### Medical Cost Prediction

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Github: https://github.com/sampurna-project

```
#importing the libraries
import numpy as np
import pandas as pd
import matplotlib.pyplot as plt
import seaborn as sns

Insurance = pd.read_csv('/content/insurance.csv')
Insurance.head()
```

	age	sex	bmi	children	smoker	region	charges
0	19	female	27.900	0	yes	southwest	16884.92400
1	18	male	33.770	1	no	southeast	1725.55230
2	28	male	33.000	3	no	southeast	4449.46200
3	33	male	22.705	0	no	northwest	21984.47061
4	32	male	28.880	0	no	northwest	3866.85520

### **Data Preprocessing**

```
Insurance.shape
```

(1338, 7)

#checking for missing values
Insurance.info()

dtypes: float64(2), int64(2), object(3)

memory usage: 73.3+ KB

#checking discriptive statistics

Insurance.describe()

	age	bmi	children	charges
count	1338.000000	1338.000000	1338.000000	1338.000000
mean	39.207025	30.663397	1.094918	13270.422265
std	14.049960	6.098187	1.205493	12110.011237
min	18.000000	15.960000	0.000000	1121.873900
25%	27.000000	26.296250	0.000000	4740.287150
50%	39.000000	30.400000	1.000000	9382.033000
75%	51.000000	34.693750	2.000000	16639.912515
max	64.000000	53.130000	5.000000	63770.428010

#value counts for categorical variables
print(Insurance.sex.value\_counts(),'\n',Insurance.region.value\_counts())

male 676 female 662 Name: sex, dtype: int64 no 1064 274 yes Name: smoker, dtype: int64 southeast southwest 325 northwest 325 northeast 324 Name: region, dtype: int64

Replacing the categorical variables with numerical values.

sex: 1 - male, 0 - femalesmoker: 1 - yes, 0 - no

• region: 0 - northeast, 1 - northwest, 2 - southeast, 3 - southwest

```
#changing categorical variables to numerical
Insurance['sex'] = Insurance['sex'].map({'male':0})
Insurance['smoker'] = Insurance['smoker'].map({'yes':1, 'no':0})
Insurance['region'] = Insurance['region'].map({'southwest':0, 'southeast':1, 'northwest':2, 'northeast':3})
```

Insurance.head(10)

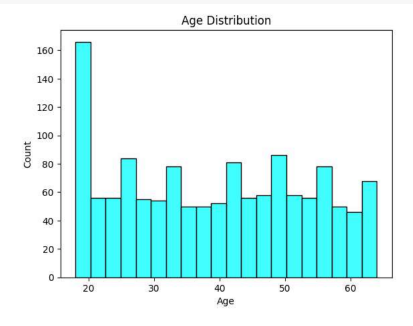
	age	sex	bmi	children	smoker	region	charges
0	19	0	27.900	0	1	0	16884.92400
1	18	1	33.770	1	0	1	1725.55230
2	28	1	33.000	3	0	1	4449.46200
3	33	1	22.705	0	0	2	21984.47061
4	32	1	28.880	0	0	2	3866.85520
5	31	0	25.740	0	0	1	3756.62160
6	46	0	33.440	1	0	1	8240.58960

## ▼ Exploratory Data Analysis

**9** 60 0 25.840 0 0 2 28923.13692

Visualization of the data is a good way to understand the data. In this section, I will plot the distribution of each variable to get an overview about their counts and distributions.

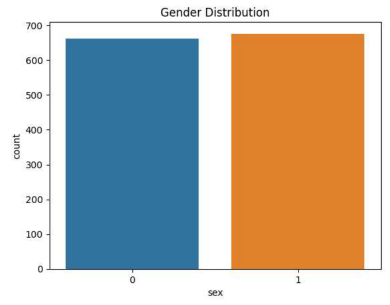
```
#age distribution
sns.histplot(Insurance.age,bins=20, kde=False,color='aqua')
plt.title('Age Distribution')
plt.xlabel('Age')
plt.ylabel('Count')
plt.show()
```



#gender plot
sns.countplot(x = 'sex', data = Insurance)

plt.title('Gender Distribution')

Text(0.5, 1.0, 'Gender Distribution')



