

A photograph of a modern building facade with dark, rectangular panels and large windows. The building has a geometric, stepped design. A dark blue diagonal overlay covers the right side of the image, where the title and author information are placed.

The Machine Learning Journey

By: Daniel Immediato

Date: 5/03/24

Preliminary: What is the Ames Dataset?

- Ames, Iowa is a college town of Iowa State University. The Ames dataset consists of the housing sale records between 2006-2010, including features like their area, number of rooms, and other attributes. The goal of this project was to create the most “accurate machine learning model with the best score”.
- 1. A machine learning model finds patterns from data. In our case, the data has 81 columns and 2580 rows. It would usually, then, make a prediction from said given data set.
- 2. A model’s “score” is a model evaluation, such as accuracy or R-squared, that explains the strength of the model.
- This can be achieved by:
 - 1. Reshaping the data into a workable state
 - 2. Picking and then creating the best models that work for you
 - 3. Tuning said models

Preprocessing: Data Types

- Before beginning any sort of analysis, one must examine whether or not the data “can” be analyzed properly to begin with. The typical dataset is split between numeric, categorical, and ordinal values. When coding, one has to ensure values are consistently classified throughout.
 - As a few examples, ‘MSSubClass’ is actually a category, not an integer, and dates, like ‘YearBuilt’ and ‘YearRemodAdd’, are also categories. PID is ultimately pointless as a feature when it comes to what we initially want to look at.
 - It is necessary to convert these to their proper data types, otherwise errors may occur or results may appear different than expected.
 - Variables like ‘OverallQual’ and ‘OverallCond’ must be manually encoded to a ‘least-to-greatest’ directory.

PID	int64
GrLivArea	int64
SalePrice	int64
MSSubClass	int64
MSZoning	object
LotFrontage	float64
LotArea	int64
Street	object
Alley	object
LotShape	object
LandContour	object
Utilities	object
LotConfig	object
LandSlope	object
Neighborhood	object
Condition1	object
Condition2	object
BldgType	object
HouseStyle	object
OverallQual	int64
OverallCond	int64
YearBuilt	int64
YearRemodAdd	int64

Preprocessing: Nulls and NA

- An NA value is likely to cause the most issues when dealing with data. When running functions, depending on the program, many can't process them and may produce an error or show results incorrectly. To deal with them you have several options, such as:
 1. Assume what the value was going to be based on related values.
 2. Replace values as a "0" or "none" if the value is legitimately lacking, or if there is no value.
 3. Replace values with most common value or median.

- Shown is an untransformed version of the "count" of all missing values in the dataset.

LotFrontage	462
Alley	2412
MasVnrType	1573
MasVnrArea	14
BsmtQual	69
BsmtCond	69
BsmtExposure	71
BsmtFinType1	69
BsmtFinSF1	1
BsmtFinType2	70
BsmtFinSF2	1
BsmtUnfSF	1
TotalBsmtSF	1
Electrical	1
BsmtFullBath	2
BsmtHalfBath	2
FireplaceQu	1241
GarageType	127
GarageYrBlt	129
GarageFinish	129
GarageCars	1
GarageArea	1
GarageQual	129
GarageCond	129
PoolQC	2571
Fence	2055
MiscFeature	2483





