# Analysis in Medicare Provider Utilization and Payment Data: From the Prospectives of Average Difference between Submitted and Charged Medicare Amount from Physician in California

# **Data Input**

```
In [1]:
```

```
import pandas as pd
import matplotlib.pyplot as plt
import numpy as np
import seaborn as sns
```

#### In [2]:

```
data = pd.read_csv("medical_insurance.csv")
data.head()

/anaconda3/lib/python3.6/site-packages/IPython/core/interactiveshell.py:2728: DtypeWarning: Columns (10)
) have mixed types. Specify dtype option on import or set low_memory=False.
interactivity=interactivity, compiler=compiler, result=result)
```

#### Out[2]:

	National Provider Identifier	Last Name/Organization Name of the Provider	First Name of the Provider	Middle Initial of the Provider	Credentials of the Provider	Gender of the Provider	Entity Type of the Provider		Address 2 of the	City of Provi
0	1003000126	ENKESHAFI	ARDALAN	NaN	M.D.	М	I	900 SETON DR	NaN	CUMBERL
1	1003000126	ENKESHAFI	ARDALAN	NaN	M.D.	M	I	900 SETON DR	NaN	CUMBERL,
2	1003000126	ENKESHAFI	ARDALAN	NaN	M.D.	М	1	900 SETON DR	NaN	CUMBERL
3	1003000126	ENKESHAFI	ARDALAN	NaN	M.D.	М	1	900 SETON DR	NaN	CUMBERL
4	1003000126	ENKESHAFI	ARDALAN	NaN	M.D.	М	I	900 SETON DR	NaN	CUMBERL

5 rows × 26 columns

```
data.info()
<class 'pandas.core.frame.DataFrame'>
RangeIndex: 9497892 entries, 0 to 9497891
Data columns (total 26 columns):
National Provider Identifier
                                                             int64
Last Name/Organization Name of the Provider
                                                             object
First Name of the Provider
                                                             object
Middle Initial of the Provider
                                                             object
Credentials of the Provider
                                                             object
Gender of the Provider
                                                             object
Entity Type of the Provider
                                                             object
Street Address 1 of the Provider
                                                             object
Street Address 2 of the Provider
                                                             object
City of the Provider
                                                             object
Zip Code of the Provider
                                                             object
State Code of the Provider
                                                             object
Country Code of the Provider
                                                             object
Provider Type
                                                             object
Medicare Participation Indicator
                                                             object
Place of Service
                                                             object
HCPCS Code
                                                             object
HCPCS Description
                                                             object
HCPCS Drug Indicator
                                                             object
Number of Services
                                                             float64
Number of Medicare Beneficiaries
                                                             float64
Number of Distinct Medicare Beneficiary/Per Day Services
                                                             float.64
Average Medicare Allowed Amount
                                                             float64
Average Submitted Charge Amount
                                                             float64
Average Medicare Payment Amount
                                                             float64
Average Medicare Standardized Amount
                                                             float64
dtypes: float64(7), int64(1), object(18)
memory usage: 1.8+ GB
In [4]:
# subset dataset to only Californa data
data_ca = data.loc[data['State Code of the Provider'] == 'CA']
display(data ca.shape)
(731564, 26)
```

We can see that number of samples are reduced from 9+ millions to 700 thousands. Let's start to dive more into the dataset

# **Exploratory Data Analysis**

# Missing Value

Since there's small portion of missing values in the remaining columns, these missing values are just dropped out.

```
In [5]:

# function to calculate missing value percentage

def get_missing_percentage(column):
    num = column.isnull().sum()
    total_n = len(column)
    return round(num/total_n, 2)
```

```
In [6]:

data_ca_drop = data_ca.drop(['Middle Initial of the Provider', 'National Provider Identifier','First Na
me of the Provider','Credentials of the Provider','Street Address 2 of the Provider','State Code of the
Provider','Country Code of the Provider'], axis=1)
```

rame. Conder or one rrovider, despect object

we can see that gender in male accounts for almost 74% of the data. It can be inferred from the fact that generally there are more male physicians (provider) than female physicians.

#### In [8]:

```
data_ca_drop_null = data_ca_drop[data_ca_drop.columns[data_ca_drop.isnull().any()].tolist()]
get_missing_percentage(data_ca_drop_null)
```

#### Out[8]:

```
Last Name/Organization Name of the Provider 0.00 Gender of the Provider 0.06 dtype: float64
```

Since there's only missing value in gender of the provider with missing at around 6%, therefore missing values were dropped.

#### In [9]:

```
data_ca_drop = data_ca_drop.dropna()
data_ca_drop.isnull().sum()
```

#### Out[9]:

```
Last Name/Organization Name of the Provider
                                                             Λ
                                                             0
Gender of the Provider
Entity Type of the Provider
                                                             0
Street Address 1 of the Provider
                                                             0
City of the Provider
                                                             0
Zip Code of the Provider
                                                             0
Provider Type
                                                             0
                                                             0
Medicare Participation Indicator
Place of Service
                                                             0
HCPCS Code
                                                             0
HCPCS Description
HCPCS Drug Indicator
                                                             0
                                                             0
Number of Services
Number of Medicare Beneficiaries
Number of Distinct Medicare Beneficiary/Per Day Services
                                                             0
Average Medicare Allowed Amount
Average Submitted Charge Amount
Average Medicare Payment Amount
                                                             0
Average Medicare Standardized Amount
dtype: int64
```

Now we don't have any missing value in the dataframe, let's look at more details in this data. First, let's create a column to show the difference between "Avergae submitted Charged Amount" and "Average Medicare Allowed Amount"

```
In [10]:
```

```
data_ca_drop['Average Medicare Difference'] = data_ca_drop['Average Submitted Charge Amount']- data_ca_
drop['Average Medicare Allowed Amount']
```

# Subset dataframe based on Provider Type with Top 5 highest Average Medicare Difference

Let's focus on the top 5 provider types that have the most average medicare difference.

```
In [11]:
```

```
data_pt_amaa = data_ca_drop.groupby('Provider Type')['Average Medicare Difference'].mean().reset_index(
).sort_values('Average Medicare Difference', ascending = False).head(5)
data_pt_amaa
```

# Out[11]:

	Provider Type	Average Medicare Difference
75	Thoracic Surgery	1192.946294
43	Neurosurgery	1149.126776
6	Cardiac Surgery	1146.053486
		_

