**Linear Regression**

Linear Regression is one of the most fundamental algorithms in the Machine Learning world. It is the door to the magical world ahead. But before proceeding with the algorithm, let’s first discuss the lifecycle of any machine learning model. This diagram explains the creation of a Machine Learning model from scratch and then taking the same model further with hyper parameter tuning to increase its accuracy, deciding the deployment strategies for that model and once deployed setting up the logging and monitoring frameworks to generate reports and dashboards based on the client requirements. A typical lifecycle diagram for a machine learning model looks like:

**What is Regression Analysis?**

Regression in statistics is the process of predicting a Label(or Dependent Variable) based on the features(Independent Variables) at hand. Regression is used for time series modelling and finding the causal effect relationship between the variables and forecasting. For example, the relationship between the stock prices of the company and various factors like customer reputation and company annual performance etc. can be studied using regression.

Regression analysis is an important tool for analysing and modelling data. Here, we fit a curve/line to the data points, in such a manner that the differences between the distance of the actual data points from the plotted curve/line is minimum. The topic will be explained in detail in the coming sections

**The use of Regression**

Regression analyses the relationship between two or more features. Let’s take an example:

Let’s suppose we want to make an application which predicts the chances of admission a student to a foreign university. In that case, the

The benefits of using Regression analysis are as follows:

* It shows the significant relationships between the Label (dependent variable) and the features(independent variable).
* It shows the extent of the impact of multiple independent variables on the dependent variable.
* It can also measure these effects even if the variables are on a different scale.

These features enable the data scientists to find the best set of independent variables for predictions.

**Linear Regression**

Linear Regression is one of the most fundamental and widely known Machine Learning Algorithms which people start with. Building blocks of a Linear Regression Model are:

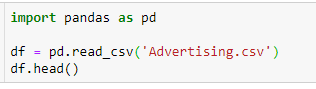
* Discreet/continuous independent variables
* A best-fit regression line.
* Minimizing the error
* Continuous dependent variable. i.e., A Linear Regression model predicts the dependent variable using a regression line based on the independent variables. The equation of the Linear Regression is:

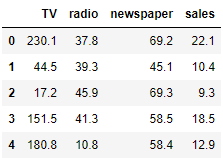
y = mx + c + e

Where, c is the intercept, m is the slope of the line, and e is the error term. The equation above is used to predict the value of the target variable based on the given predictor variable(s).

**The Problem statement:**

This data is about the amount spent on advertising through different channels like TV, Radio and Newspaper. The goal is to predict how the expense on each channel affects the sales and is there a way to optimise that sale?





What are the **features**?

* TV: Advertising dollars spent on TV for a single product in a given market (in thousands of dollars)
* Radio: Advertising dollars spent on Radio
* Newspaper: Advertising dollars spent on Newspaper

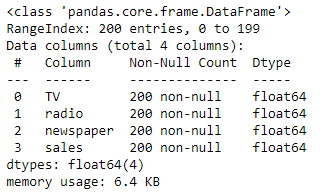
What is the **response**?

* Sales: sales of a single product in a given market (in thousands of widgets)

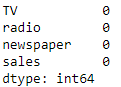


(200, 4)



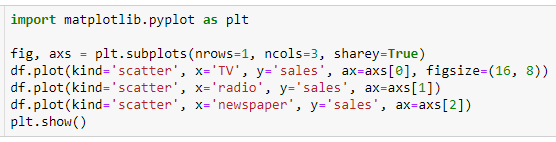


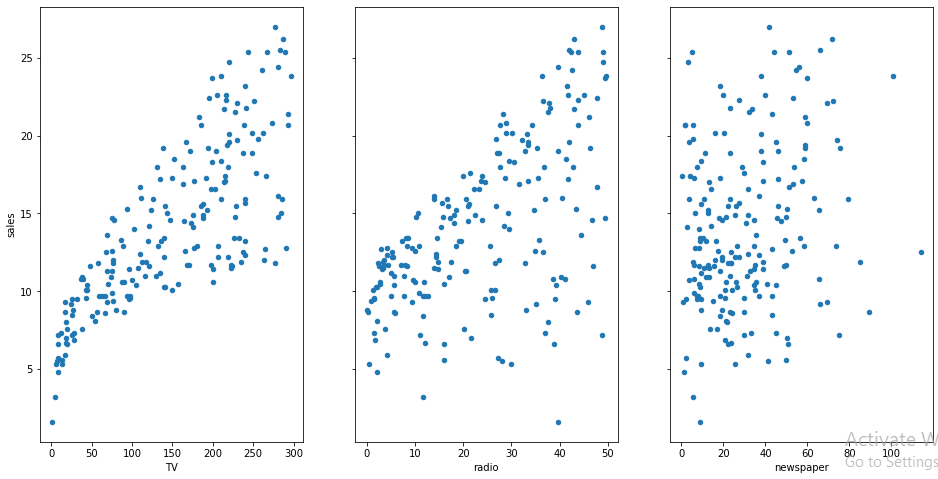
**finding the count of missing values from different columns**



Now, let's showcase the relationship between the feature and target column

**visualize the relationship between the features and the response using scatterplots**



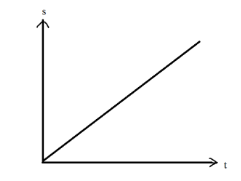


**Questions about the data**

A generic question shall be: How the company should optimise the spends on advertising to maximise the sales?

These general questions might lead you to more specific questions:

1. What’s the relationship between ads and sales?
2. How prominent is that relationship?
3. Which ad types contribute to sales?
4. How each ad contributes to sales?
5. Can sales be predicted based on the expense of the advertisement?



From the relationship diagrams above, it can be observed that there seems to be a linear relationship between the features TV ad, Radio ad and the sales is almost a linear one. A linear relationship typically looks like:

Hence, we can build a model using the Linear Regression Algorithm.

**Simple Linear Regression**

Simple Linear regression is a method for predicting a **quantitative response** using a **single feature** ("input variable").

The mathematical equation is: y = β0 + β1x

What do terms represent?

* y is the response or the target variable
* x is the feature
* β1 is the coefficient of x
* β0 is the intercept

β0 and β1 are the **model coefficients**. To create a model, we must "learn" the values of these coefficients. And once we have the value of these coefficients, we can use the model to predict the Sales!

**Estimating ("Learning") Model Coefficients**

The coefficients are estimated using the **least-squares criterion**, i.e., the best fit line has to be calculated that minimizes the **sum of squared residuals** (or "sum of squared errors").

3 methods to calculate co-efficient and bias:

* Ordinary Least Square



* From sklearn.linear\_model import LinearRegression

lr\_model = LinearRegression()

lr\_model.fit(X, y)

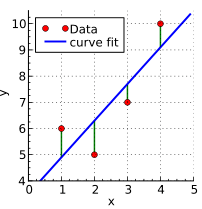
lr\_model.intercept\_

lr\_model.coef\_

* From scipy.linalg import lstsq

sp\_params, residue, rank, s = lstsq(X, y)

print(sp\_params)

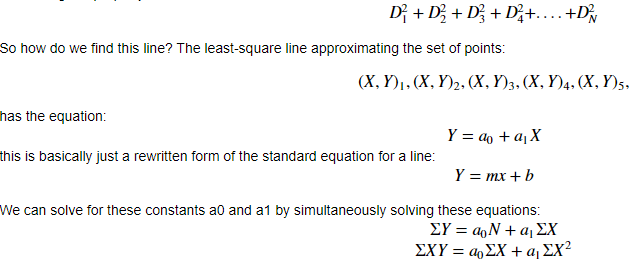
**The mathematics involved**

Take a quick look at the plot created. Now consider each point, and know that each of them has a coordinate in the form (X, Y). Now draw an imaginary line between each point and the current "best-fit" line. We'll call the distance between each point and the current best-fit line as D. To get a quick image of what we're trying to visualize, take a look at the picture below:

What elements are present in the diagram?