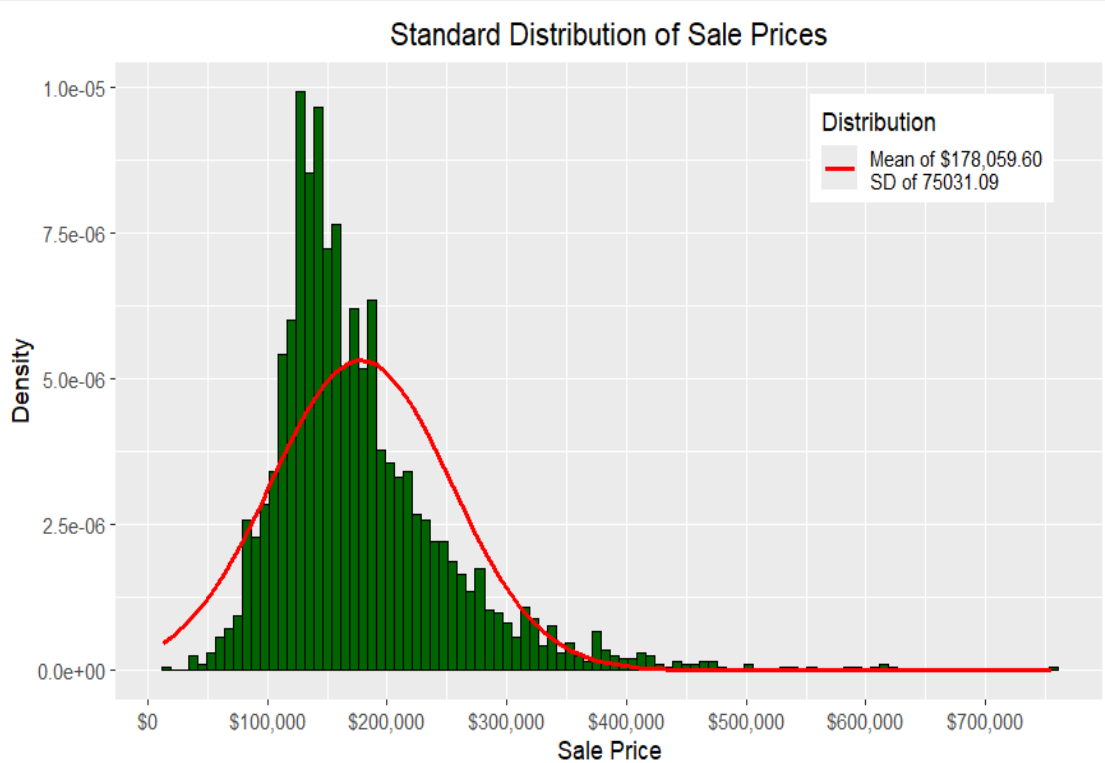
Preprocessing: The Pipeline

- The problem with preprocessing independently of each other is that it can lead not only to data leakage, but as the code becomes increasingly complex, so does the requirement for automation. Pipelines can be used to condense “heavier” data processing into more easily digestible, and, more importantly, recallable steps.
 - For our purposes, the basic pipeline consisted of: separating the data types (and classifying them appropriately), deal with the “NA” values using a SimpleImputer, scaling the numeric values, and then encoding the categorical and ordinal values. These can then be combined into a preprocessor using ColumnTransformer.

```
electrical_transformer = Pipeline(steps=[
    ('impute_electrical', SimpleImputer(strategy='most_frequent')),
    ('ordinal_electrical', OrdinalEncoder(categories=[ordinal_order['Electrical']])))
])
numeric_transformer = Pipeline(steps=[
    ('impute_mean', SimpleImputer(strategy='mean')),
    ('scaler', StandardScaler())
])
ordinal_transformer = Pipeline(steps=[
    ('impute_none', SimpleImputer(strategy='constant', fill_value='None')), # Using SimpleImputer
    ('ordinal', OrdinalEncoder(categories=[ordinal_order[feature] for feature in ordinal_except_electrical])))
])
nominal_features = [feature for feature in categorical_features if feature not in ordinal_features]
categorical_transformer = Pipeline(steps=[
    ('impute_none', SimpleImputer(strategy='constant', fill_value='None')), # Using SimpleImputer
    ('onehot', OneHotEncoder(handle_unknown='ignore'))
])
```

