Housing_Kaggle

August 30, 2023

1 Predicting Housing Prices - Kaggle

In this notebook, I walk through the process of predicting housing prices in Kaggle's housing data competition. I prepare the data using various transformers. After processing the data, I use multiple algorithms and a bayesian optimizer to find the hyperparameters that minimze the error. Let's get started!

1.1 Libraries and Initial Set-Up

```
[1]: # General libraries
     from math import ceil
     import pandas as pd
     import numpy as np
     import random as ran
     # Plats
     import matplotlib.pyplot as plt
     # Processing
     from sklearn.preprocessing import OrdinalEncoder, OneHotEncoder, StandardScaler
     from sklearn.impute import KNNImputer
     from sklearn.pipeline import Pipeline
     from sklearn.compose import ColumnTransformer
     from sklearn.feature selection import SelectKBest, mutual info regression
     from sklearn.model_selection import cross_val_score, train_test_split
     # Optimizer
     from bayes_opt import BayesianOptimization
     # ML Models
     from sklearn.linear_model import Ridge, Lasso, ElasticNet
     from sklearn.svm import SVR
     from sklearn.ensemble import RandomForestRegressor as RFR
     from xgboost import XGBRegressor
     from sklearn.neighbors import KNeighborsRegressor
     from tensorflow.keras import Sequential
     from tensorflow.keras.layers import Dense, BatchNormalization, Dropout
     from tensorflow.keras import optimizers
```

```
from scikeras.wrappers import KerasRegressor
from tensorflow.keras.callbacks import EarlyStopping
```

Above, I import all the necessary packages. I import general libraries necessary to begin the script. I then import libraries to plot, process the data, score the models, and the necessary algorithms.

Now, I'll do a little set up by pre-setting the seed, the number of crossfolds, importing the training data, finding missing values, and creating a comparison matrix. The comparison matrix will hold the name of the model, it's root mean squared error, and the best hyperparameters from the optimizer. The data frame allows me to compare model performance across each algorithm.

```
[2]: # Random draw of seed for random state #
#seed = int(ran.uniform(1, 9999))
''' Got 2095

# Set mnumber of cross folds #
cv = 5

# Import training data #
train = pd.read_csv('train.csv')

# Split into X and Y #
X = train.drop(['SalePrice', 'Id'], axis = 1)
y = pd.DataFrame(train['SalePrice'])

# Find missing values #
missing = X.isnull().sum().sort_values(ascending = False)

# Make matrix to compare models #
train_compare = pd.DataFrame(columns = ['Model', 'RMSE', 'hypers'])
```

1.2 Initial Alteration of the Data Frame

The data frame has many missing values due to not having the feature that is being measured. For example, the frame has a variable call 'PoolQC' that measures the quality of the swiming pool. If the home does not have a pool, then it it labeled as missing. To avoid missing values in the frame, missing values were altered to fit the variable of interest.

```
[3]: # Assign no pool (NP) to PoolQC #
X['PoolQC'].fillna('NP', inplace = True)
```

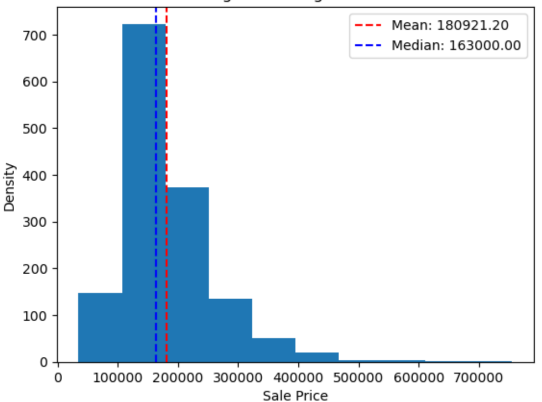
```
# Assign no feature (NF) to MiscFeature #
X['MiscFeature'].fillna('NF', inplace = True)
# Assign no ally (NAL) to Alley #
X['Alley'].fillna('NAL', inplace = True)
# Assign no fence (NF) to Fence #
X['Fence'].fillna('NF', inplace = True)
# Assign no fire place (NFP) to FireplaceQu #
X['FireplaceQu'].fillna('NFP', inplace = True)
# Assign no garage (NG) to GarageType #
X['GarageType'].fillna('NG', inplace = True)
# Fill garage variables with NG if no garage #
garage = ['GarageYrBlt', 'GarageFinish', 'GarageQual', 'GarageCond']
for x in garage:
   X[x].fillna('NG', inplace = True)
del x, garage
# Fix GarageYrBlt since it was mixed type #
X['GarageYrBlt'] = X['GarageYrBlt'].astype(str)
# Fill basement varaibles with no basement (NB) #
basement = ['BsmtExposure', 'BsmtFinType2', 'BsmtQual', 'BsmtCond', |
for x in basement:
   X[x].fillna('NB', inplace = True)
del x, basement
```

1.3 Plot Target Variable

After initial set-up, I'll plot the target variable to check for non-normality.

```
[4]: # Plot of Y #
plt.hist(y)
mean_value = y.mean()
median_value = y.median()
plt.axvline(mean_value.item(), color='red', linestyle='--', label='Mean')
```

Histogram of Target Variable



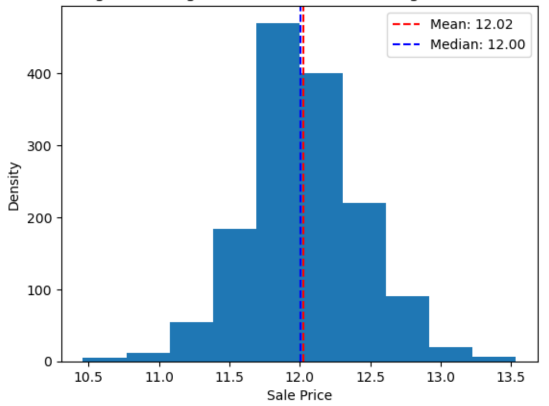
The plot above demonstrates a right skew in the target due to the right tail. In addition, there is distance between the mean and median values of the variable. I'll transform the variable with a natural log function and re-plot the data to ensure normality.

```
[5]: # Natural log the target variable #
    y_log = np.log(y)
    plt.hist(y_log)
    mean_value = y_log.mean()
    median_value = y_log.median()
    plt.axvline(mean_value.item(), color='red', linestyle='--', label='Mean')
    plt.axvline(median_value.item(), color='blue', linestyle='--', label='Median')
    plt.xlabel('Sale Price')
    plt.ylabel('Density')
    plt.title('Histogram of Target Variable with Natural Log Transformation')
    # Add the text to the legend
    mean_legend = plt.Line2D([], [], color='red', linestyle='--', label=f"Mean:__

√{mean_value.item():.2f}")

    median_legend = plt.Line2D([], [], color='blue', linestyle='--', label=f"Median:
     plt.legend(handles=[mean_legend, median_legend])
    plt.show()
    del mean_legend, mean_value, median_legend, median_value
```

Histogram of Target Variable with Natural Log Transformation



After applying the transformation, the target variable is now noramlly distributed.