79	Vascular Surgery <b>Provider Type</b>	1042.064660 Average Medicare Difference
4	CRNA	882.815789

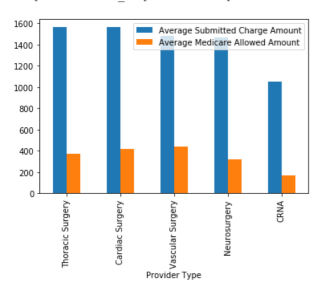
#### In [12]:

```
provider_list = ['Thoracic Surgery', 'Neurosurgery', 'Cardiac Surgery', 'Vascular Surgery', 'CRNA']
data_ca_drop_pro = data_ca_drop.loc[data_ca_drop['Provider Type'].isin(provider_list)]

# group by provider type and plot bar plot
data_ca_drop_pro.groupby('Provider Type')['Average Submitted Charge Amount', 'Average Medicare Allowed A mount'].mean().sort_values(by = 'Average Submitted Charge Amount', ascending = False).plot(kind = 'bar')
```

#### Out[12]:

<matplotlib.axes. subplots.AxesSubplot at 0x10c29b978>



# In [13]:

data\_ca\_drop\_pro.groupby('Provider Type')['Average Medicare Difference'].mean().sort\_values(ascending =
False)

# Out[13]:

```
Provider Type
Thoracic Surgery 1192.946294
Neurosurgery 1149.126776
Cardiac Surgery 1146.053486
Vascular Surgery 1042.064660
CRNA 882.815789
```

Name: Average Medicare Difference, dtype: float64

# In [14]:

```
### Top 5 Average Medicare Difference Cities
```

We may also curious about what would average medicare difference varies among cities, here, we extract top 5 cities that have most average medicare difference:

# In [15]:

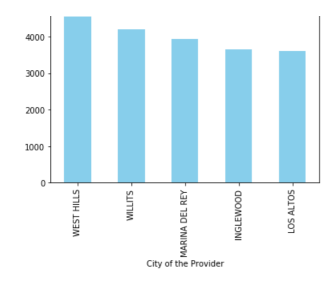
```
data_ca_drop_pro.groupby('City of the Provider')['Average Medicare Difference'].mean().sort_values(asce
nding = False).head(5).plot(kind = 'bar', color = 'skyblue')
data_ca_drop_pro.groupby('City of the Provider')['Average Medicare Difference'].mean().sort_values(asce
nding = False).head(5)
```

#### Out[15]:

```
City of the Provider
```

```
WEST HILLS 4544.530351
WILLITS 4189.692122
MARINA DEL REY 3933.719806
INGLEWOOD 3641.784906
LOS ALTOS 3608.698382
```

Name: Average Medicare Difference, dtype: float64



We can see that West Hills has the most difference in average medicare amount, with around \$4500 Now, let's subset dataframe based on top 5 provider types.

#### **ZIPCODE**

This section of code is referred from: https://www.christianpeccei.com/zipmap/

#### In [16]:

```
def read ascii boundary(filestem):
   Reads polygon data from an ASCII boundary file.
    Returns a dictionary with polygon IDs for keys. The value for each
    key is another dictionary with three keys:
    'name' - the name of the polygon
    'polygon' - list of (longitude, latitude) pairs defining the main
   polygon boundary
    'exclusions' - list of lists of (lon, lat) pairs for any exclusions in
    the main polygon
   metadata file = filestem + 'a.dat'
   data file = filestem + '.dat'
    # Read metadata
   lines = [line.strip().strip('"') for line in open(metadata file)]
   polygon ids = lines[::6]
   polygon names = lines[2::6]
   polygon data = {}
   for polygon_id, polygon_name in zip(polygon_ids, polygon_names):
        # Initialize entry with name of polygon.
        # In this case the polygon name will be the 5-digit ZIP code.
        polygon data[polygon id] = { 'name': polygon name}
   del polygon data['0']
    # Read lon and lat.
    f = open(data file)
   for line in f:
        fields = line.split()
        if len(fields) == 3:
            # Initialize new polygon
            polygon_id = fields[0]
            polygon data[polygon id]['polygon'] = []
            polygon_data[polygon_id]['exclusions'] = []
        elif len(fields) == 1:
            # -99999 denotes the start of a new sub-polygon
            if fields[0] == '-999999':
                polygon data[polygon id]['exclusions'].append([])
        else:
            # Add lon/lat pair to main polygon or exclusion
            lon = float(fields[0])
            lat = float(fields[1])
            if polygon_data[polygon_id]['exclusions']:
                polygon data[polygon id]['exclusions'][-1].append((lon, lat))
                polygon_data[polygon_id]['polygon'].append((lon, lat))
   return polygon data
```

```
In [17]:
```

```
#data_ca_drop_pro['Zip Code of the Provider'] = data_ca_drop_pro['Zip Code of the Provider'].astype('st
r')
data_ca_drop_pro['Zip Code of the Provider'] = data_ca_drop_pro['Zip Code of the Provider'].apply(lambd
a x: str(x)[:5])

/anaconda3/lib/python3.6/site-packages/ipykernel_launcher.py:2: SettingWithCopyWarning:
A value is trying to be set on a copy of a slice from a DataFrame.
Try using .loc[row_indexer,col_indexer] = value instead

See the caveats in the documentation: http://pandas.pydata.org/pandas-docs/stable/indexing.html#indexing-view-versus-copy
```

#### In [18]:

```
# group by Zip Code of the Provider
provider_zipcode = data_ca_drop_pro.groupby('Zip Code of the Provider')['Average Medicare Difference'].
mean().sort_values(ascending = False)
```

#### In [19]:

```
from pylab import *
avg_med_diff = {}

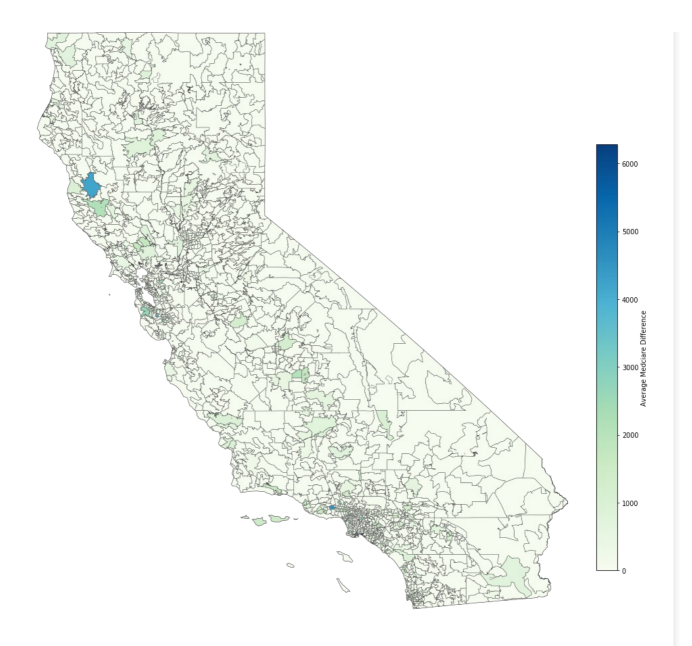
# Add data for each ZIP code
for i in range(provider_zipcode.shape[0]):
    avg_med_diff[provider_zipcode.index[i]] = provider_zipcode[i]
max_avg_med_diff = max(avg_med_diff.values())
```

