# **Medical Insurance Cost prediction**

This exercise is about performing some of the steps described in the notebook for the California Housing Data on another dataset for Medical Insurance Cost prediction.

#### Get the Data

```
In []: import pandas as pd
        medical = pd.read_csv("https://bit.ly/44evDuW")
```

## Take a Quick Look at the Data Structure

```
# display the first 5 rows of the dataset by calling the head() function on medical
medical.head()
                 bmi children smoker
  age
          sex
                                         region
                                                    charges
    19
       female 27.900
                            0
                                  yes southwest 16884.92400
               33.770
                                                  1725.55230
         male
                                      southeast
                                                 4449.46200
2
    28
         male
              33.000
                            3
                                   no southeast
3
    33
         male
              22.705
                            0
                                   no
                                       northwest 21984.47061
    32
              28.880
                                                 3866.85520
         male
```

Each row represents one patient. There are 7 attributes.

The info() method is useful to get a quick description of the data, in particular the total number of rows, each attribute's type, and the number of non-null values:

```
In [ ]: # get the number of rows, columns, and data types by using the info() method
        medical.info()
        <class 'pandas.core.frame.DataFrame'>
        RangeIndex: 1338 entries, 0 to 1337
        Data columns (total 7 columns):
             Column
                       Non-Null Count Dtype
        - - -
                       1338 non-null
         0
             age
                       1338 non-null
         1
             sex
                                        object
             bmi
                       1338 non-null
                                        float64
         3
             children 1338 non-null
                                        int64
                       1338 non-null
                                        object
             smoker
                       1338 non-null
             region
                                        object
         6
             charges
                       1338 non-null
                                        float64
        dtypes: float64(2), int64(2), object(3)
        memory usage: 73.3+ KB
        # show the number of patients in each region by using the value counts() method on the "region" column
In [ ]:
        medical["region"].value counts()
        southeast
                      364
Out[]:
                      325
        southwest
                     325
        northwest
        northeast
                     324
        Name: region, dtype: int64
```

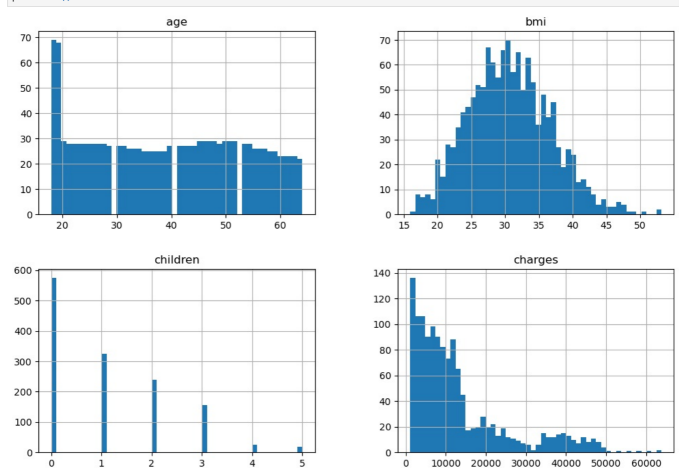
Let's look at the other fields. The describe() method shows a summary of the numerical attributes.

```
In [ ]: # show descriptive statistics for the dataset by calling the describe() method on medical
        medical.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

Out[ ]:

```
In []: # show histograms for the numerical columns by using the hist() method on medical
import matplotlib.pyplot as plt
medical.hist(bins=50, figsize=(12, 8))
plt.show()
```



Briefly write here what you observe from these histograms.

Based on the histograms, we can see that there is a spike for the group of people who are under the age of 20, which also contributes a lot to the category of people who have 0 children. It is also noticeable that a significant number of people have a BMI in between 25 and 35. Many people have medical charges between 0 and 10000. I am not sure what the unit should be for the charges since no specific details are given.

### Create a Test Set