1.4 Preprocessing

I'll now process the data to ensure is compatible for ML models. I'll split the data into training and validation data, which will be analyzed at the end. After, I encode my categorical variables and implement a KNN imputer. Last, I make a few variables into integer values.

```
[6]: # Split into train and validation sets #
     X_train, X_val, y_train, y_val = train_test_split(X, y_log, test_size = 0.25,
                                                       random_state = seed)
     # Mask of numerical features #
     numeric_feats = X_train.select_dtypes(include = ['int64', 'float64']).columns
     # Mask categorical features #
     cat_feats = X_train.select_dtypes(include = ['object']).columns
     # Encode object variables #
     ord_enc = OrdinalEncoder(handle_unknown = 'use_encoded_value', unknown_value=np.
      ⇒nan)
     X_train[cat_feats] = ord_enc.fit_transform(X_train[cat_feats])
     # Transform X_val with encoder #
     X_val[cat_feats] = ord_enc.transform(X_val[cat_feats])
     # KNN Imputer #
     knn_im = KNNImputer(n_neighbors = 10, weights = 'distance')
     X_train_imp = pd.DataFrame(knn_im.fit_transform(X_train), columns = X_train.
      ⇔columns)
     # KNN Imputer for X_val #
     X_val_imp = pd.DataFrame(knn_im.transform(X_val), columns = X_val.columns)
     # Variables need to be made integer #
     X_train_imp[['Electrical', 'MasVnrType', 'GarageYrBlt', 'Exterior1st']] =

      AX_train_imp[['Electrical', 'MasVnrType', 'GarageYrBlt', 'Exterior1st']]\
                                             .apply(lambda x: x.apply(ceil))
     X_val_imp[['Electrical', 'MasVnrType', 'GarageYrBlt', 'Exterior1st']] =

      →X_val_imp[['Electrical', 'MasVnrType', 'GarageYrBlt', 'Exterior1st']]\
```

```
.apply(lambda x: x.apply(ceil))
```

1.5 Feature Selection

The housing data includes numerous features, and many features are likely to be uncorrelated with the outcome. I limit the number of numeric features with a correlation threshold and the categorical features based on mutual information.

```
[7]: # Correlation of numeric feats and target #
     corr_val = pd.concat([X_train_imp[numeric_feats], y_train], axis =1).
      ⇔corr()['SalePrice']
     # Mask of features with high correlation to target #
     select_num = list(corr_val[(corr_val >= 0.50) | (corr_val <= -0.50)].index)</pre>
     select_num.remove('SalePrice')
     # Get mutual information of categorical features and outcome #
     fs = SelectKBest(score_func = mutual_info_regression, k = 'all')
     fs.fit(X_train_imp[cat_feats], np.ravel(y_train))
     # Sort the scores and corresponding feature names in descending order
     sorted_scores, sorted_features = zip(*sorted(zip(fs.scores_,
                                                       X_train_imp[cat_feats].
      ⇔columns),
                                                  reverse=True))
     # Select the top 20 features
     top_20_cat_features = list(sorted_features[:20])
     # Combine feature selection list #
     selected_feats = select_num + top_20_cat_features
     # DF of only selected features #
     X train imp = X train imp[selected feats]
     X_val_imp = X_val_imp[selected_feats]
     del corr val, fs, sorted features, sorted scores
```

1.6 Transformation

I now transform my data through categorical and numeric transformers based on the features in the limited data frame. I fit the transformer on the training data and then transform the training and validation data sets.

```
#### Transformation ####
    ########################
     # Categorical transformer #
    cat_trans = Pipeline(steps = [
         ('encode', OneHotEncoder(sparse_output = False))])
     # Numeric tranformer #
    num trans = Pipeline(steps = [
        ('stand', StandardScaler())])
    # Construct processor #
    processor = ColumnTransformer(
        transformers = [
             ('num', num_trans, select_num),
             ('cat', cat_trans, top_20_cat_features)],
        remainder = 'passthrough')
     # Apply processor to training data #
    temp = processor.fit_transform(X_train_imp)
     # Get categorical feature names #
    enc_cat_features = list(processor.named_transformers_['cat']['encode']\
                            .get_feature_names_out())
    # Concat label names #
    labels = select_num + enc_cat_features
    # Make df of processed data #
    X_train = pd.DataFrame(temp, columns = labels)
    del temp
     # Apply processor to validation data #
    temp = processor.transform(X_val_imp)
    # Get categorical feature names #
    enc_cat_features = list(processor.named_transformers_['cat']['encode']\
                            .get_feature_names_out())
```

```
# Concat label names #
labels = select_num + enc_cat_features

# Make df of processed data #
X_val = pd.DataFrame(temp, columns = labels)
del temp, y_log, X_val_imp, X_train_imp
```

2 Start Estimating

After preparing the training and validation datasets, the next few sections outline my strategies for predicting the outcome of interest. I'll use seven different estimation strategies that include Ridge, Lasso, Elastic Net, Support Vector Machine, Random Forest, XGBoost, KNN, and a Neural Net in their regression forms. I use a bayesian optimizer to search for the model with the best predictive power.