#### In [20]:

```
# Read in ZIP code boundaries for California
d = read_ascii_boundary('zip5/zt06_d00')
```

#### In [21]:

```
# Create figure and two axes: one to hold the map and one to hold
# the colorbar
figure(figsize=(15, 15), dpi=70)
map axis = axes([0.0, 0.0, 0.8, 0.9])
cb axis = axes([0.83, 0.1, 0.03, 0.6])
# Define colormap to color the ZIP codes.
# You can try changing this to cm.Blues or any other colormap
# to get a different effect
cmap = cm.GnBu
# Create the map axis
axes (map axis)
axis([-125, -114, 32, 42.5])
gca().set axis off()
# Loop over the ZIP codes in the boundary file
for polygon id in d:
   polygon data = array(d[polygon id]['polygon'])
   zipcode = d[polygon_id]['name']
   avg med diff mean = avg med diff[zipcode] if zipcode in avg med diff else 0.
    # Define the color for the ZIP code
   fc = cmap(avg med diff mean/max avg med diff)
    # Draw the ZIP code
   patch = Polygon(array(polygon data), facecolor=fc,
       edgecolor=(.3, .3, .3, 1), linewidth=.4)
   gca().add patch(patch)
title('Average Medciare Difference per ZIP Code in California (2014)')
# Draw colorbar
cb = mpl.colorbar.ColorbarBase(cb_axis, cmap=cmap,
   norm = mpl.colors.Normalize(vmin=0, vmax=max avg med diff))
cb.set label('Average Medciare Difference')
```



## In [22]:

# group by city of the provider and zipcode of the provider
data\_ca\_drop\_pro.groupby(['City of the Provider','Zip Code of the Provider'])['Average Medicare Differe
nce'].mean().sort\_values(ascending = False).head(5)

#### Out[22]:

City of the Provider Zip Code of the Provider SAN DIEGO 92093 17248.377187 LONG BEACH 90803 6280.067273 WEST HILLS 91307 4544.530351 95490 4189.692122 WILLITS MARINA DEL REY 90292 3933.719806 Name: Average Medicare Difference, dtype: float64

# **Procedures in each Provider Type**

#### In [23]:

```
#subset dataframe based on provider type
data_crna = data_ca_drop_pro.loc[data_ca_drop_pro['Provider Type']=='CRNA']
data_vas = data_ca_drop_pro.loc[data_ca_drop_pro['Provider Type']=='Vascular Surgery']
data_cardiac = data_ca_drop_pro.loc[data_ca_drop_pro['Provider Type']=='Cardiac Surgery']
data_thora = data_ca_drop_pro.loc[data_ca_drop_pro['Provider Type']=='Thoracic Surgery']
data_neuro = data_ca_drop_pro.loc[data_ca_drop_pro['Provider Type']=='Neurosurgery']
```

# **Top 10 Procedures in CRNA**

#### In [24]:

```
data_crna.groupby('HCPCS Description')['Average Medicare Difference'].mean().sort_values(ascending = Fa
lse).head(10).plot(kind ="barh")
data_crna.groupby('HCPCS Description')['Average Medicare Difference'].mean().sort_values(ascending = Fa
lse).head(10)
```

#### Out[24]:

HCPCS Description

Anesthesia for procedure on heart and great blood vessels on heart-lung machine, age 1 year or older, or re-operation more than 1 month after original procedure 5002.128809

Anesthesia for heart artery bypass grafting on heart-lung machine

4772.314838

Anesthesia for kidney transplant

4252.883077

Anesthesia for procedure on spine and spinal cord

3170.178024

Anesthesia for procedure on heart and great blood vessels

3118.022305

Anesthesia for removal of prostate

3040.789101

Anesthesia for open total hip joint replacement

2489.418778

Anesthesia for procedure on heart to correct abnormal rhythm

2407.988031

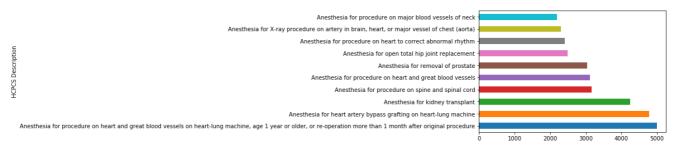
Anesthesia for X-ray procedure on artery in brain, heart, or major vessel of chest (aorta)

2299.052803

Anesthesia for procedure on major blood vessels of neck

2191.525714

Name: Average Medicare Difference, dtype: float64



It seems that anesthesia for procedures on hear and great blood vessel cost has the most differences in average medicare amount, which is around \$5000 USD. The procedure includes heart-lung usage, re-operation after original procedures. Further details about the frequency of these re-operating procedures and risk factors that contirbute to the re-operation can be discussed. But these questions will be left opened in the project.

# **Top 10 Vascular Surgery**

## In [25]:

```
data_vas.groupby('HCPCS Description')['Average Medicare Difference'].mean().sort_values(ascending = Fal
se).head(10).plot(kind ="barh")
data_vas.groupby('HCPCS Description')['Average Medicare Difference'].mean().sort_values(ascending = Fal
se).head(10)
```

#### Out[25]:

HCPCS Description

Removal of plaque and insertion of stents into artery in one leg, endovascular, accessed through the sk in or open procedure 23740.022667

Removal of plaque and insertion of stents into arteries in one leg, endovascular, accessed through the skin or open procedure 19768.319407

Removal of plaque in arteries in one leg, endovascular, accessed through the skin or open procedure 17688.368755

Removal of plaque in artery in one leg, endovascular, accessed through the skin or open procedure 14947.196645

Repair of defect of aorta in chest

9370.670909

Balloon dilation of artery of one leg, endovascular, accessed through the skin or open procedure 8738.647956

Removal of blood clot and injections (accessed through the skin) to dissolve blood clot from veins usin q fluoroscopic quidance 8160.098750

Fusion of spine bones with removal of disc at lower spinal column, anterior approach



Insertion of stents in artery in one side of groin, endovascular, accessed through the skin or open pro cedure 7086.245425

Balloon dilation of artery in one side of groin, endovascular, accessed through the skin or open proced ure 7069.189548

Name: Average Medicare Difference, dtype: float64



It seems that cost in removal of plaque and insection of stents into artery has the most differences in average medicare amount, which is \$23700 USD, followed by the procedures involving removal of plaque and insection of stents into arteriers. We can see that removel of plaque and insection of stents account for top 4 average medicare amount difference procedures in Vascular Surgery.