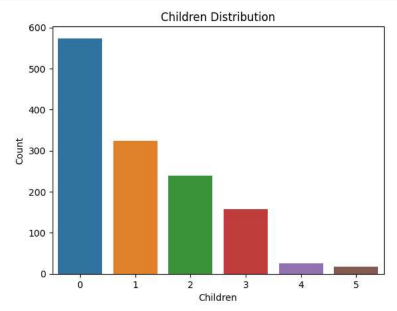
It is clear that number of males and females are almost equal in the dataset.

```
#bmi distribution
sns.histplot(Insurance.bmi,bins=20, kde=True,color='blue')
plt.title('BMI Distribution')
plt.xlabel('BMI')
plt.ylabel('Count')
plt.show()
```

# BMI Distribution

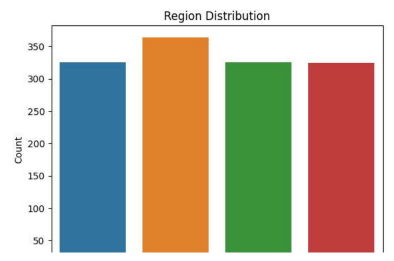
The majority of the patients have BMI between 25 and 40 which is considered as overweight and could be a major factor in increasing the medical cost.

```
#child count distribution
sns.countplot(x = 'children', data = Insurance)
plt.title('Children Distribution')
plt.xlabel('Children')
plt.ylabel('Count')
plt.show()
```



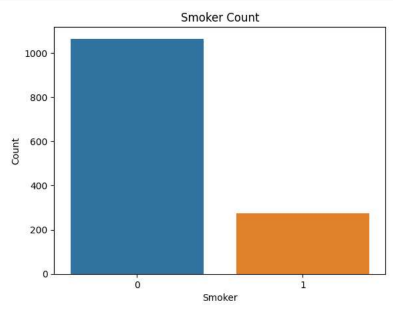
The graph clearly shows that most of the patients have no children and very few patients have more than 3 children.

```
#regionwise plot
sns.countplot(x = 'region', data = Insurance)
plt.title('Region Distribution')
plt.xlabel('Region')
plt.ylabel('Count')
plt.show()
```



The count of patient from northwest is slighltly higher than the other regions, but the number of patients from other regions are almost equal.

```
#count of smokers
sns.countplot(x = 'smoker', data = Insurance)
plt.title('Smoker Count')
plt.xlabel('Smoker')
plt.ylabel('Count')
plt.ylabel('Count')
```

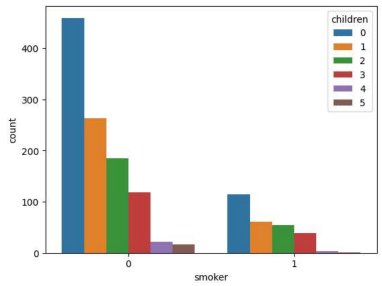


→ Smokers are very few in the dataset. Nearly 80% of the patients are non-smokers.

# Smoker count with respect to the children count.

```
sns.countplot(x = Insurance.smoker, hue = Insurance.children)
```



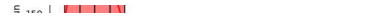


```
#charges distribution
sns.histplot(Insurance.charges,bins=20, kde=True,color='red')
plt.title('Charges Distribution')
plt.xlabel('Medical Expense')
plt.ylabel('Count')
plt.show()
```

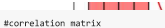
# Charges Distribution

Most of the medical expenses are below 20000, with negligible number of patients having medical expenses above 50000.

From all the above plots, we have a clear understanding about the count of patients under each category of the variables. Now I will look into the coorelation between the variables.



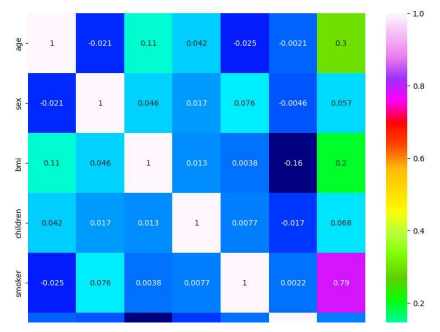
### Correlation



Insurance.corr()

	age	sex	bmi	children	smoker	region	charges
age	1.000000	-0.020856	0.109272	0.042469	-0.025019	-0.002127	0.299008
sex	-0.020856	1.000000	0.046371	0.017163	0.076185	-0.004588	0.057292
bmi	0.109272	0.046371	1.000000	0.012759	0.003750	-0.157566	0.198341
children	0.042469	0.017163	0.012759	1.000000	0.007673	-0.016569	0.067998
smoker	-0.025019	0.076185	0.003750	0.007673	1.000000	0.002181	0.787251
region	-0.002127	-0.004588	-0.157566	-0.016569	0.002181	1.000000	0.006208
charges	0.299008	0.057292	0.198341	0.067998	0.787251	0.006208	1.000000

```
#plotting the coorelation heatmap
plt.figure(figsize=(10,10))
sns.heatmap(Insurance.corr(),annot=True,cmap='gist_ncar')
plt.show()
```