2.1 Ridge

```
[9]: # Define objective for ridge #
     def obj_ridge(alpha, fit_intercept, solver):
         Objective function to minimize the error of the
         ridge regression.
         Parameters
         alpha: L2 Regularization term.
             Regularizes the coefficients. Values stipulated
             in phounds.
         fit_intercept : Boolean of fit intercept.
             Indicator of whether or not the model
             fits an intercept.
         solver: Solving method of ridge regression.
             Continuous variable for selecting the best
             solver for the regression.
         Returns
         _____
         error: Mean squared error.
             Cross validation returns root mean error that is later
             convereted into RMSE in the comparison frame.
         HHHH
```

```
# Fit intercept #
    fit_intercept = bool(round(fit_intercept))
    # Solver #
    if solver <= 1.0:</pre>
        solver = 'auto'
    elif solver <= 2.0:</pre>
        solver = 'svd'
    elif solver <= 3.0:</pre>
        solver = 'cholesky'
    elif solver <= 4.0:</pre>
       solver = 'lsgr'
    elif solver <= 5.0:</pre>
        solver = 'sparse_cg'
    elif solver <= 6.0:</pre>
        solver = 'sag'
    else:
        solver = 'saga'
    # Instantiate ridge model #
    model = Ridge(alpha=alpha, fit_intercept=fit_intercept, solver=solver,
                   max_iter = 20000, random_state=seed)
    # Cross validation and mean MSE #
    error = cross_val_score(model, X_train, y_train, cv=cv,
                             scoring='neg_mean_squared_error').mean()
    # Return error #
    return error
# Define search space #
pbounds = {
    'alpha': (0.00000001, 100),
    'fit_intercept': (0, 1),
    'solver': (0, 8),
}
# Set the optimizer #
optimizer = BayesianOptimization(
    f=obj_ridge, pbounds=pbounds, random_state=seed,
    verbose = 0)
# Call maximizer #
optimizer.maximize(init_points=50, n_iter=450)
```

```
# Pull best info #
best_hypers = optimizer.max['params']
best_mse = optimizer.max['target']
# Replace solver with string #
if best_hypers['solver'] <= 1.0:</pre>
    best_hypers['solver'] = 'auto'
elif best_hypers['solver'] <= 2.0:</pre>
    best_hypers['solver'] = 'svd'
elif best_hypers['solver'] <= 3.0:</pre>
    best_hypers['solver'] = 'cholesky'
elif best_hypers['solver'] <= 4.0:</pre>
    best_hypers['solver'] = 'lsqr'
elif best_hypers['solver'] <= 5.0:</pre>
    best_hypers['solver'] = 'sparse_cg'
elif best_hypers['solver'] <= 6.0:</pre>
    best_hypers['solver'] = 'sag'
else:
    best_hypers['solver'] = 'saga'
# Fill comparison matrix #
train_compare = pd.concat([train_compare,
                            pd.DataFrame({'Model' : 'Ridge',
                             'RMSE': np.sqrt(best_mse * -1),
                             'hypers': [best_hypers]})], ignore_index = True)
# Sort by smallest RMSE #
train_compare = train_compare.sort_values('RMSE')
```

2.2 Lasso

```
fit_intercept : Boolean of fit intercept.
        Indicator of whether or not the model
        fits an intercept.
    selection : Dictates coefficient updates.
        Continuous variable of using either cycle or
        random for coefficient update.
    Returns
    _____
    error: Mean squared error.
        Cross validation returns root mean error that is later
        convereted into RMSE in the comparison frame.
    11 11 11
    # Fit intercept #
    fit_intercept = bool(round(fit_intercept))
    # selection #
    if selection <= 0.5:
        selection = 'cyclic'
    else:
        selection = 'random'
    # Instantiate model #
    model = Lasso(alpha = alpha, fit_intercept = fit_intercept,
                  selection = selection,
                  random_state = seed, max_iter = 20000)
    # Cross validation and mean MSE #
    error = cross_val_score(model, X_train, y_train, cv=cv,
                            scoring='neg_mean_squared_error').mean()
    # Return error #
    return error
# Define search space #
pbounds = {
    'alpha': (0.0000001, 100),
    'fit_intercept': (0, 1),
    'selection': (0, 1)
}
# Set the optimizer #
optimizer = BayesianOptimization(
```

```
f=obj_lasso, pbounds=pbounds, random_state=seed,
    verbose = 0)
# Call maximizer #
optimizer.maximize(init_points = 50, n_iter = 450)
# Pull best info #
best hypers = optimizer.max['params']
best_mse = optimizer.max['target']
# Replace selection with string #
if best_hypers['selection'] <= 0.5:</pre>
    best_hypers['selection'] = 'cyclic'
else:
    best_hypers['selection'] = 'random'
# Fill comparison matrix #
train_compare = pd.concat([train_compare,
                           pd.DataFrame({'Model' : 'Lasso',
                             'RMSE': np.sqrt(best_mse * -1),
                             'hypers': [best_hypers]})], ignore_index = True)
train_compare = train_compare.sort_values('RMSE')
/Users/treywood/opt/anaconda3/lib/python3.10/site-
packages/sklearn/linear_model/_coordinate_descent.py:631: ConvergenceWarning:
Objective did not converge. You might want to increase the number of iterations,
check the scale of the features or consider increasing regularisation. Duality
gap: 3.199e+00, tolerance: 1.399e-02
 model = cd_fast.enet_coordinate_descent(
/Users/treywood/opt/anaconda3/lib/python3.10/site-
packages/sklearn/linear_model/_coordinate_descent.py:631: ConvergenceWarning:
Objective did not converge. You might want to increase the number of iterations,
check the scale of the features or consider increasing regularisation. Duality
gap: 3.078e+00, tolerance: 1.440e-02
 model = cd_fast.enet_coordinate_descent(
/Users/treywood/opt/anaconda3/lib/python3.10/site-
packages/sklearn/linear_model/_coordinate_descent.py:631: ConvergenceWarning:
Objective did not converge. You might want to increase the number of iterations,
check the scale of the features or consider increasing regularisation. Duality
gap: 3.468e+00, tolerance: 1.408e-02
 model = cd_fast.enet_coordinate_descent(
/Users/treywood/opt/anaconda3/lib/python3.10/site-
packages/sklearn/linear_model/_coordinate_descent.py:631: ConvergenceWarning:
Objective did not converge. You might want to increase the number of iterations,
```

```
check the scale of the features or consider increasing regularisation. Duality
gap: 2.884e+00, tolerance: 1.348e-02
   model = cd_fast.enet_coordinate_descent(
/Users/treywood/opt/anaconda3/lib/python3.10/site-
packages/sklearn/linear_model/_coordinate_descent.py:631: ConvergenceWarning:
Objective did not converge. You might want to increase the number of iterations,
check the scale of the features or consider increasing regularisation. Duality
gap: 2.813e+00, tolerance: 1.389e-02
   model = cd_fast.enet_coordinate_descent(
```