```
In [ ]: # use train_test_split() to split the data into training and test sets
    from sklearn.model_selection import train_test_split

train_set, test_set = train_test_split(medical, test_size=0.2, random_state=36)
```

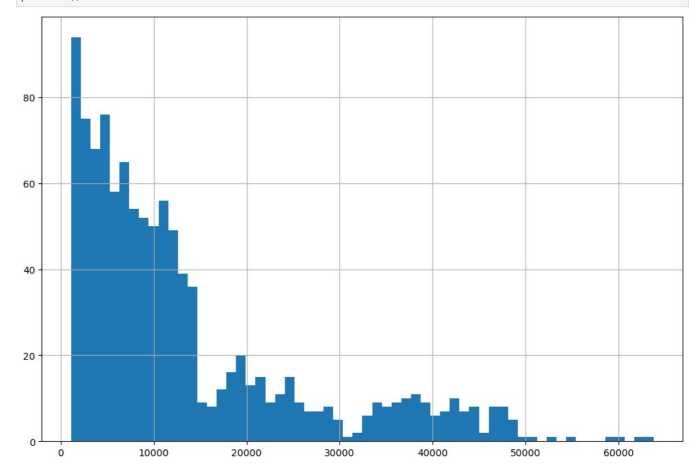
# Explore and Visualize the Data to Gain Insights

So far you have only taken a quick glance at the data to get a general understanding of the kind of data you are manipulating. Now the goal is to go into a little more depth.

First, make sure you have put the test set aside and you are only exploring the training set.

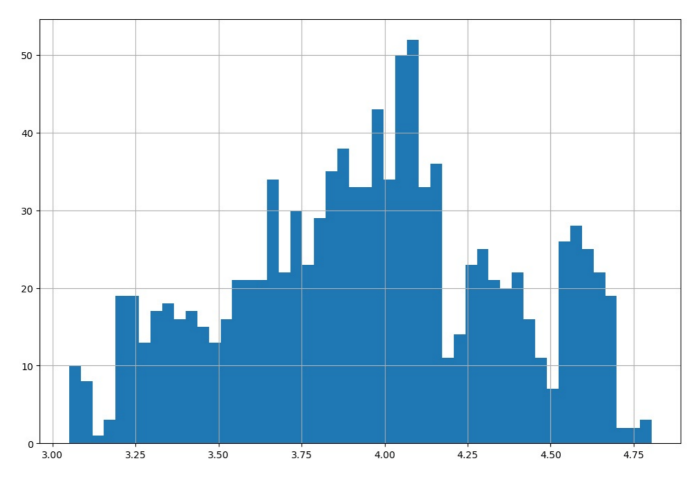
```
medical = train_set.copy()

In []: # build a histogram of the charges column
    medical["charges"].hist(bins=60, figsize=(12, 8))
    plt.show()
```



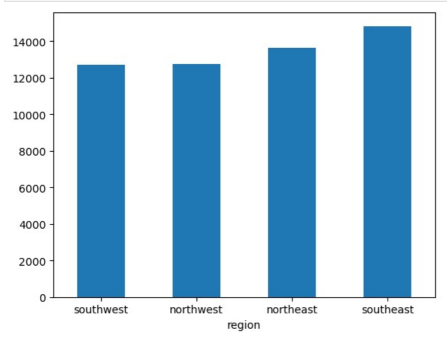
This distribution is right-skewed. To make it closer to normal we can apply natural log

```
In []: # apply a log transformation to the charges column using the np.log10() function
    import numpy as np
    transformed_charges = np.log10(medical["charges"])
    # build a histogram of the transformed column
    transformed_charges.hist(bins=50, figsize=(12, 8))
    plt.show()
```



Now let's look at the mean charges by region

```
In []: # compute the average insurance cost for each region
    # sort the charges_by_region Series from the lowest to highest cost
# plot the sorted Series using the plot.bar() method
medical.groupby(["region"]).mean()["charges"].sort_values().plot.bar(rot=0)
plt.show()
```



Overall the highest medical charges are in the Southeast and the lowest are in the Southwest. Taking into account certain factors (sex, smoking, having children) let's see how it changes by region.

Now, create three grouped barcharts for average charges by region grouped by sex, smoking, and number of children.

#### How to create grouped barcharts?

Creating grouped bar charts with Seaborn is a bit more intuitive compared to Matplotlib. You can use the catplot function with kind='bar' to create grouped bar charts. Here is an example on the tips datasets that comes with Seaborn. The tips dataset contains information about the total bill and tip amount for different meals, along with additional information such as the sex of the individual paying for the meal, whether they are a smoker, the day and time of the meal, and the size of the party.

We will create a grouped bar chart showing the average total bill for each day, grouped by whether the meal took place at lunch or dinner.