The variable smoker shows a significant coorelation with the medical expenses. Now I will explore more into patients' smoking habits and their relationa with other factors.

```
Plotting the smoker count with patient's gender

sns.catplot(x="smoker", kind="count",hue = 'sex', data=Insurance)
plt.title('Smoker Count with gender')
plt.xlabel('Smoker')
plt.ylabel('Count')
plt.show()
```

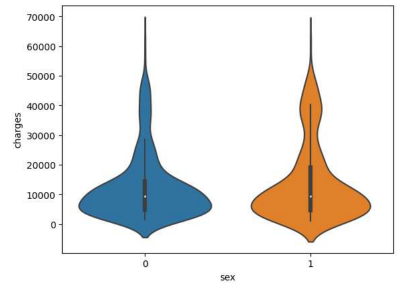
### Smoker Count with gender



We can notice more male smokers than female smokers. So, I will assume that medical treatment expense for males would be more than females, given the impact of smoking on the medical expenses.



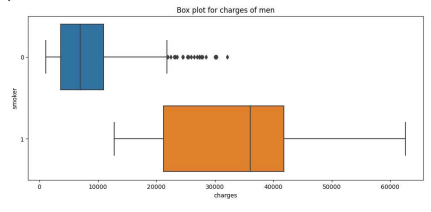
<Axes: xlabel='sex', ylabel='charges'>



```
plt.figure(figsize=(12,5))
plt.title("Box plot for charges of women")
sns.boxplot(y="smoker", x="charges", data = Insurance[(Insurance.sex == 0)] , orient="h")
```

```
plt.figure(figsize=(12,5))
plt.title("Box plot for charges of men")
sns.boxplot(y="smoker", x="charges", data = Insurance[(Insurance.sex == 1)] , orient="h")
```

<Axes: title={'center': 'Box plot for charges of men'}, xlabel='charges',
ylabel='smoker'>

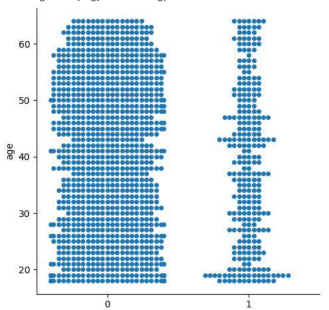


The assumption is true, that the medical expense of males is greater than that of females. In addituion to that medical expense of smokers is greater than that of non-smokers.

Smokers and age distribution

```
#smokers and age distribution
sns.catplot(x="smoker", y="age", kind="swarm", data=Insurance)
```

```
<seaborn.axisgrid.FacetGrid at
0x7f995f704a00>/usr/local/lib/python3.10/dist-packages/seaborn/categorical.py:3544
warnings.warn(msg, UserWarning)
```



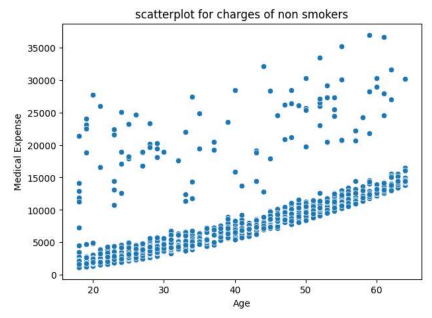
From the graph, we can see that there significant number of smokers of age 19. Now I will study the medical expense of smokers of age 19.

```
#smokers of age 19
plt.figure(figsize=(12,5))
plt.title("Box plot for charges of smokers of age 19")
sns.boxplot(y="smoker", x="charges", data = Insurance[(Insurance.age == 19)] , orient="h")
plt.xlabel('Medical Expense')
plt.ylabel('Smoker')
plt.show()
```

Surprisingly the medical expense of smokers of age 19 is very high in comparison to non smokers. In non smokers we can see some outliners, which may be due to illness or accidents.

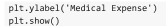
It is clear that the medical expense of smokers is higher than that of non-smokers. Now I will plot the charges distribution with repect to patients age of smokers and non-smokers.