2.3 Elastic Net

```
[]: # Define objective function for Net #
     def obj_net(alpha, l1_ratio, fit_intercept,
                 selection):
         .....
         The objective of this function is to minimize the error
         of the elastic net model.
         Parameters
         alpha: Float
             Constant the multiplies the penalty terms. O is equal to OLS.
         l1 ratio : Float
             Ratio of 11 or 12 regularization. O is 12. 1 is 11.
         fit_intercept : bool
             Option to fit an intercept.
         selection : String
             Specify how coefficients are updated across iterations.
         Returns
         _____
         error : Float
             Cross validation returns root mean error that is later
             convereted into RMSE in the comparison frame.
         11 11 11
         # Vary fit intercept #
         fit_intercept = bool(round(fit_intercept))
         # Vary selection #
         if selection <= 0.5:</pre>
             selection = 'cyclic'
             selection = 'random'
```

```
# Instantiate the model #
    model = ElasticNet(alpha = alpha, l1_ratio = l1_ratio,
                       fit_intercept = fit_intercept,
                       selection = selection, random_state = seed,
                       max_iter = 20000)
    # Cross validation and mean MSE #
    error = cross_val_score(model, X_train, np.ravel(y_train), cv=cv,
                            scoring='neg mean squared error').mean()
    # Return error #
    return error
# Define search space #
pbounds = {
    'alpha': (0.00001, 100),
    'l1_ratio': (0.001, 0.99),
    'fit_intercept': (0, 1),
    'selection': (0, 1)
}
# Set the optimizer #
optimizer = BayesianOptimization(
    f=obj_net, pbounds=pbounds, random_state=seed,
    verbose = 0)
# Call maximizer #
optimizer.maximize(init_points = 50, n_iter = 450)
# Pull best info #
best_hypers = optimizer.max['params']
best_mse = optimizer.max['target']
# Replace selection with string #
if best hypers['selection'] <= 0.5:</pre>
   best_hypers['selection'] = 'cyclic'
else:
    best_hypers['selection'] = 'random'
# Fill comparison matrix #
train_compare = pd.concat([train_compare,
                           pd.DataFrame({'Model' : 'Elastic_Net',
```

```
'RMSE': np.sqrt(best_mse * -1),
                             'hypers': [best_hypers]})], ignore_index = True)
train_compare = train_compare.sort_values('RMSE')
/Users/treywood/opt/anaconda3/lib/python3.10/site-
packages/sklearn/linear_model/_coordinate_descent.py:631: ConvergenceWarning:
Objective did not converge. You might want to increase the number of iterations,
check the scale of the features or consider increasing regularisation. Duality
gap: 8.741e+00, tolerance: 1.399e-02
 model = cd_fast.enet_coordinate_descent(
/Users/treywood/opt/anaconda3/lib/python3.10/site-
packages/sklearn/linear_model/_coordinate_descent.py:631: ConvergenceWarning:
Objective did not converge. You might want to increase the number of iterations,
check the scale of the features or consider increasing regularisation. Duality
gap: 8.823e+00, tolerance: 1.440e-02
 model = cd_fast.enet_coordinate_descent(
/Users/treywood/opt/anaconda3/lib/python3.10/site-
packages/sklearn/linear_model/_coordinate_descent.py:631: ConvergenceWarning:
Objective did not converge. You might want to increase the number of iterations,
check the scale of the features or consider increasing regularisation. Duality
gap: 8.958e+00, tolerance: 1.408e-02
 model = cd_fast.enet_coordinate_descent(
/Users/treywood/opt/anaconda3/lib/python3.10/site-
packages/sklearn/linear_model/_coordinate_descent.py:631: ConvergenceWarning:
Objective did not converge. You might want to increase the number of iterations,
check the scale of the features or consider increasing regularisation. Duality
gap: 7.971e+00, tolerance: 1.348e-02
 model = cd_fast.enet_coordinate_descent(
/Users/treywood/opt/anaconda3/lib/python3.10/site-
packages/sklearn/linear_model/_coordinate_descent.py:631: ConvergenceWarning:
Objective did not converge. You might want to increase the number of iterations,
check the scale of the features or consider increasing regularisation. Duality
gap: 8.942e+00, tolerance: 1.389e-02
 model = cd_fast.enet_coordinate_descent(
/Users/treywood/opt/anaconda3/lib/python3.10/site-
packages/sklearn/linear_model/_coordinate_descent.py:631: ConvergenceWarning:
Objective did not converge. You might want to increase the number of iterations,
check the scale of the features or consider increasing regularisation. Duality
gap: 8.315e+00, tolerance: 1.399e-02
 model = cd_fast.enet_coordinate_descent(
/Users/treywood/opt/anaconda3/lib/python3.10/site-
packages/sklearn/linear_model/_coordinate_descent.py:631: ConvergenceWarning:
Objective did not converge. You might want to increase the number of iterations,
check the scale of the features or consider increasing regularisation. Duality
gap: 8.664e+00, tolerance: 1.440e-02
 model = cd_fast.enet_coordinate_descent(
/Users/treywood/opt/anaconda3/lib/python3.10/site-
```

```
packages/sklearn/linear_model/_coordinate_descent.py:631: ConvergenceWarning:
Objective did not converge. You might want to increase the number of iterations,
check the scale of the features or consider increasing regularisation. Duality
gap: 8.603e+00, tolerance: 1.408e-02
 model = cd fast.enet coordinate descent(
/Users/treywood/opt/anaconda3/lib/python3.10/site-
packages/sklearn/linear_model/_coordinate_descent.py:631: ConvergenceWarning:
Objective did not converge. You might want to increase the number of iterations,
check the scale of the features or consider increasing regularisation. Duality
gap: 7.799e+00, tolerance: 1.348e-02
 model = cd_fast.enet_coordinate_descent(
/Users/treywood/opt/anaconda3/lib/python3.10/site-
packages/sklearn/linear_model/_coordinate_descent.py:631: ConvergenceWarning:
Objective did not converge. You might want to increase the number of iterations,
check the scale of the features or consider increasing regularisation. Duality
gap: 8.533e+00, tolerance: 1.389e-02
 model = cd_fast.enet_coordinate_descent(
```