```
import seaborn as sns
import matplotlib.pyplot as plt

# Load the 'tips' dataset
tips = sns.load_dataset("tips")

# Create a grouped bar chart
sns.catplot(data=tips, x="day", y="total_bill", hue="time", kind="bar")
plt.show()
```

In this plot, the height of the bars represents the average total bill for meals on each day, with separate bars for lunch and dinner.

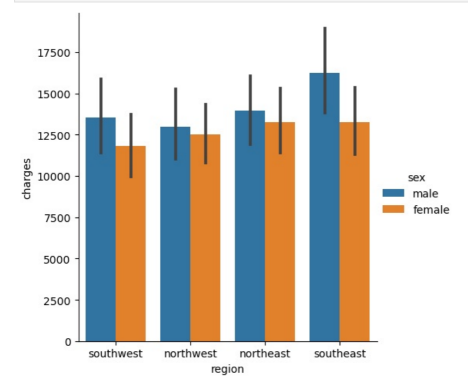
The catplot function is a flexible function that can create a variety of different plot types. By setting kind='bar', we specify that we want a bar chart. The x and y arguments specify the data for the x and y axes, and the hue argument specifies a third variable that is used to group the data.

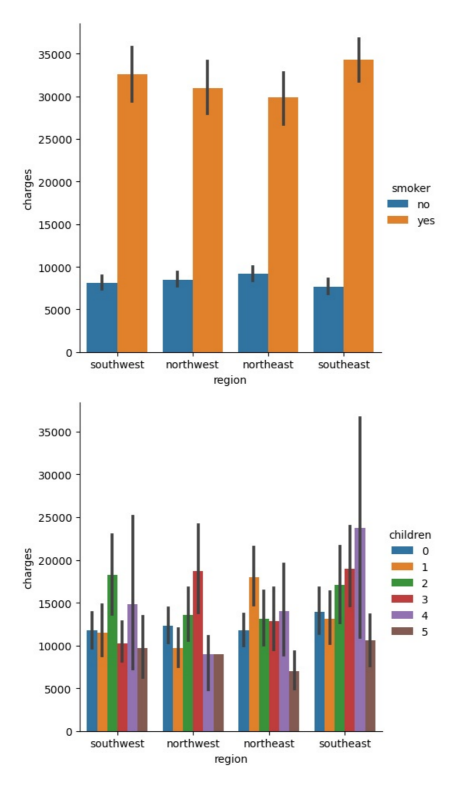
In the context of Seaborn and many other statistical visualization libraries, error bars commonly represent one standard deviation or standard error of the mean.

```
In []: # plot grouped bar charts of region and insurance costs hue by sex, smoker, and number of children (three separ
# use the catplot() function to create the bar charts
# set the kind parameter to "bar" and the data parameter to medical
import seaborn as sns
sns.catplot(data=medical, x="region", y="charges", hue="sex", kind="bar")
plt.show()

sns.catplot(data=medical, x="region", y="charges", hue="smoker", kind="bar")
plt.show()