* The red points are the **observed values** of x and y.
* The blue line is the **least squares line**.
* The green lines are the **residuals**, which is the distance between the observed values and the least squares line.

Before, we're labelling each green line as having a distance D, and each red point as having a coordinate of (X, Y). Then we can define our best fit line as the line having the property were:



These are called the normal equations for the least-squares line. There are further steps that can be taken in rearranging these equations to solve for y, but we'll let SciKit-learn do the rest of the heavy lifting here.

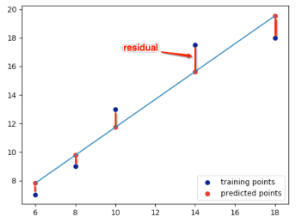
**Let’s see the underlying assumptions: -**

* The regression model is linear in terms of coefficients and error term.
* The mean of the residuals is zero.
* The error terms are not correlated with each other, i.e. given an error value; we cannot predict the next error value.
* The independent variables(x) are uncorrelated with the residual term, also termed as **exogeneity**. This, in layman term, generalises that in no way should the error term be predicted given the value of independent variables.
* The error terms have a constant variance, i.e. **homoscedasticity**.
* No Multicollinearity, i.e. no independent variables should be correlated with each other or affect one another. If there is multicollinearity, the precision of prediction by the OLS model decreases.
* The error terms are normally distributed.

The general equation of a straight line is:

𝑦 = mx + c

It means that if we have the value of m and b, we can predict all the values of y for corresponding x. During construction of a Linear Regression Model, the computer tries to calculate the values of m and c to get a straight line. But the question is:

**How do you Know this is the best fit line?**

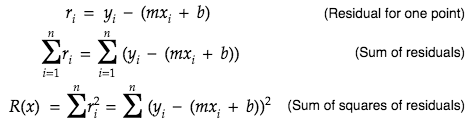
The best fit line is obtained by minimizing the residual.

Residual is the distance between the actual Y and the predicted Y, as shown below:

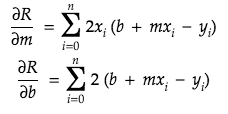
Mathematically, Residual is:

r = y − (mx + b)

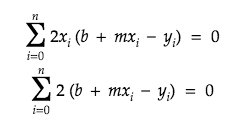
Hence, the sum of the square of residuals is:

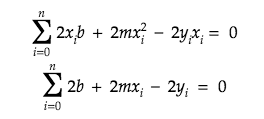


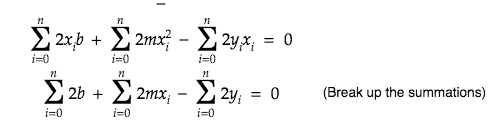
As we can see that the residual is both a function of m and b, so differentiating partially with respect to m and b will give us:

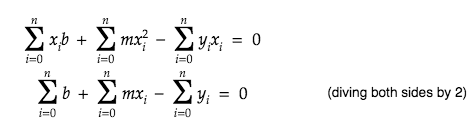


For getting the best fit line, residual should be minimum. The minima of a function occurs where the derivative=0. So, equating our corresponding derivatives to 0, we get:

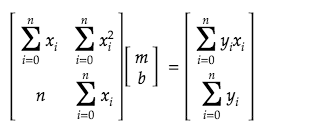






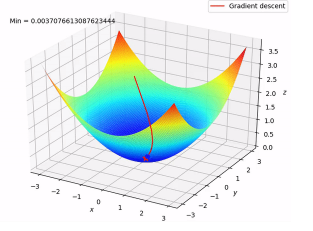


This same equation can be written in matrix form as:

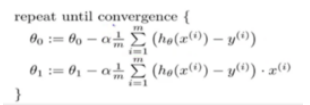


Ideally, if we'd have an equation of one dependent and one independent variable the minima will look as follows:

But as the residual's minima is dependent on two variables m and b, it becomes a Paraboloid and the appropriate m and b are calculated using Gradient Descent as shown below:



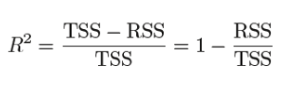
Now, let’s understand how to check, how well the model fits our data. The new values for 'slope' and 'intercept' are calculated as follows:



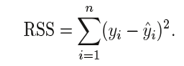
where, θ0 is 'intercept' , θ1 is the slope, αα is the learning rate, m is the total number of observations and the term after the ∑ sign is the loss. Google Tensor board recommends a Learning rate between 0.00001 and 10. Generally a smaller learning rate is recommended to avoid overshooting while creating a model.

### **R2 statistics**

The R-squared statistic provides a measure of fit. It takes the form of a proportion—the proportion of variance explained—and so it always takes on a value between 0 and 1. In simple words, it represents how much of our data is being explained by our model. For example, R2 statistic = 0.75, it says that our model fits 75 % of the total data set. Similarly, if it is 0, it means none of the data points is being explained and a value of 1 represents 100% data explanation. Mathematically R2 statistic is calculated as :

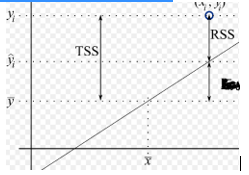


Where RSS: is the Residual Sum of squares and is given as :



RSS is the residual(error) term we have been talking about so far. And, TSS: is the Total sum of squares and given as :



TSS is calculated when we consider the line passing through the mean value of y, to be the best fit line. Just like RSS, we calculate the error term when the best fit line is the line passing through the mean value of y and we get the value of TSS.

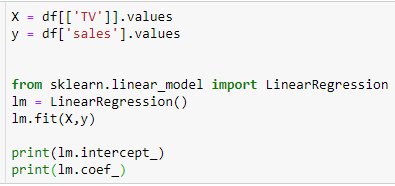
The closer the value of R2 is to 1 the better the model fits our data. If R2 comes below 0(which is a possibility) that means the model is so bad that it is performing even worse than the average best fit line.

**Adjusted R2 statistics**

As we increase the number of independent variables in our equation, the R2 increases as well. But that doesn’t mean that the new independent variables have any correlation with the output variable. In other words, even with the addition of new features in our model, it is not necessary that our model will yield better results but R2 value will increase. To rectify this problem, we use Adjusted R2 value which penalises excessive use of such features which do not correlate with the output data. Let’s understand this with an example:

We can see that R2 always increases with an increase in the number of independent variables. Thus, it doesn’t give a better picture and so we need Adjusted R2 value to keep this in check. Mathematically, it is calculated as:

In the equation above, when p = 0, we can see that adjusted R2 becomes equal to R2. Thus, adjusted R2 will always be less than or equal to R2, and it penalises the excess of independent variables which do not affect the dependent variable.



7.032593549127695

[0.04753664]

**Interpreting the model**

How do we interpret the coefficient for spends on TV ad (β1)?