# **Top 10 Cardiac Surgery**

#### In [26]:

```
data_cardiac.groupby('HCPCS Description')['Average Medicare Difference'].mean().sort_values(ascending =
False).head(10).plot(kind ="barh")
data_cardiac.groupby('HCPCS Description')['Average Medicare Difference'].mean().sort_values(ascending =
False).head(10)
```

#### Out[26]:

HCPCS Description

Insertion of vena cava filter by endovascular approach, including radiological supervision and interpre tation 30757.930833

Insertion of stents in artery in one side of groin, endovascular, accessed through the skin or open procedure 17229.114615

Heart surgery procedure

16159.088667

Transplantation of donor heart

14659.190000

Complete removal of inside lining of chest cavity and lung using an endoscope 7801.992500

Repair of hole between upper heart chambers on heart-lung machine 7540.277238

Insertion of lower heart chamber blood flow assist device

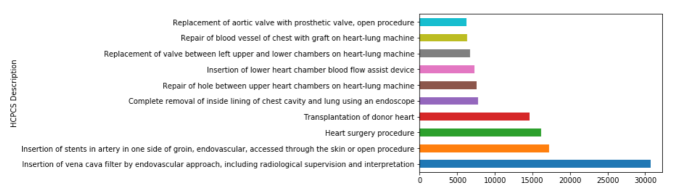
7303.768788

Replacement of valve between left upper and lower chambers on heart-lung machine 6728.743013

Repair of blood vessel of chest with graft on heart-lung machine 6352.985000

Replacement of aortic valve with prosthetic valve, open procedure 6235.826208

Name: Average Medicare Difference, dtype: float64

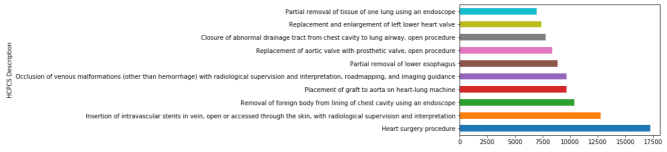


The bar chart shows that insection of vena cava by endovascular approach has the most difference in average medicare amount in cardiac surgery, which is \$30700 USD.

# **Top 10 Thoracic Surgery**

```
In [27]:
```

```
data thora.groupby('HCPCS Description')['Average Medicare Difference'].mean().sort values(ascending = F
alse) .head(10) .plot(kind ="barh")
data thora.groupby('HCPCS Description')['Average Medicare Difference'].mean().sort values(ascending = F
alse) .head (10)
Out [27]:
HCPCS Description
Heart surgery procedure
17248.377187
Insertion of intravascular stents in vein, open or accessed through the skin, with radiological supervi
sion and interpretation
                                          12753.594117
Removal of foreign body from lining of chest cavity using an endoscope
10403.839235
Placement of graft to aorta on heart-lung machine
9683.436511
Occlusion of venous malformations (other than hemorrhage) with radiological supervision and interpretat
ion, roadmapping, and imaging guidance
                                           9681.330000
Partial removal of lower esophagus
8841.641538
Replacement of aortic valve with prosthetic valve, open procedure
8396.858084
Closure of abnormal drainage tract from chest cavity to lung airway, open procedure
7774.056667
Replacement and enlargement of left lower heart valve
Partial removal of tissue of one lung using an endoscope
6982,266265
Name: Average Medicare Difference, dtype: float64
```



The procedures that involves heart surgery shows the most difference between submitted charged amount and allowed medicare amount, which is around \$17200 USD.

# **Top 10 Neurosurgery**

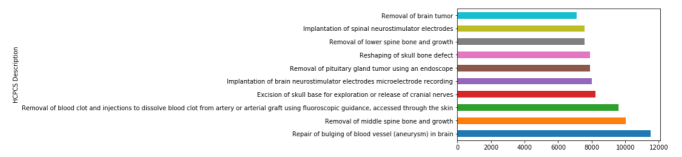
#### In [28]:

```
data_neuro.groupby('HCPCS Description')['Average Medicare Difference'].mean().sort_values(ascending = F
alse).head(10).plot(kind ="barh")
data_neuro.groupby('HCPCS Description')['Average Medicare Difference'].mean().sort_values(ascending = F
alse).head(10)
```

```
Out[28]:

HCPCS Description
Repair of bulging of blood vessel (aneurysm) in brain
11521.145454
Removal of middle spine bone and growth
10031.506923
Removal of blood clot and injections to dissolve blood clot from artery or arterial graft using fluoros
copic guidance, accessed through the skin 9610.130833
Excision of skull base for exploration or release of cranial nerves
8217.257069
Implantation of brain neurostimulator electrodes microelectrode recording
8025.686634
Removal of pituitary gland tumor using an endoscope
7914.086006
Reshaping of skull bone defect
```

7901.622421
Removal of lower spine bone and growth
7581.249946
Implantation of spinal neurostimulator electrodes
7562.391111
Removal of brain tumor
7122.118343
Name: Average Medicare Difference, dtype: float64



The largest difference in neruosurgery is the procedure of repairing of bulging of blood vessel in brain, which is around 11520 USD.

# **Feature Exploration**

#### **Correlation Matrix**

To investigate what factors would contribute to major impact on the dependent variable - average medicare difference, and understand if independent variables have correlations, correlation matrix is applied.

#### In [29]:

#### Pearson Correlation of Features



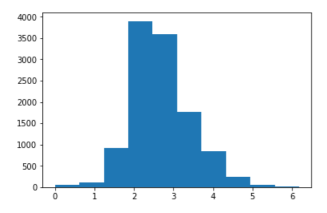
Number of Services Number of Distinct Medicare Beneficiary/Per Day Services Average Submitted Charge Amount Number of Medicare Beneficiaries

We can see that Average Medicare Allowed Amount, Average Submitted Charge Amount, Average Medicare Payment Amount, and Average Medicare Standardized Amount have higher correlation with Average Medicare Difference. We will focus on visualizing these features.

