Coefficients:

 $\begin{array}{l} {\rm charges = -12033.25 + (age)} \ 261.14 + (children) \\ {\rm 532.76 + (bmi)} \ 353.35 + (region_northwest) \\ {\rm (-1545.53)} + (region_southeast) \\ {\rm (-1505.35)} + (regionsouthwest) \\ {\rm (-1719.90)} + (sexmale) \\ {\rm (607.78)} \\ {\rm + (smokeryes)^*(22876.38)} \\ \end{array}$

• The **Estimate column** lists the estimated coefficients of the predictor variables included in the model. For instance, the "age" coefficient has an estimated value of 261.14, which means that for every one-unit increase in age, the outcome variable (presumably a medical cost) is estimated to increase by \$261.14, holding all other variables constant.

5.2 Standard Errors:

• The **Std Error** column lists the standard errors of the estimated coefficients. These are measures of the uncertainty or variability in the estimated coefficients. Smaller standard errors indicate more precise estimates.

5.3 T-values:

• The **t-value** column lists the t-statistics for each coefficient. These values represent the estimated coefficients divided by their standard errors. T-values are used to test the null hypothesis that the true coefficient is zero. Larger t-values indicate a stronger evidence against the null hypothesis.

5.4 P-values

- The pr column lists the p-values associated with the t-values. P-values represent the probability of observing the t-value or a more extreme value if the true coefficient is zero. Smaller p-values indicate stronger evidence against the null hypothesis.
- The significance codes provided in the table help to quickly identify significant coefficients; for instance, age,bmi,smokers represents p-value less than two decimal places and children represents p-value less than 0.05. One can say that region represents bigger p-value than significant level.

5.5 Residual Standard Error:

 The Residual standard error provides an estimate of the variability of the errors or unexplained variance in the model. It measures the average distance that the observed values fall from the predicted values.

5.6 R-squared:

• The "Multiple R-squared" and "Adjusted R-squared" measures how well the model fits the data. R-squared ranges from 0 to 1 and represents the proportion of variance in the outcome variable that is explained by the predictor variables. Adjusted R-squared is a corrected version of R-squared that takes into account the number of predictor variables in the model. In full model one can see 0.7379.

5.7 summary

• This data summary provides information on the estimated coefficients, their standard errors, t-values, and corresponding p-values, as well as model diagnostics such as residual standard error, multiple R-squared, adjusted R-squared, and F-statistic. These measures can be used to interpret the strength and significance of the relationship between the predictor variables and the outcome variable, as well as the overall goodness-of-fit of the model. Still working to find the best parameters to be included in the model.

6 Confidence intervals for all variables used in full model

- These intervals provide a range of plausible values for the true population coefficients based on the sample data. The first column, "2.5%", represents the lower bound of the interval, and the second column, "97.5%", represents the upper bound of the interval.
- The 95% confidence interval for the "age" variable is [213.81, 308.48]. This means that we are 95% confident that the true population coefficient for age falls within this range. Similarly, the confidence interval for the intercept is [-16276.39, -7790.11], which suggests that the expected value of the response variable (when all predictor variables are 0) is likely to be within this range.

```
confint(AB, level = 0.95)
##
                           2.5 %
                                      97.5 %
## (Intercept)
                    -16276.39288 -7790.1053
                       213.80684
                                    308.4817
## age
## children
                       -16.88546
                                   1082.3959
## bmi
                       231.26578
                                    475.4327
## regionnorthwest
                     -3498.86525
                                    407.8049
## regionsoutheast
                     -3544.78722
                                    534.0830
## regionsouthwest
                     -3668.78747
                                    228.9853
```