* A "unit" increase in spends on a TV ad is **associated with** a 0.047537 "unit" increase in Sales.
* Or, an additional $1,000 on TV ads is **translated to** an increase in sales by 47.53 Dollars.

As an increase in TV ad expenditure is associated with a **decrease** in sales, β1 would be **negative**.

**Prediction using the model**

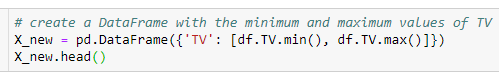
If the expense on TV ad is $50000, what will be the sales prediction for that market?

y = β0 + β1x

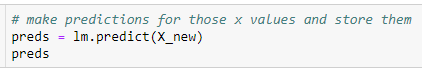
y = 7.032594+0.047537×50 = 9.409444

Thus, we would predict Sales of **9,409 widgets** in that market.

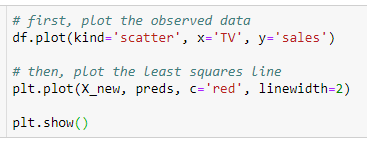
**Plotting the Least Squares Line**

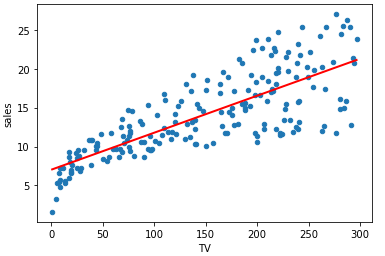






array([ 7.0658692 , 21.12245377])



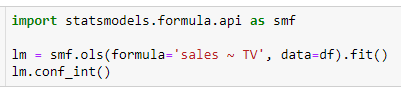


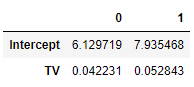
**Model Confidence**

**Question:** Is linear regression a low bias/high variance model or a high bias/low variance model?

**Answer:** It's a High bias/low variance model. Even after repeated sampling, the best fit line will stay roughly in the same position (low variance), but the average of the models created after repeated sampling won't do a great job in capturing the perfect relationship (high bias). Low variance is helpful when we don't have less training data!

If the model has calculated a 95% confidence for our model coefficients, it can be interpreted as follows: If the population from which this sample is drawn, is **sampled 100 times**, then approximately **95 (out of 100) of those confidence intervals** shall contain the "true" coefficients.





Keep in mind that we only have a **single sample of data**, and not the **entire population of data**. The "true" coefficient is either within this interval or it isn't, but there's no way actually to know. We estimate the coefficient with the data we do have, and we show uncertainty about that estimate by giving a range that the coefficient is **probably** within.

Note that using 95% confidence intervals is just a convention. You can create 90% confidence intervals (which will be more narrow), 99% confidence intervals (which will be wider), or whatever intervals you like.

**Hypothesis Testing and p-values**

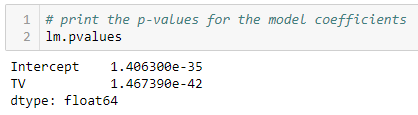
**Hypothesis testing** is Closely related to confidence intervals. We start with a **null hypothesis** and an **alternate hypothesis** (that is opposite to the null). Then, we check whether the data **rejects the null hypothesis** or **fails to reject the null hypothesis**.

("Failing to reject" the null hypothesis does not mean "accepting" the null hypothesis. The alternative hypothesis might indeed be true, but that we just don't have enough data to prove that.)

The conventional hypothesis test is as follows:

* **Null hypothesis:** No relationship exists between TV advertisements and Sales (and hence β1β1 equals zero).
* **Alternative hypothesis:** There exists a relationship between TV advertisements and Sales (and hence, β1β1 is not equal to zero).

How do we test this? We reject the null hypothesis (and thus believe the alternative hypothesis) if the 95% confidence interval **does not include zero**. The **p-value** represents the probability of the coefficient actually being zero.



If the 95% confidence interval **includes zero**, the p-value for that coefficient will be **greater than 0.05**. If the 95% confidence interval **does not include zero**, the p-value will be **less than 0.05**.

Thus, a p-value of less than 0.05 is a way to decide whether there is any relationship between the feature in consideration and the response or not. Using 0.05 as the cutoff is just a convention.

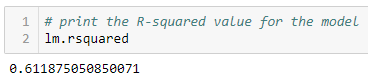
In this case, the p-value for TV ads is way less than 0.05, and so we **believe** that there is a relationship between TV advertisements and Sales.

Note that we generally ignore the p-value for the intercept.

### How Well Does the Model Fit the data?

One of the most generic way to evaluate the fit of a linear model is by computing the **R-squared** value. R-squared explains the **proportion of variance**, i.e., the proportion of variance in the observed data which the model explains, or the reduction in error over the **null model**. (A null model only predicts the mean of all the observed responses, and thus it only has an intercept and no slope.)

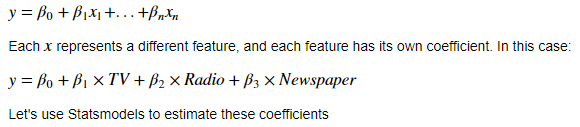
The value of R-squared lies between 0 and 1. A value closer to 1 is better as it means that more variance is explained by the model.

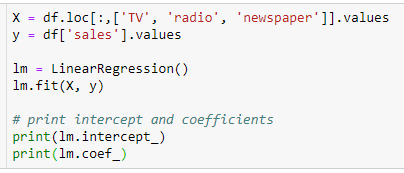


Is it a "good" R-squared value? Now, that’s hard to say. In reality, the domain to which the data belongs to plays a significant role in deciding the threshold for the R-squared value. Therefore, it's a tool for **comparing different models**.

**Multiple Linear Regression**

Till now, we have created the model based on only one feature. Now, we’ll include multiple features and create a model to see the relationship between those features and the label column. This is called **Multiple Linear Regression**.



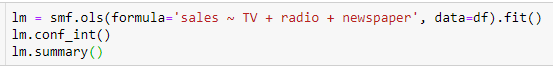


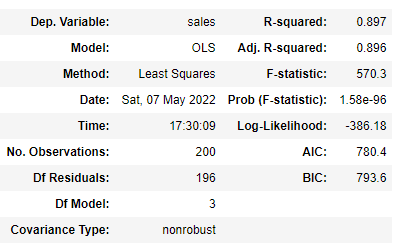
2.938889369459412

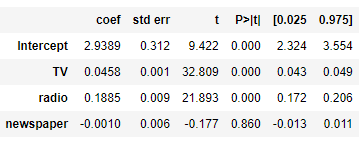
[ 0.04576465 0.18853002 -0.00103749]

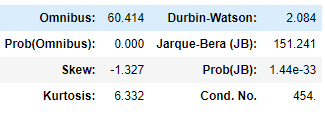
How do we interpret these coefficients? If we look at the coefficients, the coefficient for the newspaper spends is negative. It means that the money spent for newspaper advertisements is not contributing in a positive way to the sales.

A lot of the information we have been reviewing piece-by-piece is available in the model summary output:









Notes:  
[1] Standard Errors assume that the covariance matrix of the errors is correctly specified.