#### **Feature Transformation**

#### Average Medicare Difference

```
In [30]:
plt.hist(pow(data ca drop pro['Average Medicare Difference'], 1/6))
Out[30]:
(array([ 45., 101., 918., 3897., 3586., 1767., 847., 232.,
         20.]),
                 , 0.618311 , 1.236622 , 1.85493301, 2.47324401,
 array([0.
        3.09155501, 3.70986601, 4.32817702, 4.94648802, 5.56479902,
        6.18311002]),
 <a list of 10 Patch objects>)
```



#### **Average Medicare Allowed Amount**

4000

```
# log transformation of Average Medicare Allowed Amount
plt.hist(np.log(data_ca_drop_pro['Average Medicare Allowed Amount']))
(array([1.000e+00, 3.600e+01, 9.000e+00, 4.000e+01, 3.650e+02, 2.414e+03,
         6.335e+03, 1.611e+03, 5.960e+02, 6.800e+01]),
array([-3.91202301, -2.53018528, -1.14834756, 0.23349017, 1.61532789, 2.99716562, 4.37900334, 5.76084107, 7.14267879, 8.52451652,
          9.90635424]),
 <a list of 10 Patch objects>)
 6000
 5000
```

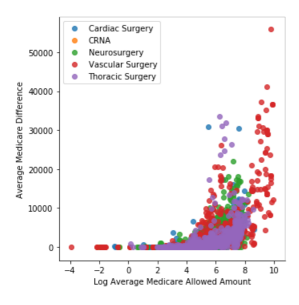
```
3000 -
2000 -
1000 -
0 -4 -2 0 2 4 6 8 10
```

#### In [32]:

```
# copy df
data_pro_copy = data_ca_drop_pro.copy()
# log transformation of
data_pro_copy['Log Average Medicare Allowed Amount'] = np.log(data_pro_copy['Average Medicare Allowed Amount'])
sns.lmplot(x = 'Log Average Medicare Allowed Amount', y='Average Medicare Difference', data = data_pro_copy, fit_reg = False, hue = 'Provider Type', legend = False)
plt.legend(loc='upper left')
```

#### Out[32]:

<matplotlib.legend.Legend at 0x1a19b03c18>



We could see there's more variations in vascular surgery. With more allowed average medicare amount, there's a increasing trend in variations in average medicare difference and the amount of average medicare difference.

# **Average Medicare Payment Amount**

<a list of 10 Patch objects>)

```
In [33]:
```

1000

```
6000 -
5000 -
4000 -
3000 -
2000 -
```

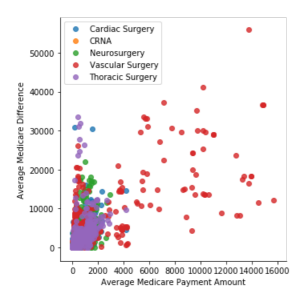
```
-4 -2 0 2 4 6 8 10
```

#### In [34]:

```
# scatter plot
sns.lmplot(x = 'Average Medicare Payment Amount', y='Average Medicare Difference', data = data_ca_drop_
pro, fit_reg = False, hue = 'Provider Type', legend = False)
plt.legend(loc='upper left')
```

#### Out[34]:

<matplotlib.legend.Legend at 0x1a1af55198>



The scatter plot showed that vascular surgery varied the most in average medicare payment amount that medicare covered after coinsurance amount deducted compared to other surgery, and it can be visualized that there's slightly positive correlation between average medicare payment amount and average medicare difference.

# **Creating Dummy Variables**

Now, I want to know that what factors would affect the average submitted charge amount. To do so, I need to get dummies for each categorical value to save space and ease computatinal complexity.

```
In [35]:
```

```
# convert categorical variable to dummy variable
data_ca_drop_dummy = pd.get_dummies(data_ca_drop_pro[['Zip Code of the Provider','Entity Type of the Pr
ovider','Provider Type','Medicare Participation Indicator','HCPCS Description','HCPCS Drug Indicator','
City of the Provider']])
```

#### In [36]:

```
# build continuous variable dataframe
data_ca_drop_continue = data_ca_drop_pro[['Number of Services','Number of Medicare Beneficiaries','Numb
er of Distinct Medicare Beneficiary/Per Day Services','Average Medicare Payment Amount','Average Medica
re Standardized Amount','Average Medicare Difference']]
```

#### In [37]:

```
# normalize continuous dataframe
data_ca_drop_continue = (data_ca_drop_continue-data_ca_drop_continue.min())/(data_ca_drop_continue.max())-data_ca_drop_continue.min())
```

#### In [38]:

```
# concat binary dataframe and continuous dataframe
data_ca_drop_dummy = pd.concat([data_ca_drop_continue,data_ca_drop_dummy], axis = 1)
```

#### In [39]:

```
from sklearn.preprocessing import StandardScaler
# run lineagr regression model
data_ca_drop_dummy.reset_index(drop = True)
V = data_ca_drop_dummy.l_Average_Medicare_Difference!l
```

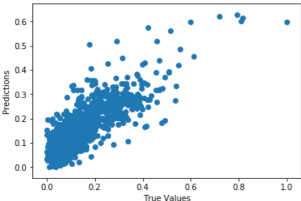
```
X = data_ca_drop_dummy.drop(['Average Medicare Difference'], axis = 1)
In [401:
from sklearn.model_selection import train_test_split
In [41]:
x_train, x_test, y_train, y_test = train_test_split(X, Y, test_size=0.2)
Linear Regression Model as a baseline model
In [42]:
from sklearn import linear model
from sklearn.linear model import LinearRegression
from sklearn import metrics
from sklearn.model selection import GridSearchCV
from sklearn.linear model import ElasticNet, ElasticNetCV
from sklearn.metrics import mean squared error, r2 score
In [43]:
model = linear model.LinearRegression()
model.fit(x train, y train)
y pred lr = model.predict(x_test)
print ("R Squared: ", model.score(x test, y test))
print("Residual sum of squares: %.2f"
             % np.mean((model.predict(x test) - y test) ** 2))
print ("Linear Regression MSE: %.4f"%mean_squared_error(y_test, y_pred_lr))
R Squared: -4.722544063284281e+21
Residual sum of squares: 10821449170476201984.00
Linear Regression MSE: 10821449170476197888.0000
Elastic Regression Model
In [44]:
cv enet = ElasticNetCV(11 ratio = np.linspace(0.1,1,40.), cv = 10, eps = 0.001, n alphas = 100, fit int
ercept = True, normalize = True, max iter = 2000)
/anaconda3/lib/python3.6/site-packages/ipykernel launcher.py:1: DeprecationWarning: object of type <cla
ss 'float'> cannot be safely interpreted as an integer.
 """Entry point for launching an IPython kernel.
```

```
In [45]:
cv enet.fit(x train, y train)
ElasticNetCV(alphas=None, copy_X=True, cv=10, eps=0.001, fit_intercept=True,
                            , 0.12308, 0.14615, 0.16923, 0.19231, 0.21538, 0.23846,
      ll ratio=array([0.1
      0.26154, 0.28462, 0.30769, 0.33077, 0.35385, 0.37692, 0.4
      0.42308, 0.44615, 0.46923, 0.49231, 0.51538, 0.53846, 0.56154,
      0.58462, 0.60769, 0.63077, 0.65385, 0.67692, 0.7 , 0.72308,
      0.74615, 0.76923, 0.79231, 0.81538, 0.83846, 0.86154, 0.88462,
      0.90769, 0.93077, 0.95385, 0.97692, 1.
                                                ]),
      max_iter=2000, n_alphas=100, n_jobs=1, normalize=True,
      positive=False, precompute='auto', random state=None,
      selection='cyclic', tol=0.0001, verbose=0)
In [46]:
print ("optimal l1 ratio: %.3f"%cv enet.l1 ratio )
print ("optimal alpha: %.6f"%cv_enet.alpha )
print ("number of iterations : %d"%cv_enet.n_iter_)
optimal l1_ratio: 0.100
optimal alpha: 0.000009
number of iterations: 42
```

