```
In [10]: ## importing the required libraries
         import pandas as pd
         from sklearn.model selection import train test split
         from sklearn.metrics import mean absolute error
         from sklearn.impute import SimpleImputer
         from sklearn.tree import DecisionTreeRegressor
         from sklearn.ensemble import RandomForestRegressor
         from xgboost import XGBRegressor
In [11]: ## Load the data
         train data = pd.read csv('train.csv')
         ## Drop rows with missing target values
         train data.dropna(axis=0, subset=['SalePrice'], inplace=True)
         ## Define target (y) and features (X)
         v = train data['SalePrice']
         X = train data.drop(['SalePrice'], axis=1).select dtypes(exclude=['object'])
         ## Now, ensure v is aligned with X after dropping missing values
         y = y[X.index]
In [12]: ## Split the data into training and testing sets
         ## 25% of the data will be used for testing, the rest for training
         ## Feature (x) and target (v) splits
         train_X, test_X, train_y, test_y = train_test_split(X.values, y.values, test_size=0.25, random_state=42)
         ## Initialise the imputer (you can choose strategy: 'mean', 'median', or 'most frequent')
         my imputer = SimpleImputer() # Default is 'mean', but can choose others
         ## Fit and transform the training data (impute missing values in the training set)
         train X = my imputer.fit transform(train X)
         ## Transform the test data using the same imputer (to avoid data Leakage)
         test X = my imputer.transform(test X)
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## Optionally, you can convert back to DataFrame with the original column names after imputation
         train X = pd.DataFrame(train X, columns=X.columns)
         test X = pd.DataFrame(test X, columns=X.columns)
In [13]: ## Predictions using the Decision Tree algorithm
         decision model = DecisionTreeRegressor()
         decision model.fit(train X, train y)
         predicted decision trees = decision model.predict(test X)
         print ("Mean Absolute Error using Decision Tress :", mean absolute error(test y, predicted decision trees))
        Mean Absolute Error using Decision Tress: 24394.780821917808
In [14]: ## Predictions using the Random Forest algorithm
         forest model = RandomForestRegressor(n estimators=100, max depth=10)
         forest model.fit(train X, train y )
         predicted random forest = forest model.predict(test X)
         print("Mean Absolute Error using Random Forest:", mean_absolute_error(test_y, predicted_random_forest))
        Mean Absolute Error using Random Forest: 17481.22066004496
In [15]: ## Predictions using the XGBoost algorithm
         xg model = XGBRegressor(n estimators=100)
         xg model.fit(train X, train y)
         predicted XGBoost = xg model.predict(test X)
         print("Mean Absolute Error using XGBoost: ", mean_absolute_error(test_y, predicted_XGBoost))
        Mean Absolute Error using XGBoost: 18624.489811643834
In [16]: ## Let's improve the RandomForest Model as it was the best performing in MAE
         ## We will use GridSearchCV to tune the hyperparameters of the RandomForest Model
         from sklearn.model selection import GridSearchCV
         ## Define the hyperparameter grid to search
         param grid = {
             'n estimators': [100, 200, 300],
             'max_depth': [10, 20, 30, None],
             'min_samples_split': [2, 5, 10],
             'min_samples_leaf': [1, 2, 4],
```

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```
'max features': ['sqrt', 'log2']
         ## Create a RandomForestRegressor model
         rf model = RandomForestRegressor()
         ## Perform grid search with cross-validation
         grid search = GridSearchCV(estimator=rf model, param grid=param grid, cv=2, n jobs=-1, verbose=2)
         grid search.fit(train X, train y)
         ## Get the best parameters and best score
         print("Best parameters:", grid_search.best_params_)
         print("Best score:", grid search.best score )
         ## Predict with the best model
         best rf model = grid search.best estimator
         predicted random forest = best rf model.predict(test X)
         print("Mean Absolute Error using optimised Random Forest:", mean absolute error(test y, predicted random forest))
        Fitting 2 folds for each of 216 candidates, totalling 432 fits
        c:\Users\samuel.mcdonnell\AppData\Local\anaconda3\envs\HousePrices\Lib\site-packages\numpy\ma\core.py:2820: RuntimeWarning: inv
        alid value encountered in cast
          data = np.array(data, dtype=dtype, copy=copy,
        Best parameters: {'max depth': None, 'max features': 'sqrt', 'min samples leaf': 1, 'min samples split': 5, 'n estimators': 30
        Best score: 0.8350334919908053
        Mean Absolute Error using optimised Random Forest: 16769.477760700804
In [17]: ## Get the feature importance scores
         feature_importances = forest_model.feature_importances_
         ## Create a list of features with their corresponding importance scores
         feature importance dict = {
             feature: importance for feature, importance in zip(X.columns, feature importances)
         ## Sort the features by their importance score (in descending order)
         sorted features = sorted(feature importance dict.items(), key=lambda x: x[1], reverse=True)
         ## Display the top 10 most important features
```

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top 10 features = sorted features[:10]
 print("Top 10 Features by Importance:")
 for feature, importance in top 10 features:
     print(f"{feature}: {importance}")
 ## Random Forest inherently handles feature importance so it isn't necessary to include in tuning process. Still interesting t
Top 10 Features by Importance:
OverallQual: 0.5624306342744222
GrLivArea: 0.12098009777652968
TotalBsmtSF: 0.03809611172661414
2ndFlrSF: 0.03260436035765927
1stFlrSF: 0.030584864830592924
BsmtFinSF1: 0.029765704475803872
GarageCars: 0.0265242748256848
GarageArea: 0.022851367791071046
LotArea: 0.020380813784723593
YearBuilt: 0.015883262828223846
```

In [19]: mean_absolute_error(test_y, predicted_random_forest)/train_data['SalePrice'].median()

Out[19]: 0.10288023166074113

In conclusion, this Random Forest model, after hyperparameter tuning, achieved a Mean Absolute Error of 16,769. The median of the house price data is 163,000. This gives us a relative error of 10.3%, a great result considering the complexity of the data.