**What are the things to be learnt from this summary?**

* TV and Radio have positive **p-values**, whereas Newspaper has a negative one. Hence, we can reject the null hypothesis for TV and Radio that there is no relation between those features and Sales, but we fail to reject the null hypothesis for Newspaper that there is no relationship between newspaper spends and sales.
* The expenses on bot TV and Radio ads are**positively associated** with Sales, whereas the expense on newspaper ad is **slightly negatively associated** with the Sales.
* This model has a higher value of **R-squared** (0.897) than the previous model, which means that this model explains more variance and provides a better fit to the data than a model that only includes the TV.

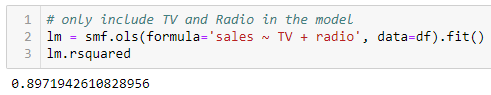
**Feature Selection**

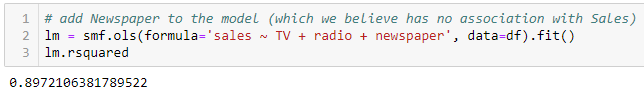
How do I decide **which features have to be included** in a linear model? Here's one idea:

* Try different models, and only keep predictors in the model if they have small p-values.
* Check if the R-squared value goes up when you add new predictors to the model.

What are the **drawbacks** in this approach? -If the underlying assumptions for creating a Linear model(the features being independent) are violated(which usually is the case),p-values and R-squared values are less reliable.

* Using a p-value cutoff of 0.05 means that adding 100 predictors to a model that are **pure noise**, still 5 of them (on average) will be counted as significant.
* R-squared is susceptible to **model overfitting**, and thus there is no guarantee that a model with a high R-squared value will generalise. Following is an example:





Selecting the model with the highest value of R-squared is not a correct approach as the value of R-squared shall always increase whenever a new feature is taken for consideration even if the feature is unrelated to the response.

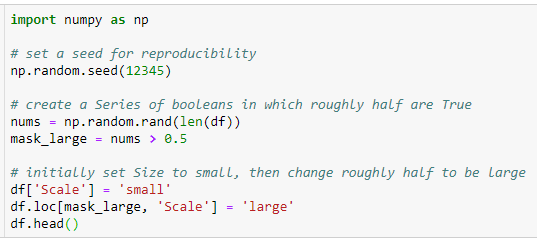
The alternative is to use **adjusted R-squared** which penalises the model complexity (to control overfitting), but this again generally [under-penalizes complexity](http://scott.fortmann-roe.com/docs/MeasuringError.html).

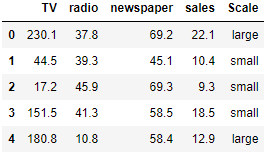
a better approach to feature selection is **Cross-validation.** It provides a more reliable way to choose which of the created models will best **generalise** as it better estimates of out-of-sample error. An advantage is that the cross-validation method can be applied to any machine learning model and the scikit-learn package provides extensive functionality for that.

**Handling Categorical Predictors with Two Categories**

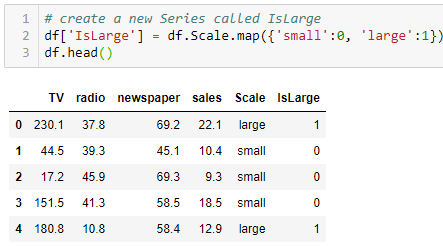
Till now, all the predictors have been numeric. What if one of the predictors is categorical?

We’ll create a new feature called **Scale**, and shall randomly assign observations as **small or large**:

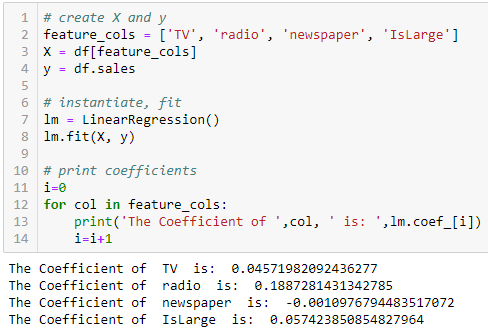




For the scikit-learn library, all data must be represented **numerically**. If the feature only has two categories, we can simply create a **dummy variable** that represents the categories as a combination of binary value:



Let's redo the multiple linear regression problem and include the **IsLarge** predictor:

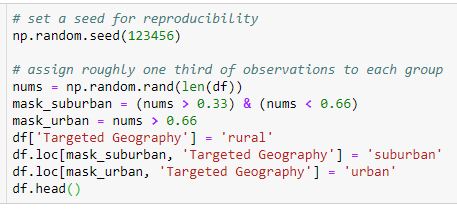


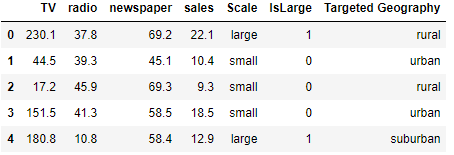
How do we interpret the coefficient for **IsLarge**? For a given TV/Radio/Newspaper ad expenditure if the average sales increases by 57.42 widgets, it’s considered as a large market.

What if the 0/1encoding is reversed? Still, the value of the coefficient shall be same, the only difference being the sign. It’ll be a negative number instead of positive.

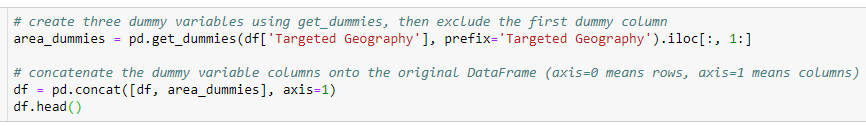
**Handling Categorical variables with More than Two Categories**

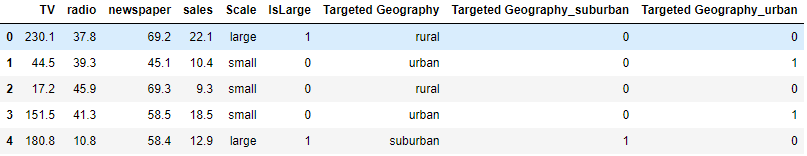
Let's create a new column called **Targeted Geography**, and randomly assign observations to be **rural, suburban, or urban**:





We need to represent the ‘Targeted Geography’ column numerically. But mapping urban=0, suburban=1 and rural=2 will mean that rural is two times suburban which is not the case. Hence, we’ll create another **dummy variable**:





**What does the encoding say?**

* rural is encoded as Targeted Geography\_suburban=0 and Targeted Geography\_urban=0
* suburban is encoded as Targeted Geography\_suburban=1 and Targeted Geography\_urban=0
* urban is encoded as Targeted Geography\_suburban=0 and Targeted Geography\_urban=1

Now the question is: **Why have we used two dummy columns instead of three?**

Because using only two dummy columns, we can capture the information of all the 3 columns. For example, if the value for Targeted Geography\_urban as well as Targeted Geography\_rural is 0, it automatically means that the data belongs to Targeted Geography\_suburban.

This is called handling the **dummy variable trap**. If there are N dummy variable columns, then the same information can be conveyed by N-1 columns. Let's include the two new dummy variables in the model:



['TV', 'radio', 'newspaper', 'IsLarge'] [ 0.04574401 0.1878667 -0.0010877 0.07739661 -0.10656299 0.26813802]

How do we interpret the coefficients?