The I1 ratio is 0.1, which means ridge regression accounts majority part in the elastic net. This is reasonable because we may have

collinearity and ridge solve collinearity issue better than lasso regressor does.

```
In [47]:
net_model = ElasticNet(11_ratio = cv_enet.11_ratio_, alpha = cv_enet.alpha_, max_iter = cv_enet.n_iter_
, fit intercept = True, normalize = True)
In [48]:
net model.fit(x train, y train)
Out[48]:
ElasticNet(alpha=9.335318823677131e-06, copy X=True, fit intercept=True,
      11_ratio=0.1, max_iter=42, normalize=True, positive=False,
      precompute=False, random state=None, selection='cyclic', tol=0.0001,
      warm start=False)
In [49]:
# MSE
y pred net = net model.predict(x test)
print("Elastic Net MSE: {}".format(mean_squared_error(y_test, y_pred_net)))
# R squared
print ("Elastic Net R-Squared: {}".format(r2_score(y_test, y_pred_net)))
Elastic Net MSE: 0.000778326776968653
Elastic Net R-Squared: 0.6603336168785282
In [50]:
plt.scatter(np.sqrt(y_test), np.sqrt(y_pred_net))
plt.xlabel("True Values")
plt.ylabel("Predictions")
/anaconda3/lib/python3.6/site-packages/ipykernel_launcher.py:1: RuntimeWarning: invalid value encounter
  """Entry point for launching an IPython kernel.
Out[50]:
Text(0,0.5,'Predictions')
```



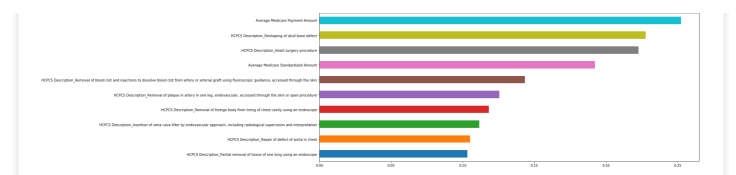
# **Feature Importance from Elastic Net**

```
In [51]:

def get_feature_importance(x_train, model):

    feature_importance = pd.Series(index = x_train.columns, data = np.abs(model.coef_))
    selected_features = (feature_importance>0).sum()
    print('{0:d} features, reduction of {1:2.2f}%'.format(
        selected_features, (1-selected_features/len(feature_importance))*100))
    feature_importance.sort_values().tail(10).plot(kind = 'barh', figsize = (20,8))
```

```
In [52]:
get_feature_importance(x_train, net_model)
684 features, reduction of 44.70%
```



# **Random Forest Training and Prediction**

```
In [53]:
```

```
from sklearn.ensemble import RandomForestRegressor

# random forest model
rf = RandomForestRegressor(n_estimators = 200, min_samples_leaf = 2, min_samples_split = 15)

# train random forest regressor
rf_model = rf.fit(x_train, y_train)

# Use the forest's predict method on the test data
y_pred_rf = rf.predict(x_test)
```

#### In [54]:

```
# MSE
print("Random Forest MSE: {}".format(mean_squared_error(y_test, y_pred_rf)))
# R squared
print ("Random Forest R-Squared: {}".format(r2_score(y_test, y_pred_rf)))
```

Random Forest MSE: 0.0007623087960162973 Random Forest R-Squared: 0.6673239579743151

# **Feature Importance from Random Forest**

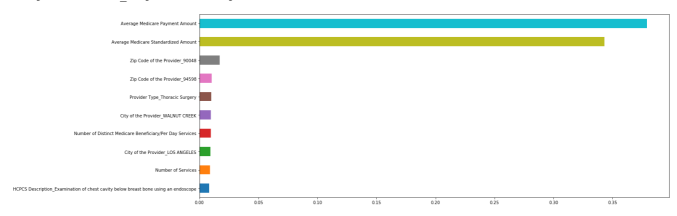
# In [55]:

```
# feature importance
feature_importance = pd.Series(index = x_train.columns, data = np.abs(rf_model.feature_importances_))
selected_features = (feature_importance>0).sum()
print('{0:d} features, reduction of {1:2.2f}%'.format(
    selected_features, (1-selected_features/len(feature_importance))*100))
feature_importance.sort_values().tail(10).plot(kind = 'barh', figsize = (20,8))
```

745 features, reduction of 39.77%

#### Out[55]:

<matplotlib.axes.\_subplots.AxesSubplot at 0x1a1dda6710>



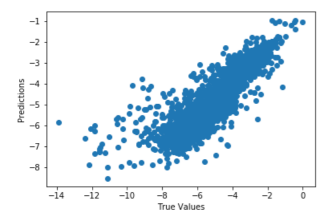
#### In [56]:

```
plt.scatter(np.log(y_test), np.log(y_pred_rf))
plt.xlabel("True Values")
plt.vlabel("Predictions")
```

```
/anaconda3/lib/python3.6/site-packages/ipykernel_launcher.py:1: RuntimeWarning: divide by zero encounte red in log
"""Entry point for launching an IPython kernel.
```

```
Out[56]:
```

```
Text(0,0.5,'Predictions')
```



# **eXtreme Gradient Boosting**

```
In [57]:
#!pip install xgboost
from xgboost import XGBRegressor
from sklearn.ensemble import GradientBoostingRegressor
In [58]:
print (x train.shape)
print(y_train.shape)
(9180, 1237)
(9180,)
In [59]:
params = { 'min_child_weight': [4,5], 'gamma': [i/10.0 for i in range(3,6)], 'subsample': [i/10.0 for i in
range(6,11)],
'colsample_bytree':[i/10.0 for i in range(6,11)], 'max_depth': [2,3,4]}
xgb = XGBRegressor(colsample bytree=0.2, gamma=0.0,
                             learning rate=0.05, max depth=6,
                             min child weight=1.5, n estimators=200,
                             reg alpha=0.9, reg lambda=0.6,
                             subsample=0.2, seed=42)
xgb_grid = GridSearchCV(xgb, params)
xgb.fit(x train,y train)
y_pred_xgb = xgb.predict(x_test)
```

### In [60]:

```
# MSE
print("Xgboost MSE: {}".format(mean_squared_error(y_test, y_pred_xgb)))

# R squared
print ("Xgboost R-Squared: {}".format(r2_score(y_test, y_pred_xgb)))
```