sns.catplot(data=medical, x="region", y="charges", hue="children", kind="bar")
plt.show()
```



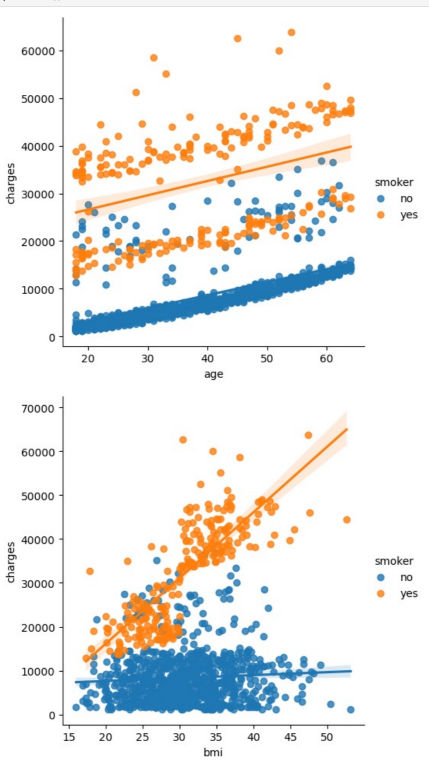


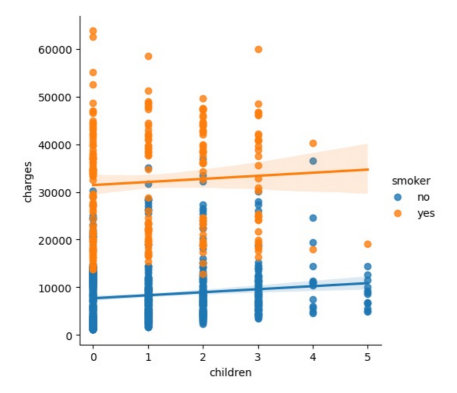
What do you observe? Briefly write what you observe from the charts.

Now let's analyze the medical charges by age, bmi and children according to the smoking factor.

```
In []: # using the lmplot() function of seaborn, build a scatter plot of age and insurance costs, hue by smoker
    sns.lmplot(data=medical, x="age", y="charges", hue="smoker")
    plt.show()
# build a second scatter plot of bmi and insurance costs, hue by smoker
```

```
sns.lmplot(data=medical, x="bmi", y="charges", hue="smoker")
plt.show()
# build a third scatter plot of children and insurance costs, hue by smoker
sns.lmplot(data=medical, x="children", y="charges", hue="smoker")
plt.show()
```





Describe in a one-liner what you obeserve from the charts.

#### Look for Correlations

```
In [ ): # compute pairwise correlation of columns using the corr() method
    medical.corr()
```

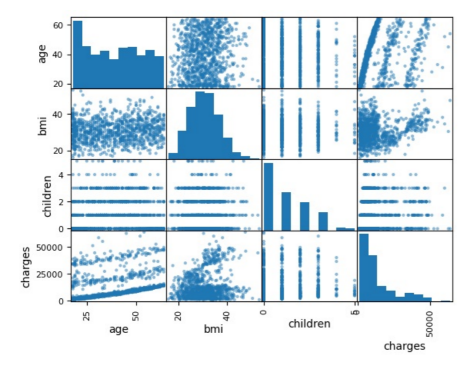
Out[ ]:	]:		age	bmi	children	charges
		age	1.000000	0.094175	0.044474	0.304674
		bmi	0.094175	1.000000	0.000124	0.182822
		children	0.044474	0.000124	1.000000	0.070805
		charges	0.304674	0.182822	0.070805	1.000000

The correlation coefficient ranges from -1 to 1. When it is close to 1, it means that there is a strong positive correlation. Finally, coefficients close to 0 mean that there is no linear correlation.

Another way to check for correlation between attributes is to use the Pandas scatter\_matrix() function, which plots every numerical attribute against every other numerical attribute.

```
In [ ]: # plot correlation matrix using scatter_matrix() function from pandas.plotting
    from pandas.plotting import scatter_matrix

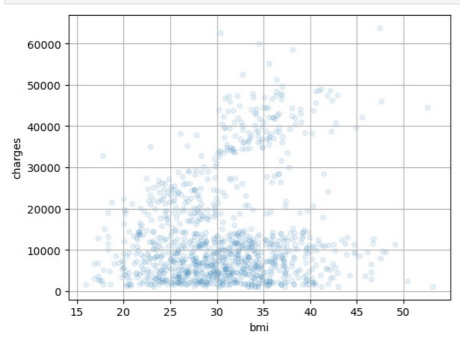
scatter_matrix(medical)
    plt.show()
```



The main diagonal would be full of straight lines if Pandas plotted each variable against itself, which would not be very useful. So instead, the Pandas displays a histogram of each attribute.

Looking at the correlation scatterplots, it seems like the most promising attribute to predict the charge value is bmi, so let's zoom in on their scatterplot.