* If all other columns are constant, the *suburban* geography is associated with an average **decrease** of 106.56 widgets in sales for $1000 spent.
* if $1000 is spent in an *urban* geography, it amounts to an average **increase** in Sales of 268.13 widgets

**A final note about dummy encoding:** If we have categories that can be ranked (i.e., worst, bad, good, better, best), we can potentially represent them numerically as (1, 2, 3, 4, 5) using a single dummy column

**Multi- Collinearity**

**Origin of the word:** The word multi-collinearity consists of two words : Multi, meaning multiple, and Collinear, meaning being linearly dependent on each other.

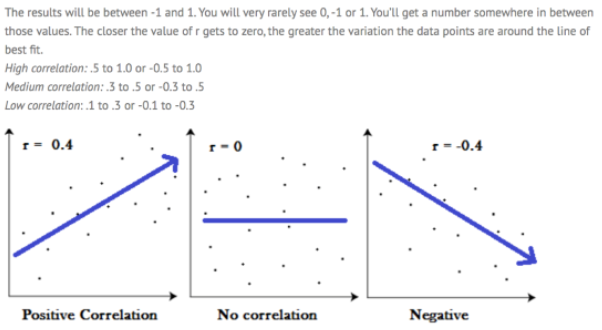
For e.g., Let’s consider this equation a + b=1=> b = 1−a

It means that ‘b’ can be represented in terms of ‘a’ i.e., if the value of ‘a’ changes, automatically the value of ‘b’ will also change. This equation denotes a simple linear relationship among two variables.

**Definition:** The purpose of executing a Linear Regression is to predict the value of a dependent variable based on certain independent variables.

So, when we perform a Linear Regression, we want our dataset to have variables which are independent i.e., we should not be able to define an independent variable with the help of another independent variable because now in our model we have two variables which can be defined based on a certain set of independent variables which defeats the entire purpose.

* Multi-collinearity is the statistical term to represent this type of a relation amongst the independent variable- when the independent variables are not so independent😊.
* We can define multi-collinearity as the situation where the independent variables (or the predictors) have strong correlation amongst themselves.



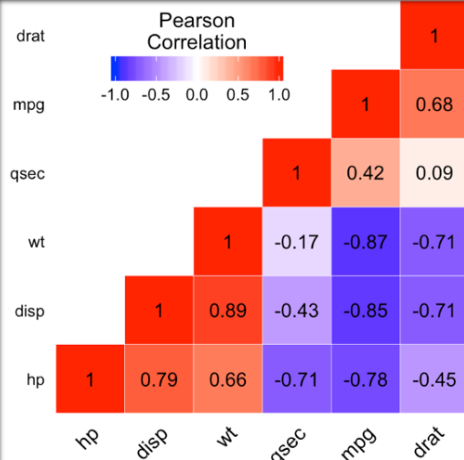
**Why Should We Care About Multi-Collinearity?**

* The coefficients in a Linear Regression model represent the extent of change in Y when a certain x (amongst X1,X2,X3…) is changed keeping others constant. But, if x1 and x2 are dependent, then this assumption itself is wrong that we are changing one variable keeping others constant as the dependent variable will also be changed. It means that our model itself becomes a bit flawed.
* We have a redundancy in our model as two variables (or more than two) are trying to convey the same information.
* As the extent of the collinearity increases, there is a chance that we might produce an overfitted model. An overfitted model works well with the test data but its accuracy fluctuates when exposed to other data sets.
* Can result in a Dummy Variable Trap.

**Detection**

**Correlation Matrices and Plots:** for correlation between all the X variables.

This plot shows the extent of correlation between the independent variable. Generally, a correlation greater than 0.9 or less than -0.9 is to be avoided.



* **Variance Inflation Factor:** Regression of one X variable against other X variables.

VIF=1/(1−Rsquared)

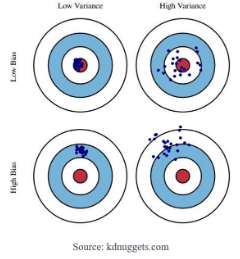
The VIF factor, if greater than 10 shows extreme correlation between the variables and then we need to take care of the correlation.

**Remedies for Multicollinearity**

* **Do Nothing:** If the Correlation is not that extreme, we can ignore it. If the correlated variables are not used in solving our business question, they can be ignored.
* **Remove One Variable**: Like in dummy variable trap
* **Combine the correlated variables:** Like creating a seniority score based on Age and Years of experience
* Principal Component Analysis

**Bias Variance trade-off**

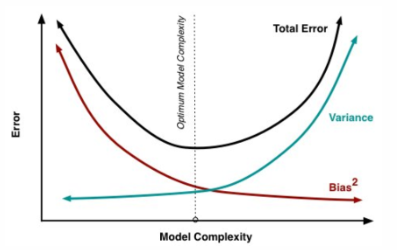
Variance is a measure of the degree of the spread of the predicted results. Bias, on the other hand, is a measure of the polarity of the results i.e., the results are favoured towards a specific outcome. The problem with Bias and Variance can be depicted as follows:

As depicted in the figure, our goal is to minimize the Variance as well as Bias to get to accurate results. If there are lesser number of predictors in the data set that too having big coefficients, it is a possibility that the model obtained is favored towards those predictors i.e., a small change in those predictors lead to a big change in the predicted result. In other words, that predictor is driving our model, or our model is biased for that predictor. An easy way to overcome this is to introduce more no. of predictors which contributes towards the end result of the prediction. This increases the complexity of the model or in other words, the variance in the model increases. Mathematically,



Irreducible error is the error due to noise in the data.

Graphically it looks like:



It can be seen that as the bias decreases, the variance increases. So, we need to find a balance so that we get a model which has low variance as well as low bias. This is called **bias-variance tradeoff.**

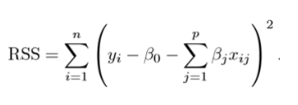
Bias can be minimized by training with more data and variance can be reduced by using regularization methods which we are going to discuss.

**Regularization**

When we use regression models to train some data, there is a good chance that the model will overfit the given training data set. Regularization helps sort this overfitting problem by restricting the degrees of freedom of a given equation i.e. simply reducing the number of degrees of a polynomial function by reducing their corresponding weights.  
In a linear equation, we do not want huge weights/coefficients as a small change in weight can make a large difference for the dependent variable (Y). So, regularization constraints the weights of such features to avoid overfitting. Simple linear regression is given as:



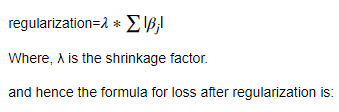
Using the OLS method, we try to minimize the cost function given as:

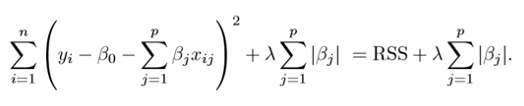


To regularize the model, a Shrinkage penalty is added to the cost function. Let’s see different types of regularizations in regression:

**LASSO(Least Absolute Shrinkage and Selection Operator) Regression (L1 Form)**

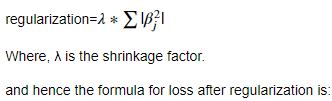
LASSO regression penalizes the model based on the sum of magnitude of the coefficients. The regularization term is given by

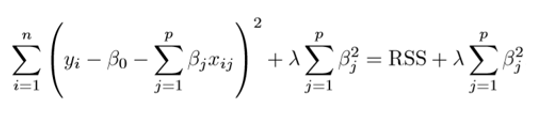




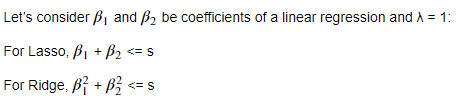
**Ridge Regression (L2 Form)**

Ridge regression penalizes the model based on the sum of squares of magnitude of the coefficients. The regularization term is given by

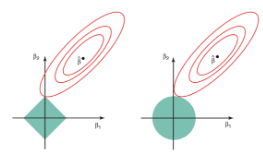




This value of lambda can be anything and should be calculated by cross validation as to what suits the model.