2.4 Support Vector Machine

```
[]: # Define objective function for SVM #
     def obj_SVR(kernel, degree,
                 gamma, C, epsilon,
                 shrinking):
         The objective of this function is to minimze the erro
         of the support vector regression.
         Parameters
         _____
         kernel: Kernel used in solver.
             String inputs that are used in optimizer.
         degree : Degree of polyomial kernel.
            Only used in poly alogrithm.
         gamma : Kernel cofficient.
             Only used in rbf, poly, and sigmoid.
         C: L2 regularizer.
             More regularization at smaller values.
         epsilon: Epplison value in SVR model.
             Specifies penalty in training loss function.
         shrinking : Boolean value.
             Dictates if the model uses shrinking heuristic.
         Returns
         error : Mean squared error.
             Cross validation returns root mean error that is later
```

```
convereted into RMSE in the comparison frame.
    n n n
    # Kernel #
    if kernel <= 1:</pre>
        kernel = 'linear'
    elif kernel <= 2:</pre>
        kernel = 'poly'
    elif kernel <= 3:</pre>
        kernel = 'rbf'
    else:
        kernel = 'sigmoid'
    # Gamma #
    if gamma <= 0.5:
        gamma = 'scale'
        gamma = 'auto'
    # Shrinking #
    shrinking = bool(round(shrinking))
    # Instantiate SVR #
    model = SVR(kernel = kernel, degree = int(degree),
                gamma = gamma, C = C,
                epsilon = epsilon, shrinking = shrinking,
                max_iter = 50000)
    # Cross validation and mean MSE #
    error = cross_val_score(model, X_train, np.ravel(y_train), cv=cv,
                             scoring='neg_mean_squared_error').mean()
    # Return error #
    return error
# Define search space #
pbounds = {
    'kernel': (0, 4),
    'degree': (1, 10),
    'gamma': (0, 1),
    'C': (0.0001, 100),
    'epsilon': (0.0001, 100),
    'shrinking': (0, 1)
}
```

```
# Set the optimizer #
optimizer = BayesianOptimization(
    f=obj_SVR, pbounds=pbounds, random_state=seed,
    verbose = 0)
# Call maximizer #
optimizer.maximize(init_points = 50, n_iter = 450)
# Pull best info #
best_hypers = optimizer.max['params']
best_mse = optimizer.max['target']
# Replace kernel with string #
if best_hypers['kernel'] <= 1:</pre>
    best_hypers['kernel'] = 'linear'
elif best_hypers['kernel'] <= 2:</pre>
    best_hypers['kernel'] = 'poly'
elif best_hypers['kernel'] <= 3:</pre>
    best_hypers['kernel'] = 'rbf'
else:
    best_hypers['kernel'] = 'sigmoid'
# Replace gamma with string #
if best_hypers['gamma'] <= 0.5:</pre>
    best_hypers['gamma'] = 'scale'
else:
    best_hypers['gamma'] = 'auto'
# Fill comparison matrix #
train_compare = pd.concat([train_compare,
                            pd.DataFrame({'Model' : 'SVR',
                             'RMSE': np.sqrt(best_mse * -1),
                             'hypers': [best_hypers]})], ignore_index = True)
train_compare = train_compare.sort_values('RMSE')
```

2.5 Random Forest

```
[]: # Define objective function for random forest #
     def obj_RF(n_estimators, criterion,
                min_samples_split, min_samples_leaf,
                max_features, bootstrap, min_impurity_decrease):
         11 11 11
         Parameters
         n estimators : Float
             Number of trees to estimate in the forest.
         criterion : String
             How the tree measures quality of the split.
         min\_samples\_split : Float
             Minimum number of samples required to split a node.
         min\_samples\_leaf : Float
             Minimum number of samples required to be a leaf.
         max_features : String
             Number of features to consider when splitting.
         bootstrap : Boolean
             Whether bootstraps are used when building trees.
         min_impurity_decrease : Float
             Node is split if it decreases the impurity.
         Returns
         error: Mean squared error.
             Cross validation returns root mean error that is later
             convereted into RMSE in the comparison frame.
         11 11 11
         # Criterion #
         if criterion <= 1.0:</pre>
             criterion = 'squared_error'
         elif criterion <= 2.0:</pre>
             criterion = 'absolute_error'
         elif criterion <= 3.0:</pre>
             criterion = 'friedman_mse'
         else:
             criterion = 'poisson'
         # Max features #
         if max_features <= 0.5:</pre>
             max_features = 'sqrt'
         else:
```

```
max_features = 'log2'
    # Bootstrap #
   bootstrap = bool(round(bootstrap))
    # instantiate random forest moel #
   model = RFR(n_estimators = int(n_estimators), criterion = criterion,
                min_samples_split = min_samples_split,
                min_samples_leaf = min_samples_leaf,
                max_features = max_features, bootstrap = bootstrap,
                min_impurity_decrease = min_impurity_decrease,
                n_jobs = -1, random_state = seed)
    # Cross validation and mean MSE #
   error = cross_val_score(model, X_train, np.ravel(y_train), cv=cv,
                            scoring='neg_mean_squared_error').mean()
    # Return error #
   return error
# Define search space #
pbounds = {
    'n estimators': (1, 1000),
    'criterion': (0, 4),
    'min_samples_split': (0.01, .70),
    'min_samples_leaf': (0.01, .70),
    'max_features': (0, 1),
    'bootstrap': (0, 1),
    'min_impurity_decrease': (0.001, 0.4)
}
# Set the optimizer #
optimizer = BayesianOptimization(
   f=obj_RF, pbounds=pbounds, random_state=seed,
   verbose = 0)
# Call maximizer #
optimizer.maximize(init_points = 50, n_iter = 450,)
# Pull best info #
best_hypers = optimizer.max['params']
best_mse = optimizer.max['target']
```

```
# Replace criterion with string #
if best_hypers['criterion'] <= 1.0:</pre>
    best_hypers['criterion'] = 'squared_error'
elif best_hypers['criterion'] <= 2.0:</pre>
    best_hypers['criterion'] = 'absolute_error'
elif best_hypers['criterion'] <= 3.0:</pre>
    best_hypers['criterion'] = 'friedman_mse'
else:
    best_hypers['criterion'] = 'poisson'
# Replace max features with string #
if best_hypers['max_features'] <= 0.5:</pre>
    best_hypers['max_features'] = 'sqrt'
else:
    best_hypers['max_features'] = 'log2'
# Fill comparison matrix #
train_compare = pd.concat([train_compare,
                            pd.DataFrame({'Model' : 'Random_Forest',
                             'RMSE': np.sqrt(best_mse * -1),
                             'hypers': [best_hypers]})], ignore_index = True)
train_compare = train_compare.sort_values('RMSE')
```