Xgboost MSE: 0.0008630282118585724 Xgboost R-Squared: 0.6233694125294127

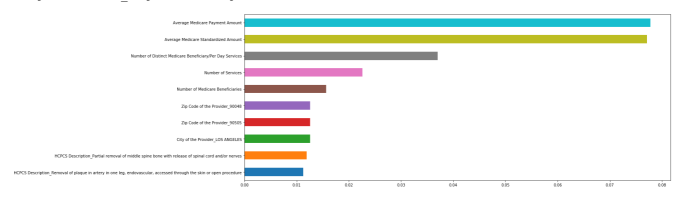
# In [61]:

```
# feature importance
feature_importance = pd.Series(index = x_train.columns, data = np.abs(xgb.feature_importances_))
selected_features = (feature_importance>0).sum()
print('{0:d} features, reduction of {1:2.2f}%'.format(
    selected_features, (1-selected_features/len(feature_importance))*100))
feature_importance.sort_values().tail(10).plot(kind = 'barh', figsize = (20,8))
```

318 features, reduction of 74.29%

#### Out[61]:

<matplotlib.axes. subplots.AxesSubplot at 0x1a18d097b8>

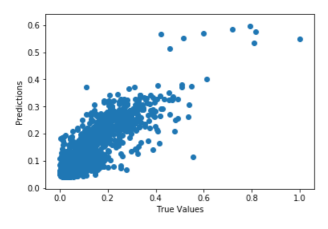


#### In [62]:

```
# plot prediciton values vs true values
plt.scatter(np.sqrt(y_test), np.sqrt(y_pred_xgb))
plt.xlabel("True Values")
plt.ylabel("Predictions")
```

#### Out[62]:

Text(0,0.5,'Predictions')



# In [63]:

```
# import MLP classifier
from sklearn.neural_network import MLPRegressor

# initialize MLP classifier
mlp = MLPRegressor(hidden_layer_sizes=(30,30,30))

# train MLP classifier
mlp.fit(x_train,y_train)

# predict
y_pred_mlp = mlp.predict(x_test)
```

#### In [64]:

```
# MSE
print("Neural Network MSE: {}".format(mean_squared_error(y_test, y_pred_mlp)))
# R squared
print ("Neural Network R-Squared: {}".format(r2_score(y_test, y_pred_mlp)))
```

Neural Network MSE: 0.000680317208914271 Neural Network R-Squared: 0.703105568811091

#### In [65]:

```
# plot prediciton values vs true values
plt.scatter(np.sqrt(y_test), np.sqrt(y_pred_mlp))
plt.xlabel("True Values")
plt.ylabel("Predictions")
```

/anaconda3/lib/python3.6/site-packages/ipykernel\_launcher.py:2: RuntimeWarning: invalid value encounter ed in sqrt

# 

# **Stacking**

```
In [116]:
```

```
from sklearn.model_selection import StratifiedKFold
from xgboost.sklearn import XGBClassifier
from sklearn.ensemble import RandomForestClassifier
from sklearn.linear_model import LogisticRegression
from sklearn.metrics import accuracy_score
from sklearn import cross_validation
from sklearn.cross_validation import KFold
import plotly.graph_objs as go
import plotly.offline as py
py.init_notebook_mode(connected=True)
```

#### In [67]:

# x's dimension and y's dimension

```
print (x_train.shape)
print(y_train.shape)

(9180, 1237)
(9180,)

In [68]:

ntrain = x_train.shape[0]
ntest = x_test.shape[0]

SEED = 43
kf = KFold(ntrain, n_folds= 10, random_state=SEED)
```

#### In [72]:

```
def get_train_test_per_model(clf, x_train, y_train, x_test):
    train = np.zeros((ntrain,))
    test = np.zeros((ntest,))
    test_skf = np.empty((10, ntest))

for i, (train_index, test_index) in enumerate(kf):
    x_tr = x_train.iloc[train_index]
    y_tr = y_train.iloc[train_index]
    x_te = x_train.iloc[test_index]

    clf.fit(x_tr, y_tr)

    train[test_index] = clf.predict(x_te)
    test_skf[i, :] = clf.predict(x_test)

test[:] = test_skf.mean(axis=0)
    return train.reshape(-1, 1), test.reshape(-1, 1)
```

```
In [73]:
# train all models
# Create our OOF train and test predictions. These base results will be used as new features
rf_train_stack, rf_test_stack = get_train_test_per_model(rf, x_train, y_train, x_test) # Random Forest
xgb_train_stack, xgb_test_stack = get_train_test_per_model(xgb,x_train, y_train, x_test) # Xgboost
mlp_train_stack, mlp_test_stack = get_train_test_per_model(mlp, x_train, y_train, x_test) # Neural Neto
In [74]:
predictions train stack = pd.DataFrame( { 'RandomForest': rf train stack.ravel(),
      'GradientBoost': xgb train stack.ravel(),
      'NeuralNetowrk': mlp_train_stack.ravel()
In [75]:
heatmap = [
   go.Heatmap(
        z= predictions train stack.astype(float).corr().values ,
        x= predictions_train_stack.columns.values,
        y= predictions_train_stack.columns.values,
         colorscale='YlGnBu',
           showscale=True,
           reversescale = True
py.iplot(heatmap, filename='labeled-heatmap')
```