Where s is the maximum value the equations can achieve . If we plot both the above equations, we get the following graph:



The red ellipse represents the cost function of the model, whereas the square (left side) represents the Lasso regression and the circle (right side) represents the Ridge regression.

**Difference between Ridge and Lasso**

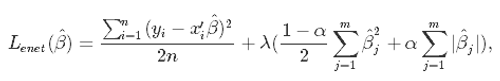
Ridge regression shrinks the coefficients for those predictors which contribute very less in the model but have huge weights, very close to zero. But it never makes them exactly zero. Thus, the final model will still contain all those predictors, though with less weights. This doesn’t help in interpreting the model very well. This is where Lasso regression differs with Ridge regression. In Lasso, the L1 penalty does reduce some coefficients exactly to zero when we use a sufficiently large tuning parameter λ. So, in addition to regularizing, lasso also performs feature selection.

**Why use Regularization?**

Regularization helps to reduce the variance of the model, without a substantial increase in the bias. If there is variance in the model that means that the model won’t fit well for dataset different that training data. The tuning parameter λ controls this bias and variance tradeoff. When the value of λ is increased up to a certain limit, it reduces the variance without losing any important properties in the data. But after a certain limit, the model will start losing some important properties which will increase the bias in the data. Thus, the selection of good value of λ is the key. The value of λ is selected using cross-validation methods. A set of λ is selected and cross-validation error is calculated for each value of λ and that value of λ is selected for which the cross-validation error is minimum.

**Elastic Net**

According to the Hands-on Machine Learning book, elastic Net is a middle ground between Ridge Regression and Lasso Regression. The regularization term is a simple mix of both Ridge and Lasso’s regularization terms, and you can control the mix ratio α.

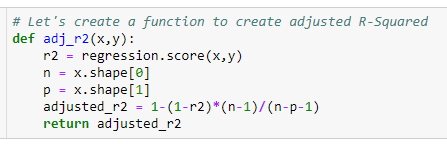


where α is the mixing parameter between ridge (α = 0) and lasso (α = 1).

**When should you use plain Linear Regression (i.e., without any regularization), Ridge, Lasso, or Elastic Net?**

According to the Hands-on Machine Learning book, it is almost always preferable to have at least a little bit of regularization, so generally you should avoid plain Linear Regression. Ridge is a good default, but if you suspect that only a few features are actually useful, you should prefer Lasso or Elastic Net since they tend to reduce the useless features’ weights down to zero as we have discussed. In general, Elastic Net is preferred over Lasso since Lasso may behave erratically when the number of features is greater than the number of training instances or when several features are strongly correlated.

No let's see the Python Implementation of these concepts:

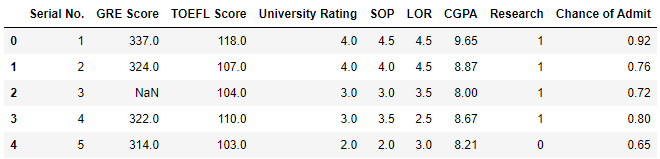


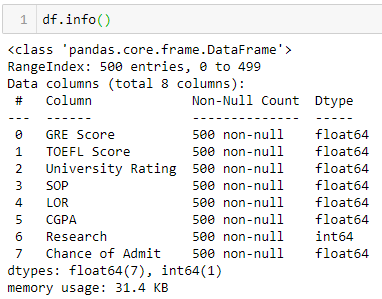
**Problem statement:** the goal here is to find the chance od admission of a candidate based on his/her

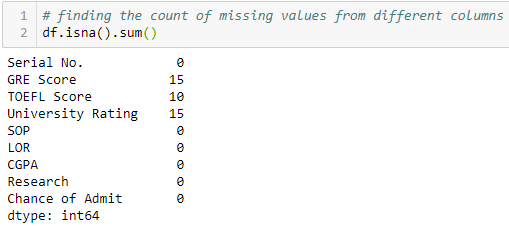
GRE Score(out of 340), TOEFL Score(out of 120), rating of the University(out of 5)

In which he/she trying to get the admission, strength of the Letter of Recommendation(out of 5), CGPA(out of 10) and the research experience(0 or 1)

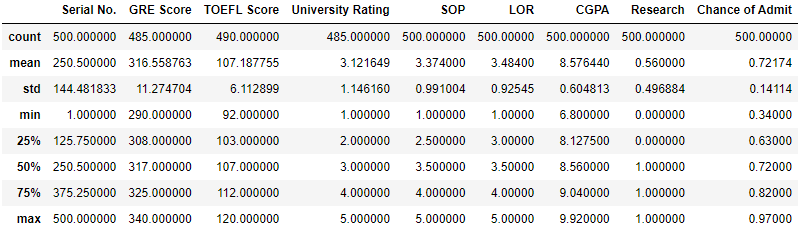


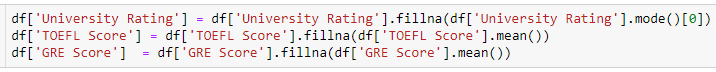




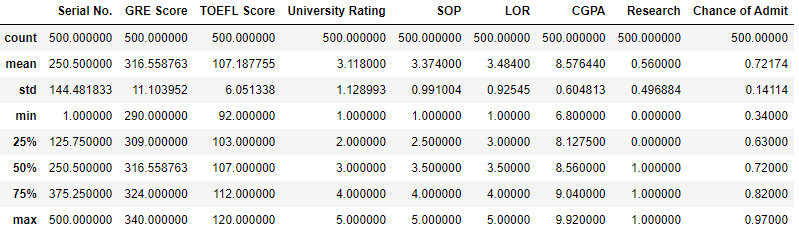






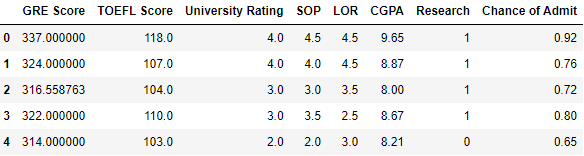




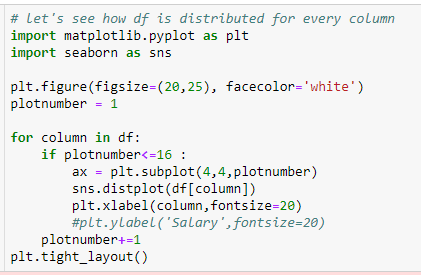


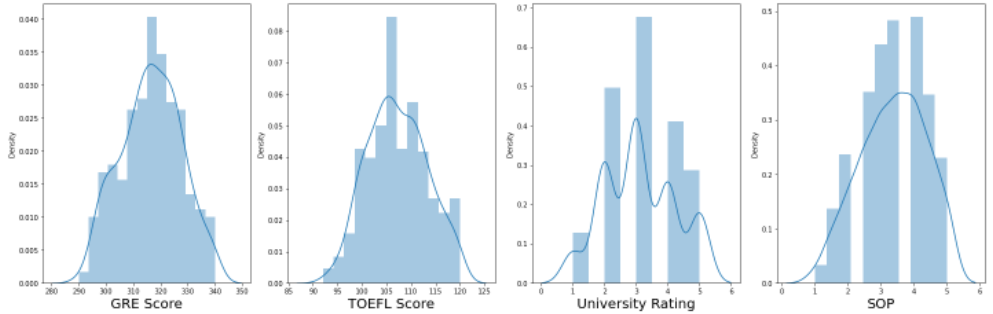
Now the data looks good and there are no missing values. Also, the first cloumn is just serial numbers, so we don' need that column. Let's drop it from data and make it more clean.