1974.8375

7 Reduced Model

sexmale

smokeryes

7.1 Evaluating Various regression models.

-759.27612

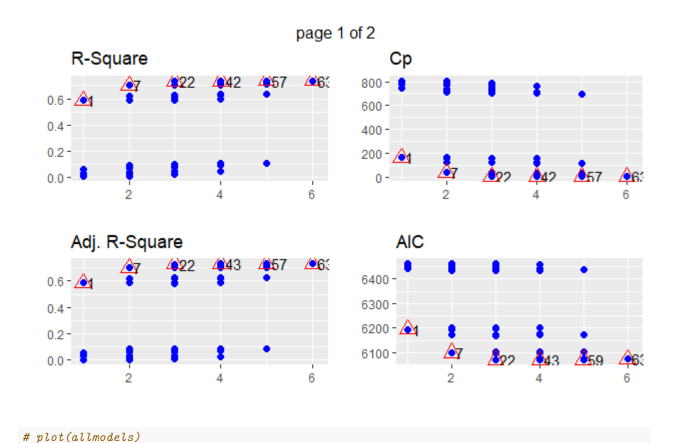
21173.19777 24579.5601

• Let us begin by using a multiple linear regression model that uses all the six variables. From the summary table we see that our R squared and Adjusted r square are around 0.73 and the residual standard error is 5915. The r squared value is high enough to be considered good but let us continue finding better fits.

```
ols_step_all_possible(AB) -> allmodels
as_tibble(allmodels) -> allmodels_1
tail(allmodels_1,8)
```

```
## # A tibble: 8 x 14
##
     mindex
                n predict~1 rsquare
                                       adjr predrsq
                                                         ср
                                                              aic
                                                                   sbic
                                                                                  msep
##
      <int> <int> <chr>
                               <dbl>
                                      <dbl>
                                               <dbl>
                                                      <dbl> <dbl> <dbl> <dbl> <dbl>
                                                                                 <dbl>
## 1
         56
                4 children~
                              0.0470 0.0275 0.00150 768.
                                                            6457. 5593. 6486. 3.77e10
## 2
         57
                              0.737
                                            0.721
                                                       3.77 6072. 5217. 6105. 1.04e10
                5 age chil~
                                     0.731
## 3
                                     0.728
                                            0.718
                                                       6.64 6075. 5220. 6108. 1.05e10
                5 age bmi ~
                              0.735
## 4
         59
                5 age chil~
                              0.735
                                     0.730
                                            0.722
                                                       6.70 6071. 5220. 6097. 1.05e10
                                     0.702
                                                            6103. 5247. 6136. 1.15e10
## 5
         60
                5 age chil~
                              0.709
                                            0.690
                                                      35.4
## 6
         61
                                            0.609
                5 children~
                              0.632
                                     0.623
                                                     121.
                                                            6173. 5315. 6207. 1.46e10
## 7
                                     0.0873 0.0600
                                                            6439. 5574. 6472. 3.53e10
                5 age chil~
                              0.109
                                                     702.
                                                            6073. 5218. 6110. 1.04e10
## 8
         63
                6 age chil~
                              0.738
                                     0.731
                                            0.720
                                                       5
     ... with 3 more variables: fpe <dbl>, apc <dbl>, hsp <dbl>, and abbreviated
       variable name 1: predictors
```

the best model is represented by including all age, children, bmi, region, smoker full model. knitr::include_graphics("C:/Users/Dhruv Jain/Desktop/1.png")