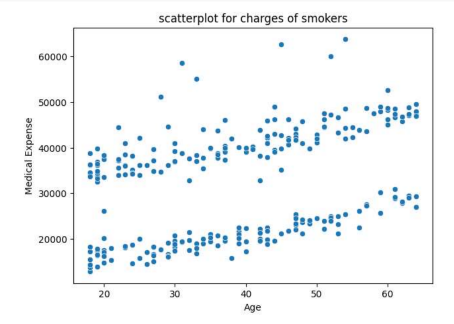
```
#non smokers charge distribution
plt.figure(figsize=(7,5))
plt.title("scatterplot for charges of non smokers")
sns.scatterplot(x="age", y="charges", data = Insurance[(Insurance.smoker == 0)])
plt.xlabel('Age')
plt.ylabel('Medical Expense')
plt.show()
```



Majority of the points shows that medical expense increases with age which may be due to the fact that older people are more prone to illness. But there are some outliners which shows that there are other illness or accidents which may increase the medical expense.

```
#smokers charge distribution
plt.figure(figsize=(7,5))
plt.title("scatterplot for charges of smokers")
sns.scatterplot(x="age", y="charges", data = Insurance[(Insurance.smoker == 1)])
plt.xlabel('Age')
```





Here we see pecularity in the graph. In the graph there are two segments, one with high medical expense which may be due to smoking related illness and the other with low medical expense which may be due age related illness.

Now, in order to get a more clear picture, I will combine these two graphs.

```
#age charges distribution

sns.lmplot(x="age", y="charges", data = Insurance, hue = 'smoker')
plt.xlabel('Age')
plt.ylabel('Medical Expense')
plt.show()
```



Now, we clearly understand the variation in charges with respect to age and smoking habits. The medical expense of smokers is higher than that of non-smokers. In non-smokers, the cost of treatment increases with age which is obvious. But in smokers, the cost of treatment is high even for younger patients, which means the smoking patients are spending upon their smoking related illness as well as age related illness.

```
#bmi charges distribution for obese people
plt.figure(figsize=(7,5))
sns.distplot(Insurance[(Insurance.bmi >= 30)]['charges'])
plt.title('Charges Distribution for Obese People')
plt.xlabel('Medical Expense')
plt.show()
```

```
<ipython-input-34-3ee980cb50b5>:3: UserWarning:
```

Charges distribution for patients with BMI less than 30 i.e. healthy patients

```
plt.figure(figsize=(7,5))
sns.distplot(Insurance[(Insurance.bmi < 30)]['charges'])
plt.title('Charges Distribution for Non Obese People')
plt.xlabel('Medical Expense')
plt.show()</pre>
```

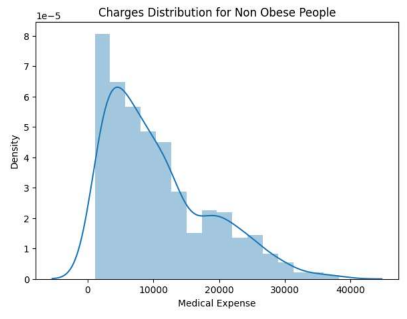
<ipython-input-35-5afd3889ee34>:2: UserWarning:

`distplot` is a deprecated function and will be removed in seaborn v0.14.0.

Please adapt your code to use either `displot` (a figure-level function with similar flexibility) or `histplot` (an axes-level function for histograms).

For a guide to updating your code to use the new functions, please see https://gist.github.com/mwaskom/de44147ed2974457ad6372750bbe5751

sns.distplot(Insurance[(Insurance.bmi < 30)]['charges'])</pre>



Therefore, patients with BMI less than 30 are spending less on medical treatment than those with BMI greater than 30.

Through the EDA, we have a clear understanding about the data and the coorelation between the variables. Now, I will build a model to predict the medical expense of patients.

Train Test Split

```
from sklearn.model_selection import train_test_split
x_train, x_test, y_train, y_test = train_test_split(Insurance.drop('charges',axis=1), Insurance['charges'], test_size=0.2, random_state=0)
Model Building
Linear Regression
#Linear Regression
from sklearn.linear_model import LinearRegression
lr = LinearRegression()
lr
     ▼ LinearRegression
     LinearRegression()
#model training
lr.fit(x_train,y_train)
#model accuracy
lr.score(x_train,y_train)
     0.7368306228430945
#model prediction
y_pred = lr.predict(x_test)
Polynomial Regression
from sklearn.preprocessing import PolynomialFeatures
poly_reg = PolynomialFeatures(degree=2)
poly_reg
     ▼ PolynomialFeatures
     PolynomialFeatures()
#transforming the features to higher degree
x_train_poly = poly_reg.fit_transform(x_train)
#splitting the data
x_train, x_test, y_train, y_test = train_test_split(x_train_poly, y_train, test_size=0.2, random_state=0)
plr = LinearRegression()
#model training
plr.fit(x_train,y_train)
#model accuracy
plr.score(x_train,y_train)
```