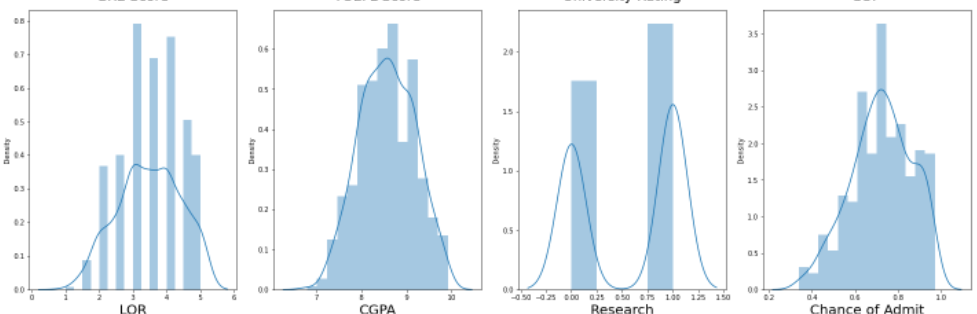




Let's visualize the data and analyze the relationship between independent and dependent variables:



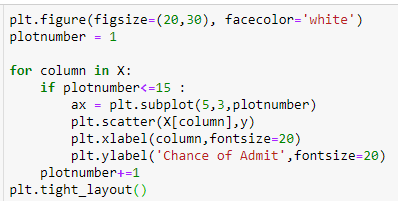


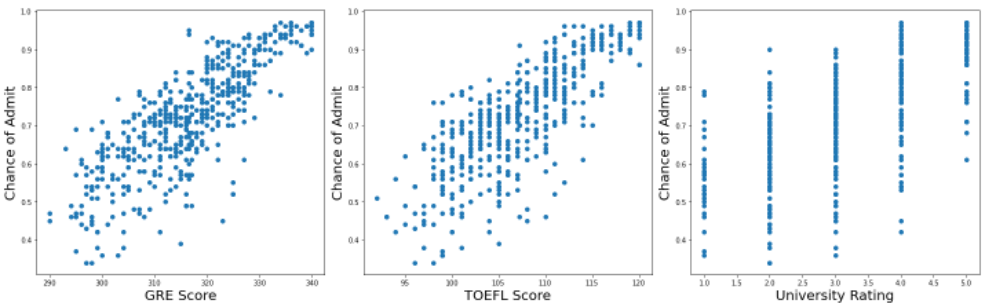


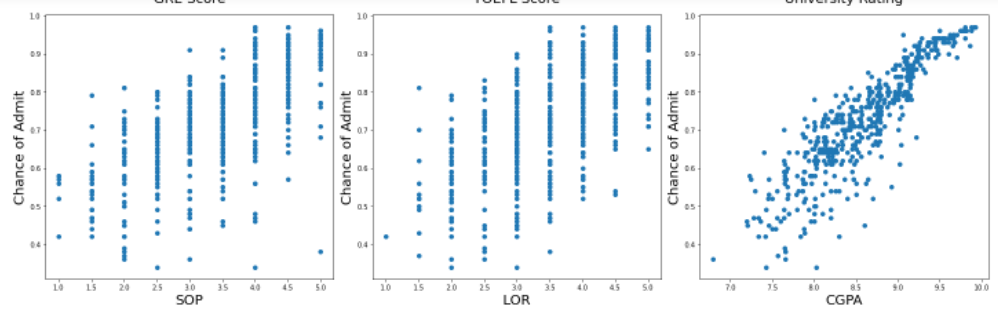
The data distribution looks decent enough and there doesn't seem to be any skewness. Great let's go ahead!

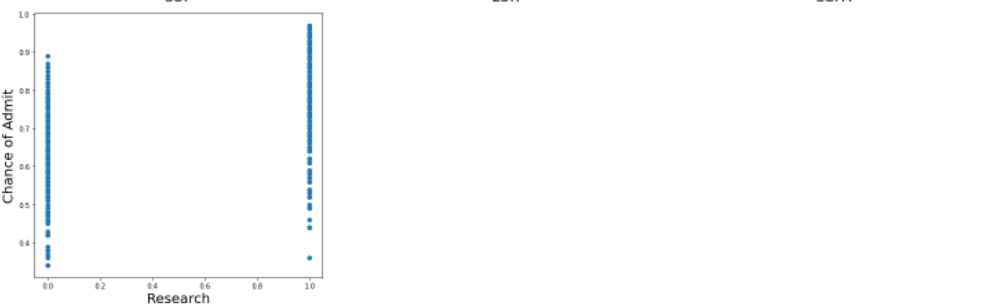
Let's observe the relationship between independent variables and dependent variable.







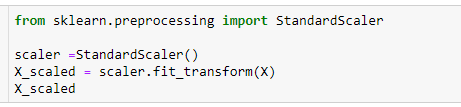


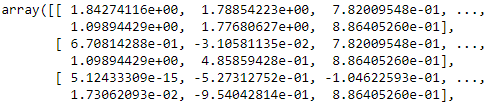


Great, the relationship between the dependent and independent variables look fairly linear.

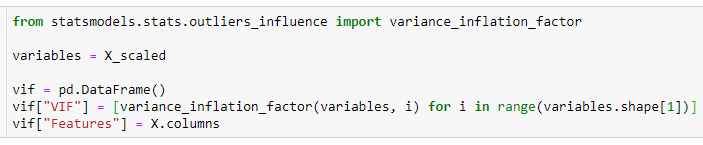
Thus, our linearity assumption is satisfied.

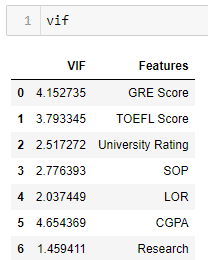
Let's move ahead and check for multicollinearity.





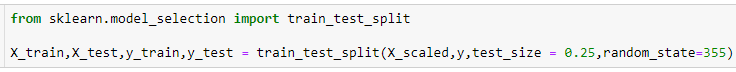
* we create a new data frame which will include all the VIFs
* Note that each variable has its own variance inflation factor as this measure is variable specific (not model specific)
* we do not include categorical values for mulitcollinearity as they do not provide much information as numerical ones do
* here we make use of the variance\_inflation\_factor, which will basically output the respective VIFs
* Finally, I like to include names so it is easier to explore the result.

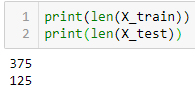


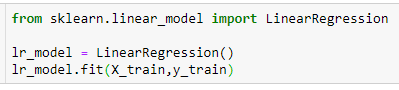


Here, we have the correlation values for all the features. As a thumb rule, a VIF value greater than 5 means a very severe multicollinearity. We don't any VIF greater than 5 , so we are good to go.