2.6 XGBoost

```
subsample : Float
        Subsample of the dataset to use in tree.
    colsample_bytree : Float
        Subsample of columns to use in each tree.
    reg\_lambda : Float
        L2 regularization on weights. Higher values make models more
 \hookrightarrow conservative.
    alpha: Float
        L1 regularization on weights. Higher values make models more ⊔
 \hookrightarrow conservative.
    Returns
    error : Float
        Cross validation returns root mean error that is later
        convereted into RMSE in the comparison frame.
    HHHH
    # instantiate XGBoost #
    model = XGBRegressor(n_estimators = int(n_estimators), eta = eta,
                          gamma = gamma, max_depth = int(max_depth),
                          subsample = subsample, colsample_bytree = __
 ⇔colsample_bytree,
                          reg_lambda = reg_lambda, alpha = alpha,
                          seed = seed, n_jobs = 1)
    # Cross validation and mean MSE #
    error = cross_val_score(model, X_train, np.ravel(y_train), cv=cv,
                             scoring='neg_mean_squared_error').mean()
    # Return error #
    return error
# Define the search space #
pbounds = {
    'n estimators': (1, 2000),
    'eta': (0, 1),
    'gamma': (0, 5),
    'max_depth': (2, 7),
    'subsample': (0.5, 1),
    'colsample_bytree': (0.2, 0.9),
    'reg_lambda': (0.05, 10),
    'alpha': (0.05, 10)
}
```

2.7 KNN

```
[]: # Define objective function for K-Nearest Neighbors #
     def obj_knn(n_neighbors, weights, algorithm,
                 leaf_size, p):
         11 11 11
         This objective function minimzes the error for k-nearest
         neighbors regression.
         Parameters
         n_neighbors: int
             Number of neighbors to use.
         weights : String
             Weight function used in prediction.
         algorithm : String
             Process used to compute the nearest neighbors.
         leaf_size : int
             Leaf size passed to specific algorithms.
         p:int
             Power parameter for Minkowski metric.
         Returns
```

```
error : Float
        Cross validation returns root mean error that is later
        convereted into RMSE in the comparison frame.
    11 11 11
    # Variation on weights #
    if weights <= 0.5:</pre>
        weights = 'uniform'
    else:
        weights = 'distance'
    # Variation on algorithm #
    if algorithm <= 1.0:</pre>
        algorithm = 'auto'
    elif algorithm <= 2.0:</pre>
        algorithm = 'ball_tree'
    elif algorithm <= 3.0:</pre>
        algorithm = 'kd_tree'
    else:
        algorithm = 'brute'
    # Variation on p #
    if p <= 1.0:</pre>
        p = 1
    elif p <= 1.0 and algorithm != 'brute':</pre>
        p = 1
    else:
        p = 2
    # Instantiate model #
    model = KNeighborsRegressor(n_neighbors = int(n_neighbors), weights = ___
 ⇔weights,
                                  algorithm = algorithm, leaf_size =
 →int(leaf_size), p = p)
    # Cross validation and mean MSE #
    error = cross_val_score(model, X_train, np.ravel(y_train), cv=cv,
                              scoring='neg_mean_squared_error').mean()
    # Return error #
    return error
# Define search space #
pbounds = {
```

```
'n_neighbors': (2, 10),
    'weights': (0, 1),
    'algorithm': (0, 4),
    'leaf_size': (2, 50),
    'p': (0.001, 2)
}
# Set the optimizer #
optimizer = BayesianOptimization(
    f=obj knn, pbounds=pbounds, random state=seed,
    verbose = 0)
# Call maximizer #
optimizer.maximize(init_points = 50, n_iter = 450)
# Pull best info #
best_hypers = optimizer.max['params']
best_mse = optimizer.max['target']
# Replace weights with string #
if best_hypers['weights'] <= 0.5:</pre>
    best_hypers['weights'] = 'uniform'
else:
    best_hypers['weights'] = 'distance'
# Replace algorithm with string #
if best_hypers['algorithm'] <= 1.0:</pre>
    best_hypers['algorithm'] = 'auto'
elif best_hypers['algorithm'] <= 2.0:</pre>
    best_hypers['algorithm'] = 'ball_tree'
elif best_hypers['algorithm'] <= 3.0:</pre>
    best_hypers['algorithm'] = 'kd_tree'
else:
    best_hypers['algorithm'] = 'brute'
# Replace p #
if best_hypers['p'] <= 1.0:</pre>
    best_hypers['p'] = 1
elif best_hypers['p'] <= 1.0 and best_hypers['algorithm'] != 'brute':</pre>
    best_hypers['p'] = 1
else:
```