7.2 Finding Reduce model

• Let us see how r will create dummy variables for us using Region variable. R will convert the categorical variables for us We can see R will do this automatically for us creating these dummy variables. Northeast is essentially the control variable. When the three variables are 0, it means that region is northeast! No need to bother here as R does this for us.

```
contrasts(as.factor(insurance_new$region))
```

```
## northwest southeast southwest
## northeast 0 0 0
## northwest 1 0 0
## southeast 0 1 0
## southwest 0 0 1
```

```
# Next step - Let us include all the interaction terms as well. Our residual standard error reduced. si
```

```
lm(charges~ age + children + bmi +
    region + sex + smoker + age:children + age:bmi + age:region + age:sex + age:smoker+
    children:bmi + children:region + children:sex + children:smoker
    + bmi:region + bmi:sex+ bmi:smoker + region:sex + region + smoker
    + sex:smoker, insurance_new) -> interactionModel
summary(interactionModel)
```

```
##
## Call:
## lm(formula = charges ~ age + children + bmi + region + sex +
       smoker + age:children + age:bmi + age:region + age:sex +
##
##
       age:smoker + children:bmi + children:region + children:sex +
##
       children:smoker + bmi:region + bmi:sex + bmi:smoker + region:sex +
       region + smoker + sex:smoker, data = insurance new)
##
##
## Residuals:
##
      Min
                1Q Median
                                3Q
                                       Max
   -8196.3 -2253.3 -1060.8
                             274.9 20420.1
##
## Coefficients:
##
                              Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                                         5950.715 -0.476 0.634228
                             -2834.483
## age
                               146.188
                                          117.074
                                                    1.249 0.212868
## children
                                         1533.089 -0.034 0.972835
                               -52.254
## bmi
                                98.895
                                         196.858
                                                    0.502 0.615821
                                                    0.911 0.362857
## regionnorthwest
                              4940.926
                                       5420.759
## regionsoutheast
                              1791.263
                                        5455.771
                                                    0.328 0.742922
## regionsouthwest
                               485.686
                                       5074.751
                                                    0.096 0.923825
## sexmale
                             -1228.344
                                         3662.175 -0.335 0.737574
## smokeryes
                            -17140.502
                                         4375.582 -3.917 0.000114 ***
## age:children
                                                    1.337 0.182338
                                25.113
                                           18.782
## age:bmi
                                1.850
                                           3.667
                                                    0.504 0.614324
## age:regionnorthwest
                                60.038
                                           59.631
                                                    1.007 0.314922
## age:regionsoutheast
                                           61.319
                                                   1.126 0.261106
                                69.054
## age:regionsouthwest
                                98.009
                                           58.561
                                                    1.674 0.095368
## age:sexmale
                                           41.292 -0.537 0.592007
                               -22.156
## age:smokeryes
                               -57.725
                                           53.308 -1.083 0.279845
## children:bmi
                               -13.009
                                           43.378 -0.300 0.764478
## children:regionnorthwest
                               105.184
                                          740.993
                                                   0.142 0.887226
## children:regionsoutheast
                              -318.240
                                          763.888 -0.417 0.677299
## children:regionsouthwest
                              -233.540
                                          712.265 -0.328 0.743255
## children:sexmale
                               553.118
                                          487.168
                                                   1.135 0.257230
                                          638.980 -1.908 0.057521 .
## children:smokeryes
                             -1218.854
## bmi:regionnorthwest
                              -260.593
                                          159.656 -1.632 0.103803
## bmi:regionsoutheast
                              -120.067
                                          154.970 -0.775 0.439153
## bmi:regionsouthwest
                                          156.511 -1.032 0.303190
                              -161.455
## bmi:sexmale
                                                   0.971 0.332616
                               105.175
                                          108.360
## bmi:smokeryes
                              1459.039
                                         134.821 10.822 < 2e-16 ***
## regionnorthwest:sexmale
                                         1733.496 -0.528 0.597929
                              -915.301
## regionsoutheast:sexmale
                             -3533.510
                                         1791.412 -1.972 0.049580 *
## regionsouthwest:sexmale
                             -2069.808
                                         1698.494 -1.219 0.224059
## sexmale:smokeryes
                              -380.218
                                         1604.912 -0.237 0.812908
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Residual standard error: 4910 on 269 degrees of freedom
## Multiple R-squared: 0.8331, Adjusted R-squared: 0.8144
## F-statistic: 44.75 on 30 and 269 DF, p-value: < 2.2e-16
```