```
In [ ]: # plot a scatter plot of bmi vs. insurance costs using the medical.plot() method, use the alpha parameter to se
medical.plot(kind="scatter", x="bmi", y="charges", alpha=0.1, grid=True)
plt.show()
```



The correlation is somewhat visible; you can clearly see the upward trend.

# Prepare the data for ML

```
# drop the charges column from the train_set and save the resulting dataset to a variable called `medical`
# create a copy of the train_set labels and save it to a variable called `medical_labels`
# replace None with the correct code

medical = train_set.drop("charges", axis=1)
medical_labels = train_set["charges"].copy()
```

# **Transformation Pipelines**

As you can see, there are many data transformation steps that need to be executed in the right order. Fortunately, Scikit-Learn provides the Pipeline class to help with such sequences of transformations.

```
In [ ]: # uncomment the following code to create a pipeline for preprocessing the data
         from sklearn.compose import ColumnTransformer
         from sklearn.impute import SimpleImputer
         from sklearn.preprocessing import StandardScaler
         from sklearn.preprocessing import OneHotEncoder
         from sklearn.pipeline import make_pipeline
         num_attribs = ["age", "bmi", "children"]
         cat_attribs = ["sex", "smoker", "region"]
         num pipeline = make pipeline(
             SimpleImputer(strategy="median"),
             StandardScaler())
         cat pipeline = make pipeline(
             SimpleImputer(strategy="most_frequent"),
             OneHotEncoder(handle_unknown="ignore"))
         preprocessing = ColumnTransformer([
             ("num", num_pipeline, num_attribs),
("cat", cat_pipeline, cat_attribs)])
         medical_prepared = preprocessing.fit_transform(medical)
         print(medical_prepared.shape)
         print(preprocessing.get_feature_names_out())
         ['num_age' 'num_bmi' 'num_children' 'cat_sex_female' 'cat_sex_male'
          'cat smoker_no' 'cat smoker_yes' 'cat region_northeast'
'cat region_northwest' 'cat region_southeast' 'cat region_southwest']
```

## Select and Train a Model

At last! You framed the problem, you got the data and explored it, you sampled a training set and a test set, and you wrote a preprocessing pipeline to automatically clean up and prepare your data for machine learning algorithms. You are now ready to select and train a machine learning model.

## Train and Evaluate on the Training Set

The good news is that thanks to all these previous steps, things are now going to be easy! You decide to train a very basic linear regression model to get started:

```
In [ ]:
        #create a pipeline for preprocessing the data and fitting a linear regression model
        from sklearn.linear_model import LinearRegression
        lin reg = make pipeline(preprocessing, LinearRegression())
        # housing labels is the column we want to predict
        # uncomment the following line to fit the model
        lin_reg.fit(medical, medical_labels)
                          Pipeline
Out[]:
          ▶ columntransformer: ColumnTransformer
                    num
                                       cat
             ▶ SimpleImputer
                                ▶ SimpleImputer
             ▶ StandardScaler
                                ▶ OneHotEncoder
                     ▶ LinearRegression
```

Try out the model on the training set, look at the first five predictions and compare them to the labels:

```
from sklearn.metrics import mean_squared_error
lin_rmse = mean_squared_error(medical_labels, medical_predictions, squared=False)
lin_rmse
Out[]: 6202.558946589081
```

Now try <code>DecisionTreeRegressor</code>, as this is a fairly powerful model capable of finding complex nonlinear relationships in the data (decision trees are covered later in the course):