2.8 Neural Network

```
[]: # Define objective function for network #
     def obj_net(batch_size, epochs, activation, num_nodes,
                 num_hidden_layers, learning_rate, rate, optimizer):
         .....
         The objective of this function is to minimize the error of the
         neural network
         Parameters
         _____
         batch\_size : Int
             The number of cases to include in each batch.
         epochs : Int
             Number of runs through the data when updating weights.
         activation : String
             Type of activation function for the layer.
         num_nodes : Int
             Number of nodes to include in the hidden layer.
         num_hidden_layers : Int
             Number of hideen layers in the model.
         learning_rate : Float
             How much to change the model with each model update.
         rate : Float
             Dropout rate for each hidden layer to prevent overfitting.
         optimizer : String
             Optimizer to use for the model.
         Returns
         _____
         error : Float
             Cross validation returns root mean error that is later
             convereted into RMSE in the comparison frame.
         11 11 11
```

```
# Set Optimizer #
if optimizer <= 0.33:</pre>
    optimizer = optimizers.Adam(learning_rate = learning_rate)
elif optimizer <= 0.66:</pre>
    optimizer = optimizers.Adagrad(learning_rate = learning_rate)
else:
    optimizer = optimizers.RMSprop(learning_rate = learning_rate)
# Set activation function #
if activation <= 0.33:</pre>
    activation = 'relu'
elif activation <= 0.66:</pre>
   activation = 'sigmoid'
else:
   activation = 'tanh'
# Instantiate model
model = Sequential()
# Set input layer #
model.add(Dense(int(num_nodes), activation = activation,
                input_shape = (X_train.shape[1],)))
# Set hidden layer with batch normalizer #
for _ in range(int(num_hidden_layers)):
    model.add(Dense(int(num_nodes), activation = activation))
    model.add(BatchNormalization())
    model.add(Dropout(rate = rate, seed = seed))
# Add output layer #
model.add(Dense(1))
# Set compiler #
model.compile(optimizer = optimizer,
              loss = 'mean_squared_error')
# Set early stopping #
early_stopping = EarlyStopping(monitor='val_loss',
                                patience=15,
                                restore_best_weights=True)
# Create model #
```

```
reg = KerasRegressor(model = lambda : model,
                        batch_size = int(batch_size),
                         epochs = int(epochs),
                        validation_split = 0.2,
                        callbacks = [early_stopping],
                        random_state = seed)
    # Cross validation and mean MSE #
   error = cross_val_score(reg, X_train, np.ravel(y_train), cv=cv,
                       scoring='neg_mean_squared_error').mean()
    # Return error #
   return error
# Define search space #
pbounds = {
    'batch_size': (50, 1460),
    'epochs': (5, 500),
    'learning_rate': (0.001, 0.15),
    'num_nodes': (5, 80),
    'num_hidden_layers': (2, 20),
    'activation': (0, 1),
    'rate': (0.0, 0.9),
    'optimizer': (0, 1)
}
# Set the optimizer #
optimizer = BayesianOptimization(f=obj_net, pbounds=pbounds,
                                random_state=seed,
                                verbose = 0)
# Call the maximizer #
optimizer.maximize(init_points=50, n_iter=450)
# Pull best info #
best_hypers = optimizer.max['params']
best_mse = optimizer.max['target']
# Replace optimizer and learning rate #
if best_hypers['optimizer'] <= 0.33:</pre>
   best_hypers['optimizer'] = optimizers.Adam(learning_rate =__
```

```
elif best_hypers['optimizer'] <= 0.66:</pre>
   best_hypers['optimizer'] = optimizers.Adagrad(learning_rate =_
 ⇔best_hypers['learning_rate'])
else:
   ⇔best_hypers['learning_rate'])
# Replace activation with string #
if best_hypers['activation'] <= 0.33:</pre>
   best hypers['activation'] = 'relu'
elif best_hypers['activation'] <= 0.66:</pre>
     best_hypers['activation'] = 'sigmoid'
else:
    best_hypers['activation'] = 'tanh'
# Fill comparison matrix #
train_compare = pd.concat([train_compare,
                         pd.DataFrame({'Model' : 'Neural_Net',
                          'RMSE': np.sqrt(best_mse * -1),
                          'hypers': [best_hypers]})], ignore_index = True)
train_compare = train_compare.sort_values('RMSE')
```

2.9 Best Models

After creating and optimizing the models, the script filled the train_compare matrix with the best models and their associated errors.

```
[]: display(train_compare)
```