Let us also see if transforming our dependent variable might help. Our R squared increased slightly # but not much

```
lm(log(charges)~ age + children + bmi +
    region + sex + smoker + age:children + age:bmi + age:region + age:sex + age:smoker+
    children:bmi + children:region + children:sex + children:smoker
  + bmi:region + bmi:sex+ bmi:smoker + region:sex + region + smoker
  + sex:smoker, insurance_new) -> interactionModel
summary(interactionModel)
##
## Call:
## lm(formula = log(charges) ~ age + children + bmi + region + sex +
##
      smoker + age:children + age:bmi + age:region + age:sex +
##
      age:smoker + children:bmi + children:region + children:sex +
##
      children:smoker + bmi:region + bmi:sex + bmi:smoker + region:sex +
##
      region + smoker + sex:smoker, data = insurance_new)
##
## Residuals:
##
       Min
                1Q
                    Median
                                 30
                                        Max
## -0.56846 -0.16881 -0.07365 0.03661 2.07636
##
## Coefficients:
##
                           Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                          6.5081702  0.4688489  13.881  < 2e-16 ***
                          0.0404944 0.0092241
                                              4.390 1.63e-05 ***
## age
## children
                          0.2323820 0.1207900 1.924 0.055428
## bmi
                          0.0373892 0.0155102 2.411 0.016596 *
## regionnorthwest
                        0.1858655 0.4270944 0.435 0.663776
                         0.0443852 0.4298529 0.103 0.917836
## regionsoutheast
                         -0.1660037 0.3998328 -0.415 0.678339
## regionsouthwest
## sexmale
                        -0.3419357 0.2885379 -1.185 0.237038
## smokeryes
                         1.5843978  0.3447462  4.596  6.63e-06 ***
## age:children
                         -0.0027142 0.0014798 -1.834 0.067741
## age:bmi
                         -0.0003435 0.0002889 -1.189 0.235491
## age:regionnorthwest
                        0.0125558 0.0046982 2.672 0.007990 **
                        0.0145449 0.0048312
## age:regionsoutheast
                                               3.011 0.002855 **
                        0.0165099 0.0046140
## age:regionsouthwest
                                               3.578 0.000410 ***
## age:sexmale
                          0.0040348 0.0032533 1.240 0.215983
## age:smokeryes
                         ## children:bmi
                         ## children:regionnorthwest -0.0022434 0.0583819 -0.038 0.969377
## children:regionsoutheast -0.0215457 0.0601857 -0.358 0.720634
## children:regionsouthwest -0.0263279 0.0561184 -0.469 0.639343
                          0.0645546 0.0383833 1.682 0.093760 .
## children:sexmale
## children:smokeryes
                         ## bmi:regionnorthwest
                         ## bmi:regionsoutheast
                         -0.0213882 0.0122099 -1.752 0.080963 .
## bmi:regionsouthwest
                         -0.0213983 0.0123313 -1.735 0.083835
                          0.0049372 0.0085376 0.578 0.563549
## bmi:sexmale
## bmi:smokerves
                        0.0495150 0.0106224 4.661 4.95e-06 ***
## regionnorthwest:sexmale 0.0331110 0.1365798 0.242 0.808632
## regionsoutheast:sexmale -0.2677983 0.1411430 -1.897 0.058852 .
## regionsouthwest:sexmale -0.1418041 0.1338221 -1.060 0.290256
## sexmale:smokeryes
                          0.0337446 0.1264488
                                              0.267 0.789778
## ---
```

```
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.3869 on 269 degrees of freedom
## Multiple R-squared: 0.8363, Adjusted R-squared: 0.818
## F-statistic: 45.81 on 30 and 269 DF, p-value: < 2.2e-16</pre>
```