0.8372892271391211

```
#model prediction
y_pred = plr.predict(x_test)
Decision Tree Regressor
#decision tree regressor
from sklearn.tree import DecisionTreeRegressor
dtree = DecisionTreeRegressor()
dtree
      ▼ DecisionTreeRegressor
     DecisionTreeRegressor()
#model training
dtree.fit(x_train,y_train)
#model accuracy
dtree.score(x_train,y_train)
     0.9993688476658964
#model prediction
dtree_pred = dtree.predict(x_test)
Random Forest Regressor
#random forest regressor
from sklearn.ensemble import RandomForestRegressor
rf = RandomForestRegressor(n_estimators=100)
     * RandomForestRegressor
     RandomForestRegressor()
#model training
rf.fit(x_train,y_train)
#model accuracy
rf.score(x_train,y_train)
     0.9760236161665283
#model prediction
rf_pred = rf.predict(x_test)
Model Evaluation
from sklearn.metrics import mean_squared_error,mean_absolute_error,r2_score
```

Linear Regression

```
#distribution of actual and predicted values
plt.figure(figsize=(7,5))
ax1 = sns.distplot(y_test,hist=False,color='r',label='Actual Value')
sns.distplot(y_pred,hist=False,color='b',label='Predicted Value',ax=ax1)
plt.title('Actual vs Predicted Values for Linear Regression')
plt.xlabel('Medical Expense')
plt.show()
```

<ipython-input-52-2d0e63236188>:3: UserWarning:

`distplot` is a deprecated function and will be removed in seaborn v0.14.0.

Please adapt your code to use either `displot` (a figure-level function with similar flexibility) or `kdeplot` (an axes-level function for kernel density plots

For a guide to updating your code to use the new functions, please see <a href="https://gist.github.com/mwaskom/de44147ed2974457ad6372750bbe5751">https://gist.github.com/mwaskom/de44147ed2974457ad6372750bbe5751</a>

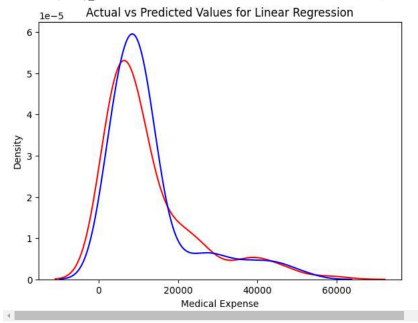
ax1 = sns.distplot(y\_test,hist=False,color='r',label='Actual Value')
<ipython-input-52-2d0e63236188>:4: UserWarning:

`distplot` is a deprecated function and will be removed in seaborn v0.14.0.

Please adapt your code to use either `displot` (a figure-level function with similar flexibility) or `kdeplot` (an axes-level function for kernel density plots

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sns.distplot(y\_pred,hist=False,color='b',label='Predicted Value',ax=ax1)



print('MAE:', mean\_absolute\_error(y\_test, y\_pred))
print('MSE:', mean\_squared\_error(y\_test, y\_pred))

```
print('RMSE:', np.sqrt(mean_squared_error(y_test, y_pred)))
print('R2 Score:', r2_score(y_test, y_pred))
```

MAE: 2988.626627897196 MSE: 24512834.56541676 RMSE: 4951.043785447344 R2 Score: 0.8221477010678055

### Polynomial Regression

```
#acutal vs predicted values for polynomial regression
plt.figure(figsize=(7,5))
ax1 = sns.distplot(y_test,hist=False,color='r',label='Actual Value')
sns.distplot(y_pred,hist=False,color='b',label='Predicted Value',ax=ax1)
plt.title('Actual vs Predicted Values for Polynomial Regression')
plt.xlabel('Medical Expense')
plt.show()
```

```
cipython-input-54-7a574536b1bb>:3: UserWarning:
    'distplot' is a deprecated function and will be removed in seaborn v0.14.0.

Please adapt your code to use either 'displot' (a figure-level function with similar flexibility) or 'kdeplot' (an axes-level function for kernel density plots

print('MAE:', mean_absolute_error(y_test, y_pred))
print('MSE:', mean_squared_error(y_test, y_pred))
print('RMSE:', np.sqrt(mean_squared_error(y_test, y_pred)))
print('RZ Score:', r2_score(y_test, y_pred))

MAE: 2988.626627897196
MSE: 24512834.56541676
RMSE: 4951.043785447344
RZ Score: 0.8221477010678055
```