2.10 Predict on the Validation Set

After training each model, I now predict on the validation set to see which model does the best job.

```
random_state = seed)
mod_ridge.fit(X_train, y_train)
# Lasso #
lasso_dict = train_compare.loc[train_compare['Model'] == 'Lasso', 'hypers'].
 yalues[0]
mod_lasso = Lasso(alpha = lasso_dict['alpha'],
                  fit_intercept = bool(lasso_dict['fit_intercept']),
                  selection = lasso_dict['selection'],
                  random_state = seed,
                  max_iter = 20000)
mod_lasso.fit(X_train, y_train)
# Elastic Net #
elastic_dict = train_compare.loc[train_compare['Model'] == 'Elastic_Net',u

    'hypers'].values[0]

mod_elastic = ElasticNet(alpha = elastic_dict['alpha'],
                         fit_intercept = bool(elastic_dict['fit_intercept']),
                         11_ratio = elastic_dict['l1_ratio'],
                         selection = elastic_dict['selection'],
                         \max iter = 20000,
                         random_state = seed)
mod_elastic.fit(X_train, y_train)
# SVR #
svr_dict = train_compare.loc[train_compare['Model'] == 'SVR', 'hypers'].
 yalues[0]
mod_SVR = SVR(C = svr_dict['C'],
              degree = int(svr_dict['degree']),
              epsilon = svr_dict['epsilon'],
              gamma = svr_dict['gamma'],
              kernel = svr_dict['kernel'],
              shrinking = bool(svr_dict['kernel']))
mod_SVR.fit(X_train, y_train)
# Random Forest #
```

```
rf_dict = train_compare.loc[train_compare['Model'] == 'Random_Forest',__
 mod_rf = RFR(bootstrap = bool(rf_dict['bootstrap']),
            criterion = rf_dict['criterion'],
            max features = rf dict['max features'],
            min_impurity_decrease = rf_dict['min_impurity_decrease'],
            min_samples_leaf = rf_dict['min_samples_leaf'],
            min_samples_split = rf_dict['min_samples_split'],
            n_estimators = int(rf_dict['n_estimators']))
mod_rf.fit(X_train, np.ravel(y_train))
# XGBoost Regression #
boost_dict = train_compare.loc[train_compare['Model'] == 'XGBoost_Reg',__
mod_boost = XGBRegressor(alpha = boost_dict['alpha'],
                        colsample_bytree = boost_dict['colsample_bytree'],
                        eta = boost_dict['eta'],
                        gamma = boost_dict['gamma'],
                        max_depth = int(boost_dict['max_depth']),
                        n_estimators = int(boost_dict['n_estimators']),
                        reg_lambda = boost_dict['reg_lambda'],
                        subsample = boost_dict['subsample'],
                        random_state = seed,
                        n jobs = 3)
mod_boost.fit(X_train, np.ravel(y_train))
# K-Nearest #
knn dict = train compare.loc[train compare['Model'] == 'KNN Reg', 'hypers'].
 ovalues[0]
mod_KNN = KNeighborsRegressor(algorithm = knn_dict['algorithm'],
                             leaf size = int(knn dict['leaf size']),
                             n_neighbors = int(knn_dict['n_neighbors']),
                             p = float(knn dict['p']),
                             weights = knn_dict['weights'])
mod_KNN.fit(X_train, y_train)
# Neutral Net #
```

```
net_dict = train_compare.loc[train_compare['Model'] == 'Neural_Net', 'hypers'].
 ⇔values[0]
mod_net = Sequential()
mod_net.add(Dense(int(net_dict['num_nodes']),
                  activation = net_dict['activation'],
                  input_shape = (X_train.shape[1],)))
# Set hidden layer with batch normalizer #
for _ in range(int(net_dict['num_hidden_layers'])):
   mod_net.add(Dense(int(net_dict['num_nodes']), activation =__
 →net_dict['activation']))
   mod_net.add(BatchNormalization())
   mod_net.add(Dropout(rate = net_dict['rate'], seed = seed))
# Add output layer #
mod_net.add(Dense(1))
# Set compiler #
mod_net.compile(optimizer = net_dict['optimizer'],
              loss = 'mean_squared_error')
mod net.fit(X train, np.ravel(y train))
# Make list of models #
mod_list = [mod_ridge, mod_lasso, mod_elastic, mod_SVR, mod_rf, mod_boost,__
→mod_KNN, mod_net]
# Make matrix to compare models #
val_compare = pd.DataFrame(columns = ['Model', 'RMSE'])
# Loop model predictions on validation set #
for x in mod list:
   pred = x.predict(X_val)
   mse = mean_squared_error(y_val, pred)
   rmse = np.sqrt(mse)
   model_name = type(x).__name__
   val_compare = pd.concat([val_compare,
                             pd.DataFrame({'Model': [model_name],
                                            'RMSE': [np.exp(rmse)],
```

```
'Model_Specs': [x]})],
ignore_index = True)

# Sort by RMSE #
val_compare = val_compare.sort_values('RMSE')

# Display Dataframe #
display(val_compare)
```

2.11 Import and Prepare Test Data

I now prepare the test dataset with the transformers created earlier.

```
[]: # Import test data #
     X_test = pd.read_csv('test.csv')
     ids = X_test['Id']
     # Drop ID #
     X_test = X_test.drop("Id", axis = 1)
     # Assign no pool (NP) to PoolQC #
     X_test['PoolQC'].fillna('NP', inplace = True)
     # Assign no feature (NF) to MiscFeature #
     X_test['MiscFeature'].fillna('NF', inplace = True)
     # Assign no ally (NAL) to Alley #
     X_test['Alley'].fillna('NAL', inplace = True)
     # Assign no fence (NF) to Fence #
     X_test['Fence'].fillna('NF', inplace = True)
     # Assign no fire place (NFP) to FireplaceQu #
     X_test['FireplaceQu'].fillna('NFP', inplace = True)
     # Assign no garage (NG) to GarageType #
     X_test['GarageType'].fillna('NG', inplace = True)
```

```
# Fill garage variables with NG if no garage #
garage = ['GarageYrBlt', 'GarageFinish', 'GarageQual', 'GarageCond']
for x in garage:
   X_test[x].fillna('NG', inplace = True)
del x, garage
# Fix GarageYrBlt since it was mixed type #
X_test['GarageYrBlt'] =X_test['GarageYrBlt'].astype(str)
# Fill basement varaibles with no basement (NB) #
basement = ['BsmtExposure', 'BsmtFinType2', 'BsmtQual', 'BsmtCond', |
for x in basement:
   X_test[x].fillna('NB', inplace = True)
del x, basement
# Transform X test with encoder #
X_test[cat_feats] = ord_enc.transform(X_test[cat_feats])
# KNN Imputer for X val #
X_test = pd.DataFrame(knn_im.transform(X_test), columns = X_test.columns)
# Variables need to be made integer #
X_test[['Electrical', 'MasVnrType', 'GarageYrBlt', 'Exterior1st',
 X_test[['Electrical', 'MasVnrType', 'GarageYrBlt', 'Exterior1st', | ]
.apply(lambda x: x.apply(ceil))
# Get pre-selected features #
X_test = X_test[selected_feats]
# Apply processor to validation data #
temp = processor.transform(X_test)
# Get categorical feature names #
enc_cat_features = list(processor.named_transformers_['cat']['encode']\
                      .get_feature_names_out())
```

```
# Concat label names #
labels = select_num + enc_cat_features

# Make df of processed data #
X_test = pd.DataFrame(temp, columns = labels)
```

2.12 Predict on Test Set

To finish up the problem, I predict using the test features and the best model from the validation testing stage. I then store these predictions in a CSV file to submit to Kaggle.

```
[]: # Pull best model from val_compare #
best_model = val_compare.iloc[0]['Model_Specs']

# Predict on test set #
predict_test = best_model.predict(X_test)

# Exponentiate to make into real dollars #
predict_test = np.exp(predict_test)

# Create new DataFrame with the IDs and the predicted sale prices #
predictions_with_id = pd.DataFrame({'Id': ids, 'SalePrice': predict_test})

# Save the predictions to a CSV file with the original IDs
predictions_with_id.to_csv('predicted_sale_prices.csv', index=False)
```