• Let us add more interaction terms that include transformed x variables our y variable has been transformed here as well. Our r squared increased again to 0.8373 and the residual error is now 4839. It is better slightly but there are a lot of variables which are not significant. For example, sex:smoke interaction variable's p value is 0.7829 which is significantly above 0.05. In the next step, let us remove all those variables that are not significant

```
lm(charges~ age + children + bmi +
     region + sex + smoker + age:children + age:bmi + age:region + age:sex + age:smoker+
      children:sex + children:smoker
   + bmi:region + bmi:sex+ bmi:smoker + region:sex + region + smoker
   + sex:smoker + log(bmi)+ bmi*bmi + log(age) + age*age + log(age)*log(bmi), insurance_new) -> interac
summary(interactionModel)
##
## Call:
## lm(formula = charges ~ age + children + bmi + region + sex +
##
       smoker + age:children + age:bmi + age:region + age:sex +
       age:smoker + children:sex + children:smoker + bmi:region +
##
##
       bmi:sex + bmi:smoker + region:sex + region + smoker + sex:smoker +
##
       log(bmi) + bmi * bmi + log(age) + age * age + log(age) *
       log(bmi), data = insurance_new)
##
##
## Residuals:
                1Q Median
##
       Min
  -7396.2 -2274.3 -911.3
                             237.6 19910.0
##
## Coefficients:
                             Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                             -496.094 225079.381 -0.002
                                                            0.9982
## age
                               99.734
                                          588.741
                                                    0.169
                                                            0.8656
## children
                             -414.881
                                          841.634 -0.493
                                                            0.6225
## bmi
                             1080.305
                                         931.509
                                                   1.160
                                                            0.2472
## regionnorthwest
                             5496.631
                                        5208.089
                                                    1.055
                                                            0.2922
## regionsoutheast
                             6158.931
                                        5612.118
                                                    1.097
                                                            0.2734
## regionsouthwest
                             1725.464
                                         4974.472
                                                    0.347
                                                            0.7290
## sexmale
                             -772.434
                                        3586.073 -0.215
                                                            0.8296
## smokeryes
                           -17224.875
                                         4309.003 -3.997 8.27e-05 ***
## log(bmi)
                            -5259.620 73591.622 -0.071
                                                            0.9431
## log(age)
                            22322.993
                                       67679.455
                                                    0.330
                                                            0.7418
                                                            0.2084
## age:children
                               22.167
                                           17.580
                                                    1.261
## age:bmi
                                8.347
                                           18.669
                                                    0.447
                                                            0.6551
## age:regionnorthwest
                               74.566
                                           59.294
                                                    1.258
                                                            0.2096
## age:regionsoutheast
                                           60.104
                                                            0.2718
                               66.190
                                                    1.101
## age:regionsouthwest
                               94.718
                                           57.672
                                                    1.642
                                                            0.1017
## age:sexmale
                              -21.656
                                           41.180 -0.526
                                                            0.5994
## age:smokeryes
                              -68.335
                                           51.950 -1.315
                                                            0.1895
## children:sexmale
                              691.627
                                          473.110
                                                    1.462
                                                            0.1449
```

```
## children:smokeryes
                            -1197.148
                                         625.331 -1.914
                                                           0.0566 .
                                         155.290 -1.940
## bmi:regionnorthwest
                             -301.338
                                                           0.0534 .
## bmi:regionsoutheast
                             -276.348
                                         162.749 -1.698
                                                           0.0907 .
## bmi:regionsouthwest
                                         155.430 -1.296
                                                           0.1962
                             -201.368
## bmi:sexmale
                               91.781
                                         105.495
                                                  0.870
                                                          0.3851
## bmi:smokeryes
                             1473.976
                                        132.003 11.166
                                                         < 2e-16 ***
## regionnorthwest:sexmale
                            -667.267
                                       1702.405 -0.392
                                                          0.6954
## regionsoutheast:sexmale
                           -3302.764
                                        1780.210 -1.855
                                                           0.0646 .
## regionsouthwest:sexmale
                            -2344.500
                                        1670.199
                                                 -1.404
                                                           0.1615
## sexmale:smokeryes
                            -430.457
                                        1560.915 -0.276
                                                           0.7829
## log(bmi):log(age)
                            -8223.943
                                      19871.918 -0.414
                                                           0.6793
## --
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 4839 on 270 degrees of freedom
## Multiple R-squared: 0.8373, Adjusted R-squared: 0.8198
## F-statistic: 47.9 on 29 and 270 DF, p-value: < 2.2e-16
```