#### Decision Tree Regressor

```
#distribution plot of actual and predicted values
plt.figure(figsize=(7,5))
ax = sns.distplot(y_test, hist=False, color="r", label="Actual Value")
sns.distplot(dtree_pred, hist=False, color="b", label="Fitted Values" , ax=ax)
plt.title('Actual vs Fitted Values for Decision Tree Regression')
plt.xlabel('Medical Expense')
plt.ylabel('Distribution')
plt.show()
```

```
<ipython-input-56-46f60f40ec0e>:3: UserWarning:
     `distplot` is a deprecated function and will be removed in seaborn v0.14.0.
     Please adapt your code to use either `displot` (a figure-level function with
     similar flexibility) or `kdeplot` (an axes-level function for kernel density plots
     For a guide to updating your code to use the new functions, please see
     https://gist.github.com/mwaskom/de44147ed2974457ad6372750bbe5751
       ax = sns.distplot(y_test, hist=False, color="r", label="Actual Value")
     <ipython-input-56-46f60f40ec0e>:4: UserWarning:
     `distplot` is a deprecated function and will be removed in seaborn v0.14.0.
     Please adapt your code to use either `displot` (a figure-level function with
     similar flexibility) or `kdeplot` (an axes-level function for kernel density plots
print('MAE:', mean_absolute_error(y_test, dtree_pred))
print('MSE:', mean squared error(y test, dtree pred))
print('RMSE:', np.sqrt(mean_squared_error(y_test, dtree_pred)))
print('Accuracy:', dtree.score(x_test,y_test))
     MAE: 3481.74540546729
     MSE: 51717484.48358668
     RMSE: 7191.486945242039
     Accuracy: 0.6247649986846959
                        1
Random Forest Regressor
#distribution plot of actual and predicted values
plt.figure(figsize=(7,5))
ax = sns.distplot(y_test, hist=False, color="r", label="Actual Value")
sns.distplot(rf pred, hist=False, color="b", label="Fitted Values" , ax=ax)
plt.title('Actual vs Fitted Values for Random Forest Regressor')
plt.xlabel('Medical Expense')
plt.ylabel('Distribution')
plt.show()
```

```
<ipython-input-58-255136d82566>:3: UserWarning:
     `distplot` is a deprecated function and will be removed in seaborn v0.14.0.
     Please adapt your code to use either `displot` (a figure-level function with
     similar flexibility) or `kdeplot` (an axes-level function for kernel density plots
     For a guide to updating your code to use the new functions, please see
     https://gist.github.com/mwaskom/de44147ed2974457ad6372750bbe5751
       ax = sns.distplot(y_test, hist=False, color="r", label="Actual Value")
     <ipython-input-58-255136d82566>:4: UserWarning:
     `distplot` is a deprecated function and will be removed in seaborn v0.14.0.
     Please adapt your code to use either `displot` (a figure-level function with
     similar flexibility) or `kdeplot` (an axes-level function for kernel density plots
     For a guide to updating your code to use the new functions, please see
     https://gist.github.com/mwaskom/de44147ed2974457ad6372750bbe5751
       sns.distplot(rf_pred, hist=False, color="b", label="Fitted Values" , ax=ax)
           <sub>1e-5</sub> Actual vs Fitted Values for Random Forest Regressor
         5
print('MAE:', mean_absolute_error(y_test, rf_pred))
print('MSE:', mean squared error(y test, rf pred))
print('RMSE:', np.sqrt(mean_squared_error(y_test, rf_pred)))
print('Accuracy:', rf.score(x_test,y_test))
     MAE: 2910.5753671795514
     MSE: 26751292,149569027
     RMSE: 5172.165131699589
     Accuracy: 0.8059066243232434
                                     11
Conclusion
                   11
```

From the above models, we can see that Decision Tree Regressor and Random Forest Regressor are giving the best results. But, Random Forest Regressor is giving the best results with the least RMSE value. Therefore, I will use Random Forest Regressor to predict the medical expense of patients.

Moreover, the medical expense of smokers is higher than that of non-smokers. The medical expense of patients with BMI greater than 30 is higher than that of patients with BMI less than 30. The medical expense of older patients is higher than that of younger patients.

Thus, from the overall analysis, we can conclude that the medical expense of patients depends on their age, BMI, smoking habits.