Great. Let's go ahead and use linear regression and see how good it fits our data. But first. let's split our data in train and test.

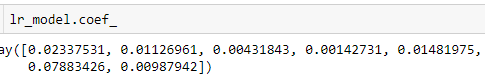




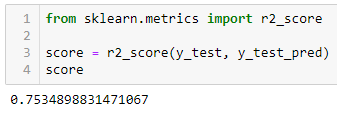


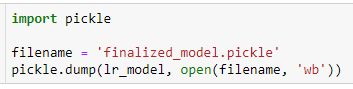
LinearRegression()

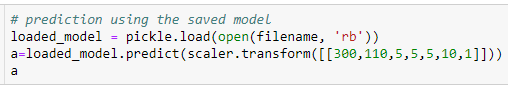








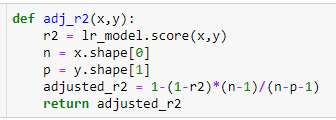




array([0.92190162])



0.8415250484247909





Our r2 score is 84.15% and adj r2 is 83.85% for our training etc., so looks like we are not being penalized by use of any feature.

Let's check how well model fits the test data.

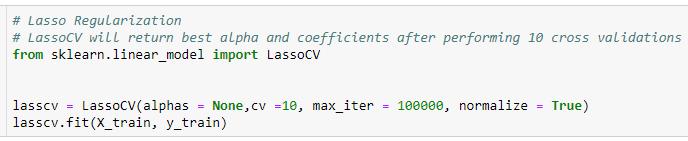
Now let's check if our model is overfitting our data using regularization.



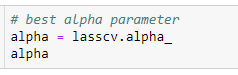
0.7534898831471067

So it looks like our model r2 score is less on the test data.

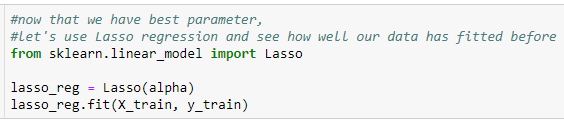
Let's see if our model is overfitting our training data.



LassoCV(cv=10, max\_iter=100000, normalize=True)



3.0341655445178153e-05

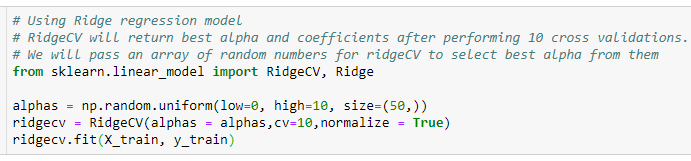


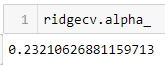
Lasso(alpha=3.0341655445178153e-05)

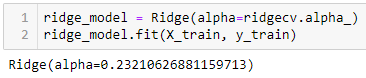


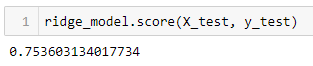
0.7534654960492284

our r2\_score for test data (75.34%) comes same as before using regularization. So, it is fair to say our OLS model did not overfit the data.

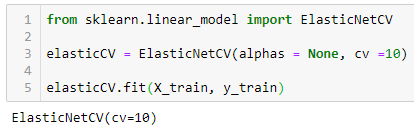




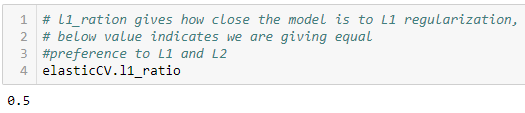


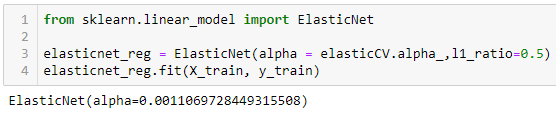


we got the same r2 square using Ridge regression as well. So, it's safe to say there is no overfitting.







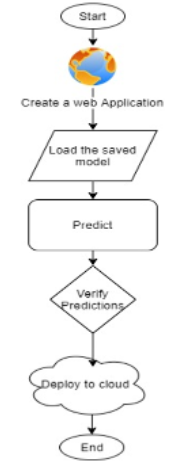




So, we can see by using different type of regularization, we still are getting the same r2 score. That means our OLS model has been well trained over the training data and there is no overfitting.

**Cloud Deployment (Google Cloud Platform)**

Once the training is completed, we need to expose the trained model as an API for the user to consume it. For prediction, the saved model is loaded first and then the predictions are made using it. If the web app works fine, the same app is deployed to the cloud platform. The application flow for cloud deployment looks like:



Pre-requisites:

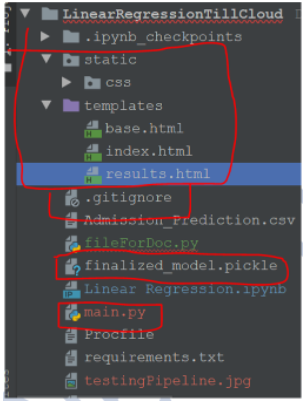
* Basic knowledge of flask framework.
* Any Python IDE installed(we are using PyCharm).
* A Google Cloud Platform account.
* Basic understanding of HTML.

Flask App

As we’ll expose the created model as a web API to be consumed by the client/client APIs, we’d do it using the flask framework. The flow of our flask app will be:



Create the project structure, as shown below:



Deployment to G-cloud:

* Go to <https://cloud.google.com/> and create an account if already haven’t created one. Then go to the console of your account.
* Go to **IAM and admin**(highlighted) and click **manage resources**
* Click CREATE PROJECT to create a new project for deployment.
* Once the project gets created, select **App Engine** and select **Dashboard**.
* Go to <https://dl.google.com/dl/cloudsdk/channels/rapid/GoogleCloudSDKInstaller.exe> to download the google cloud SDK to your machine.
* Click Start Tutorial on the screen and select Python app and click start.
* Check whether the correct project name is displayed and then click next.
* Create a file ‘app.yaml’ and put ‘runtime: python37’ in that file.
* Create a ‘requirements.txt’ file by opening the command prompt/anaconda prompt, navigate to the project folder and enter the command ‘pip freeze > requirements.txt’. It is recommended to use separate environments for different projects.
* Your python application file should be called ‘main.py’. It is a GCP specific requirement.
* Open command prompt window, navigate to the project folder and enter the command gcloud init to initialise the gcloud context.
* It asks you to select from the list of available projects
* Once the project name is selected, enter the command gcloud app deploy app.yaml --project
* After executing the above command, GCP will ask you to enter the region for your application. Choose the appropriate one.
* GCP will ask for the services to be deployed. Enter ‘y’ to deploy the services.
* And then it will give you the link for your app,and the deployed app looks like:

**Polynomial Regression**

For understanding Polynomial Regression, let's first understand a polynomial. Merriam-webster defines a polynomial as: "A mathematical expression of one or more algebraic terms each of which consists of a constant multiplied by one or more variables raised to a non-negative integral power (such as a + bx + cx^2)". Simply said, poly means many. So, a polynomial is an aggregation of many monomials(or Variables). A simple polynomial equation can be written as:



So, Polynomial Regression can be defined as a mechanism to predict a dependent variable based on the polynomial relationship with the independent variable.

In the equation,



the maximum power of 'x' is called the degree of the polynomial equation. For example, if the degree is 1, the equation becomes



which is a simple linear equation. if the degree is 2, the equation becomes



which is a quadratic equation and so on.