7.3 Let eliminate all interaction

- Terms or the single whose p value is insignificant. For this case our starting model has a lot of variables so after removing many of the variables we will have a reduced model.
- Those variables whose p value is too high should be removed and a reduced model has to be produced.we run the reduced model below our r squared slightly decreased to 0.8332 and the adjusted r squared to 0.8244.
- It is very small change to our previous step so it is fine to take this. The p values are also significant so for that reason we chose this model. This could be one candidate model for our regression.

```
lm(charges~ children+bmi+smoker+age:children+age:region+
    age:smoker+children:smoker+bmi:region+bmi:smoker+smoker+log(bmi)+
    bmi*bmi + log(age),insurance_new) -> interactionModelSignificant
summary(interactionModelSignificant)
```

```
##
## lm(formula = charges ~ children + bmi + smoker + age:children +
##
       age:region + age:smoker + children:smoker + bmi:region +
       bmi:smoker + smoker + log(bmi) + bmi * bmi + log(age), data = insurance_new)
##
##
## Residuals:
       Min
                                3Q
##
                1Q Median
                                       Max
## -7189.3 -2334.9 -1166.7
                             385.6 21603.4
##
## Coefficients:
##
                        Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                        55898.45
                                   31237.64
                                              1.789 0.07461
## children
                         -319.02
                                     745.49 -0.428 0.66902
## bmi
                         1285.83
                                     454.43
                                              2.830 0.00499 **
## smokeryes
                                    4145.36 -4.142 4.55e-05 ***
                       -17169.12
                                            -2.357 0.01911 *
## log(bmi)
                       -30700.93
                                   13026.61
## log(age)
                         5125.69
                                    1406.95
                                             3.643 0.00032 ***
```

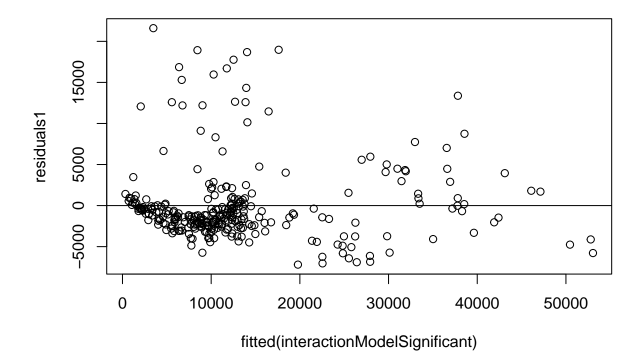
```
## children:age
                          25.99
                                     17.10
                                             1.520 0.12966
                         136.02
                                             2.664 0.00815 **
## age:regionnorthwest
                                     51.05
## age:regionsoutheast
                         122.86
                                     51.49
                                             2.386 0.01769 *
## age:regionsouthwest
                                     52.25
                                             2.602 0.00976 **
                         135.95
## smokeryes:age
                         -71.61
                                     50.30
                                            -1.424
                                                    0.15563
## children:smokeryes
                                            -1.853 0.06487
                       -1125.32
                                    607.17
## bmi:regionnorthwest
                        -218.28
                                     75.92
                                            -2.875 0.00434 **
## bmi:regionsoutheast
                        -213.74
                                     70.80
                                            -3.019 0.00277 **
## bmi:regionsouthwest
                        -246.50
                                     74.75
                                            -3.297
                                                    0.00110 **
## bmi:smokeryes
                        1468.40
                                    125.32 11.717 < 2e-16 ***
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
## Residual standard error: 4858 on 284 degrees of freedom
## Multiple R-squared: 0.8275, Adjusted R-squared: 0.8184
## F-statistic: 90.81 on 15 and 284 DF, p-value: < 2.2e-16
```