# In [119]:

```
### Do combinations of models (stacking)
from sklearn.linear_model import LogisticRegression
from sklearn.metrics import accuracy_score
from sklearn.metrics import log_loss
from sklearn import cross_validation

#define parameter for stacking
model_list = ['XGBoost', 'MLP', 'RandomForest']
model_len = len(model_list)
opt_dict = {}

# Change df to np.array
X1 = X
X2 = Y
```

```
X3 = X
y = Y
pred= np.zeros((3,9180))
pred xgb = pd.Series([])
pred_mlp = pd.Series([])
pred rf = pd.Series([])
# Do 10-fold train & evaluate
cv = cross validation. KFold(len(x train), n folds=10, shuffle=False, random state=None)
for traincy, testcy in cv:
    X train1, X test1= (X1.iloc[traincv], X1.iloc[testcv])
    X train2, X test2= (X2.iloc[traincv], X2.iloc[testcv])
    X train3, X test3 = (X3.iloc[traincv], X3.iloc[testcv])
    y_train1, y_test1= (y.iloc[traincv], y.iloc[testcv])
    # XGB
    xgb= XGBRegressor(colsample bytree=0.2, gamma=0.0,
                             learning rate=0.05, max depth=8,
                             min_child_weight=1.5, n_estimators=200,
                             reg alpha=0.9, reg lambda=0.6,
                             subsample=0.2, seed=42)
    xgb.fit(X train1,y train1)
    pred xgb temp = xgb.predict(X test1)
    pred_xgb = pd.concat([pred_xgb,pd.Series(pred_xgb_temp)], axis = 1)
    # MLP classifier
    mlp = MLPRegressor(hidden layer sizes=(30,30,30))
    mlp.fit(X train2,y train1)
    pred_mlp_temp = mlp.predict(X_test2)
    pred_mlp = pd.concat([pred_mlp,pd.Series(pred_mlp_temp)], axis = 1)
    #random forest regressor
    rf = RandomForestRegressor(n estimators = 200, min samples leaf = 2, min samples split = 15)
    rf model = rf.fit(X train3, y train1)
    pred_rf_temp = rf.predict(X_test1)
    pred rf = pd.concat([pred rf,pd.Series(pred rf temp)], axis = 1)
In [123]:
#average result from cross validation by row
xgb_result = pred_xgb.mean(axis = 1)
mlp result = pred mlp.mean(axis = 1)
rf_result = pred_rf.mean(axis = 1)
#combine three model
pred_combine = pd.concat([xgb_result, mlp_result, rf_result], axis = 1)
pred combine.columns = ['xgb','mlp','rf']
In [151]:
def get optimal MSE(allocs, pred df, eps=1e-15):
    pred df = np.clip(pred df, eps, 1 - eps)
    pred df w = np.sum(pred df*allocs,axis = 1)
    return mean squared error(y test1, pred df w)
In [152]:
```

```
model_list = ['XGBoost', 'MLP', 'RandomForest']
model_len = len(model_list)
init_vals= [1.0 / model_len] * model_len
cons = ({ 'type': 'ineq', 'fun': lambda x: 1.0-np.sum(x)})
bnds = [(0.0, 1.0)] * model_len

#optimized allocations
opts = spo.minimize(get_optimal_MSE, init_vals, args = (pred_combine,),method='SLSQP', bounds=bnds, con
straints=cons, options = { 'disp':True})
opt_allocs = opts.x
```

```
Optimization terminated successfully. (Exit mode 0)

Current function value: 0.0019119187971855295

Iterations: 1

Function evaluations: 5
```

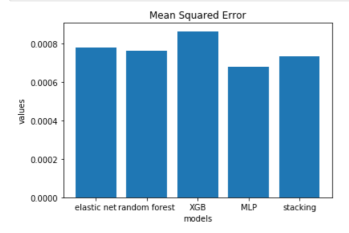
```
Gradient evaluations: 1
In [153]:
# optimal wieghts for each learner
display(opt_allocs)
# optimal result
display(opts)
array([0.33333333, 0.33333333, 0.33333333])
     fun: 0.0019119187971855295
     jac: array([-2.25775148e-05, 2.54604238e-05, -6.28369744e-05])
 message: 'Optimization terminated successfully.'
    nfev: 5
    nit: 1
    njev: 1
  status: 0
 success: True
      x: array([0.33333333, 0.33333333, 0.33333333])
In [154]:
y stack pred = 0.33333333*mlp.predict(x test) + 0.33333333*xgb.predict(x test) + 0.33333333*rf.predict
(x_test)
In [155]:
# MSE
print("Stacking MSE: {}".format(mean squared error(y test, y stack pred)))
print ("Stacking R-Squared: {}".format(r2_score(y_test, y_stack_pred)))
Stacking MSE: 0.000732264623199042
Stacking R-Squared: 0.6804354116936113
In [158]:
# plot prediciton values vs true values
\verb|plt.scatter(np.log(y_test), np.log(y_stack_pred))|\\
plt.xlabel("True Values")
plt.ylabel("Predictions")
/anaconda3/lib/python3.6/site-packages/ipykernel_launcher.py:2: RuntimeWarning:
divide by zero encountered in log
/anaconda3/lib/python3.6/site-packages/ipykernel launcher.py:2: RuntimeWarning:
invalid value encountered in log
Out[158]:
Text(0,0.5,'Predictions')
    -2
    -4
    -6
   -8
```

# -14 - -12 -10 -8 -6 -4 -2 (

# **Comparison between Models**

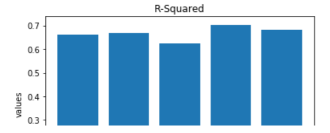
-10 -12

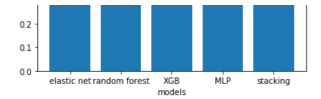
```
، رسان عالم
# Mean Squared Error
model_mse = [
            mean_squared_error(y_test, y_pred_net),\
            mean_squared_error(y_test, y_pred_rf),\
            mean_squared_error(y_test, y_pred_xgb),\
y_pos = np.arange(len(bar_cat))
# Create bars
plt.bar(y_pos, model_mse)
# Create names on the x-axis
plt.xticks(y_pos, bar_cat)
# Add title and axis names
plt.title('Mean Squared Error')
plt.xlabel('models')
plt.ylabel('values')
# Show graphic
plt.show()
```



# In [165]:

```
# R-Squared
model_r\_squared = [
              r2_score(y_test, y_pred_net),\
              r2_score(y_test, y_pred_rf),\
              r2_score(y_test, y_pred_xgb),\
              r2_score(y_test, y_pred_mlp),\
              r2_score(y_test, y_stack_pred)]
bar_cat_r = ('elastic net', 'random forest', 'XGB', 'MLP', 'stacking')
y_pos_r = np.arange(len(bar_cat_r))
# Create bars
plt.bar(y_pos_r, model_r_squared)
# Names on x-axis
plt.xticks(y_pos_r, bar_cat_r)
# title and axis names
plt.title('R-Squared')
plt.xlabel('models')
plt.ylabel('values')
# Show graphic
plt.show()
```





The result showded that in addition to baseline model linear regression, the rest five models significantly reduced mean squared error, at the same time, R-Sqaured significantly were increased among the rest five models. Among the five models, MLP model showed the lowest mean squared error and highest R-Squared. It is intersted to note that MLP perform the same or even slightly better than the stacking model.

Noted that stacking model was built based on the mean value from cross validation and the training set and testing set were resampled, it is unlikely that stacking model did not perform the best because of overfitting issue. Instead, it may be explained by the fact that the model may be suboptimal. It can be inferred that neural network performed the best because random forest regressor and xgboosting regressor did not fit to the optimal parameters, grid search can be implemented to get the optimal value and increase performace in stacking model.

Also, noted that random forest regressor also outperformed xgbkosting regressor, it is very likely that xgboosting regressor did not for to the optimal values. It is also possible that neural newtwork may better predict numerical value compared to xgboosting and random forest in this case. For the future work, optimal parameters should be achieved. In addition to MSE and R-Squared, other analysis can be also conducted to accurately define what is the "best model".