```
In []: # use DecisionTreeRegressor to train the model
    # use the make_pipeline() function to create a pipeline for preprocessing and model training
    # use the preprocessing object you created earlier
    # make predictions on the training set and compute the RMSE
    from sklearn.tree import DecisionTreeRegressor

    tree_reg = make_pipeline(preprocessing, DecisionTreeRegressor())
    tree_reg.fit(medical, medical_labels)

medical_pred = tree_reg.predict(medical)
    tree_rmse = mean_squared_error(medical_labels, medical_pred, squared=False)
    tree_rmse
```

Out[]: 270.2373612509658

# Better Evaluation Using Cross-Validation

The following code randomly splits the training set into 10 nonoverlapping subsets called folds, then it trains and evaluates the decision tree model 10 times, picking a different fold for evaluation every time and using the other 9 folds for training. The result is an array containing the 10 evaluation scores:

**Warning**. Scikit-Learn's cross-validation features expect a utility function (greater is better) rather than a cost function (lower is better), so the scoring function is actually the opposite of the RMSE. It's a negative value, so you need to switch the sign of the output to get the RMSE scores.

```
In [ ]: # uncomment the following line to compute the mean of the RMSEs

np.mean(tree_rmses)

0503.700353798569
```

Let's try one last model now: the RandomForestRegressor. As you will see later in the course, random forests work by training many decision trees on random subsets of the features, then averaging out their predictions.

### Randomized Search for Good Hyperparameters

```
In [ ]: # uncomment the following lines to search for the best hyperparameters
         from sklearn.model selection import RandomizedSearchCV
         from scipy.stats import randint
         from sklearn.pipeline import Pipeline
         1)
        param distribs = {'random forest max features': randint(low=2,high=20)}
         rnd_search = RandomizedSearchCV(full_pipeline,
                                           param distributions=param distribs,
                                           n iter=10,
                                           cv=3.
                                           scoring='neg_root_mean_squared_error',
                                           random state=42)
         rnd search.fit(medical, medical_labels)
                     RandomizedSearchCV
Out[]:
                    estimator: Pipeline
           ▶ preprocessing: ColumnTransformer
                    num
                                        cat
            ▶ SimpleImputer
                                ▶ SimpleImputer
            ▶ StandardScaler
                                ▶ OneHotEncoder
                 ▶ RandomForestRegressor
In [ ]: # uncomment the following lines to print the best search scores
         rn_res = pd.DataFrame(rnd_search.cv_results_)
         rn res.sort values(by="mean test score", ascending=False, inplace=True)
         rn res.head(5)["mean test score"]
             -4819.331482
Out[]:
            -4868.115355
        4
            -4868.115355
            -4884.079869
            -4892.429905
        Name: mean_test_score, dtype: float64
In [ ]: # uncomment the following lines to print the feature importances
         final model = rnd search.best estimator # includes preprocessing
         feature_importances = final_model["random_forest"].feature_importances_
         feature_importances
        array([0.1415512 , 0.18312249, 0.02236506, 0.00438671, 0.00459264,
Out[]:
                0.27840571, 0.34567506, 0.00534023, 0.00571929, 0.00520326,
                0.00363835])
In [ ]: # uncomment the following line to print the feature importances with the feature names
         sorted(zip(feature_importances, final_model["preprocessing"].get_feature_names_out()),reverse=True)
        [(0.3456750561140971, 'cat__smoker_yes'), (0.2784057087071407, 'cat__smoker_no'),
         (0.18312249286698468, 'num_bmi'),
(0.14155120446244887, 'num_age'),
         (0.022365059991817766, 'num__children'),
(0.005719289257578255, 'cat__region_northwest'),
(0.005340233009953545, 'cat__region_northeast'),
         (0.005203256938419487, 'cat__region_southeast'), (0.0045926399374515234, 'cat__sex_male'),
          (0.00438671058172246, 'cat__sex_female'),
         (0.0036383481323855964, 'cat region southwest')]
In [ ]: # now that you have a final model, evaluate it on the test set (find rmse)
         x_test = test_set.drop("charges", axis=1)
        y test = test set["charges"].copy()
         predictions = final model.predict(x test)
         final rmse = mean squared error(y test, predictions, squared=False)
         final_rmse
```

Out[]: 4482.4544777295905

Loading [MathJax]/jax/output/CommonHTML/fonts/TeX/fontdata.js