- let also use the variables from the last step to establish model selection like we have covered in class R will use any combination of these variables to come up with r^2, cp values etc for each of them.we will order the result by r^2, then by adjusted then by cp and inspect if we should choose any other combination of the variables from above.
- As we can see the maximum r square values are around 0.833 (from 2047 models) which aligns with our finding from above. So need to change anything. Since there are a lot of models whose r squared is 0.833 we might choose the one with the lowest cp. And that is the 6th row with cp of 5.16. The variables used are smoker log(bmi) log(age) age:region smoker:age children:smoker bmi:region bmi:smoker with 8 variables. So that could be an alternative reduced version of the model we have above.
- So going forward let me choose the model from above. We felt it was good enough so now let us investigate more by analysing the residuals and the normality.

```
lm(charges~ children + bmi +
    smoker + age:children + age:region + age:smoker+
    + children:smoker + bmi:region + bmi:smoker + smoker
    + log(bmi)+ bmi*bmi + log(age) , insurance_new) -> interactionModelSignificant
summary(interactionModelSignificant)
```

```
##
## Call:
## lm(formula = charges ~ children + bmi + smoker + age:children +
##
       age:region + age:smoker + +children:smoker + bmi:region +
       bmi:smoker + smoker + log(bmi) + bmi * bmi + log(age), data = insurance_new)
##
##
## Residuals:
                1Q Median
##
       Min
                                3Q
                                       Max
   -7189.3 -2334.9 -1166.7
                             385.6 21603.4
##
##
## Coefficients:
##
                        Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                        55898.45
                                   31237.64
                                              1.789 0.07461 .
## children
                         -319.02
                                     745.49 -0.428 0.66902
## bmi
                         1285.83
                                     454.43
                                             2.830 0.00499 **
                                    4145.36 -4.142 4.55e-05 ***
## smokeryes
                       -17169.12
```

```
## log(bmi)
                     -30700.93 13026.61 -2.357 0.01911 *
                       5125.69 1406.95 3.643 0.00032 ***
## log(age)
## children:age
                       25.99
                                    17.10 1.520 0.12966
                                    51.05 2.664 0.00815 **
## age:regionnorthwest 136.02
## age:regionsoutheast 122.86
                                    51.49
                                          2.386 0.01769 *
## age:regionsouthwest
                                    52.25 2.602 0.00976 **
                      135.95
## smokeryes:age
                                    50.30 -1.424 0.15563
                        -71.61
## children:smokeryes
                                   607.17 -1.853 0.06487 .
                      -1125.32
                                   75.92 -2.875 0.00434 **
## bmi:regionnorthwest
                      -218.28
## bmi:regionsoutheast -213.74
                                    70.80 -3.019 0.00277 **
## bmi:regionsouthwest -246.50
                                    74.75 -3.297 0.00110 **
                                   125.32 11.717 < 2e-16 ***
## bmi:smokeryes
                       1468.40
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Residual standard error: 4858 on 284 degrees of freedom
## Multiple R-squared: 0.8275, Adjusted R-squared: 0.8184
## F-statistic: 90.81 on 15 and 284 DF, p-value: < 2.2e-16
# The residual plot is not perfect but there is no clear pattern. So it should be relatively fine. It i
# Observing the residual vs fitted plot one can say that more points are plotted in one area specifical
resid(interactionModelSignificant) -> residuals1
plot(fitted(interactionModelSignificant), residuals1)+
abline(0,0)
```



integer(0)

7.4 CI for two of our chosen quantitative variables

- Using this regression model Let us take at least two variables and find confidence interval for the independent variables. Let us take two such as log(age) and bmi for example.
- The standard error for $\log(\text{age})$ is 1406.95 and the coefficient is 5125.69 and standard error for bmi is 454 and the coefficient is 1285.83.
- Degree of freedom is 284. (300-16). t is 1.9683. Which is closer to the z score actually.

```
qt(p=.025, df=284, lower.tail = FALSE) -> t
t
```

[1] 1.968352

```
# so for bmi
#upper bound 2179.438
1285.83 + 1.9683 * 454
```

[1] 2179.438

```
# lower bound 392.2218

1285.83 - 1.9683 * 454

## [1] 392.2218

# for log(age)
# upper bound ] 7894.99
5125.69 + 1.9683 * 1406.95

## [1] 7894.99

# lower bound
5125.69 - 1.9683 * 1406.95

## [1] 2356.39
```

8

8.1 Researching and Applying a model analysis.

- Here we will show how random forest model can be used on our full model or using all explanatory variables to predict the dependent variable charges.
- Random forest works by using if-else decision trees using all variables. For example one possible path could be if a person is a non smoker, male and if he has a bmi value above 90, the medical charge should approximately be 1000. This is just to. show how it works under the hood but the actual mechanisms and branching rules are not as trivial as my example. For a continuous dependent variable, we should use random forest regressor (it does regression but using random forest)
- We will compare the root mean squared error using our random forest model and the regression we had above Random Forest Model.

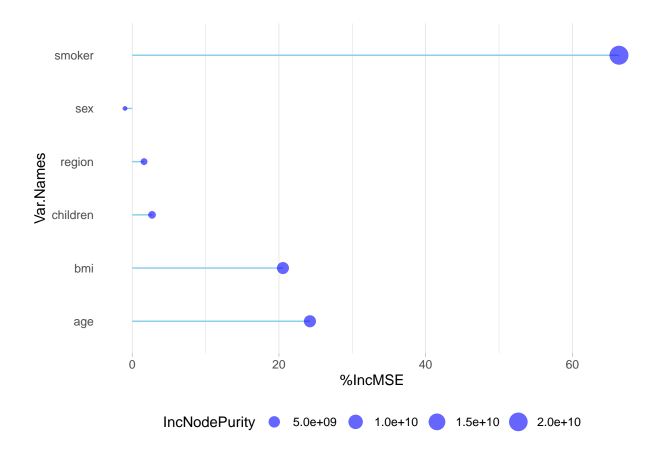
- from fitting the random forest model, we can see that the root mean squared is 37712690^(0.5). Rf.fit gives us the squared value so we have to take the root of it to find the root mean square. so 37712690^(0.5) = 6141.066. This random forest model produced worse result than our regression model.
- we got a value of 4858 as our best root mean squared error from our regressoin model.
- But also just like we can tune our regression model, we can also tune our random forest model. Let us increase the number of trees from 3 to 300 in the random forest model. we get 25902511^(0.5) = 5089.451. So as we increase our number of trees the root mean squared approached our best regression model out put. Of course we can tune a lot of things in the decision trees of random forest as well so random might give us a root mean square value less than our regression model.
- The variability explained by this random forest model was also close to what we have in the regression model. It is 80% here.

```
rf.fit
##
## Call:
    randomForest(formula = charges ~ ., data = insurance_new, ntree = 3,
                                                                                keep.forest = FALSE, impor
                  Type of random forest: regression
##
##
                        Number of trees: 3
## No. of variables tried at each split: 2
##
##
             Mean of squared residuals: 41785075
##
                       % Var explained: 67.73
rf.fit <- randomForest(charges ~ ., data=insurance_new, ntree=300,
                       keep.forest=FALSE, importance=TRUE)
rf.fit
##
## Call:
    randomForest(formula = charges ~ ., data = insurance_new, ntree = 300,
##
                                                                                  keep.forest = FALSE, im
                  Type of random forest: regression
##
##
                        Number of trees: 300
## No. of variables tried at each split: 2
##
##
             Mean of squared residuals: 26152008
                       % Var explained: 79.81
##
```

- Let also this which variables were important according to the latest random forest model.
- we had. We see that smoker variable was very important in terms of information gain (it is one of the best variable used in the decision tree and is found to be important in terms of determining medical charges/bills. The other variables that are found important are age and BMI as we see from the diagram.) children, region and sex had minimal impact compared to the other variables.

```
ImpData <- as.data.frame(importance(rf.fit))
ImpData$Var.Names <- row.names(ImpData)

ggplot(ImpData, aes(x=Var.Names, y=`%IncMSE`)) +
   geom_segment( aes(x=Var.Names, xend=Var.Names, y=0, yend=`%IncMSE`), color="skyblue") +
   geom_point(aes(size = IncNodePurity), color="blue", alpha=0.6) +
   theme_light() +
   coord_flip() +
   theme(
    legend.position="bottom",
    panel.grid.major.y = element_blank(),
    panel.border = element_blank(),
    axis.ticks.y = element_blank()
)</pre>
```



9 Final summary

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