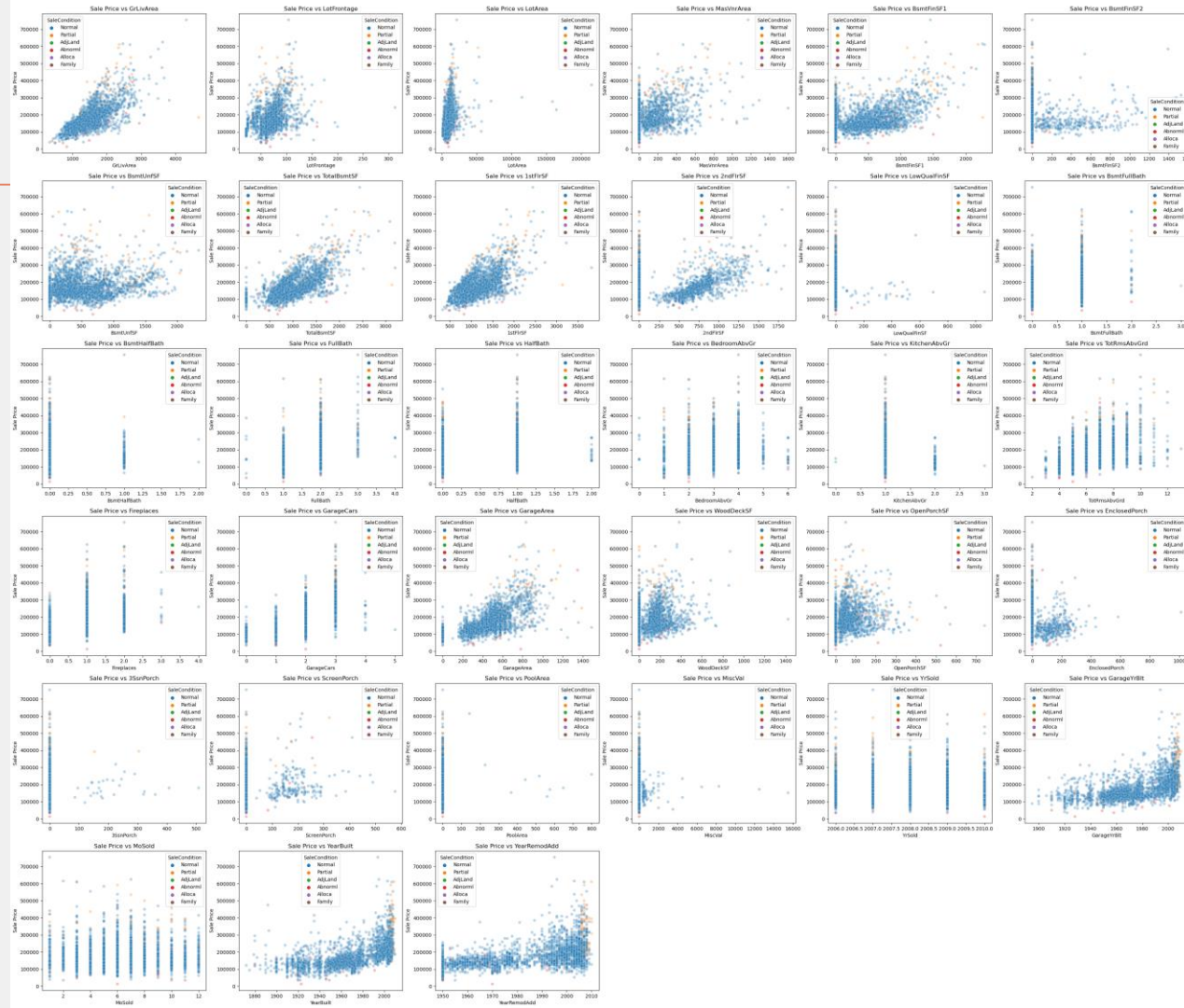
Overview: The Target

- Our target, the 'Y' value, is Sale Price. This is the value we are measuring, and eventually predicting. Using a standard distribution histogram, we can see that most houses have been sold at around \$180,000. This is fairly typical, as most people cannot afford more expensive houses.
- However, it should also be kept in mind that there are numerous outliers in the data because of this. In fact, one can even be seen here, with a house being sold at over \$700,000.

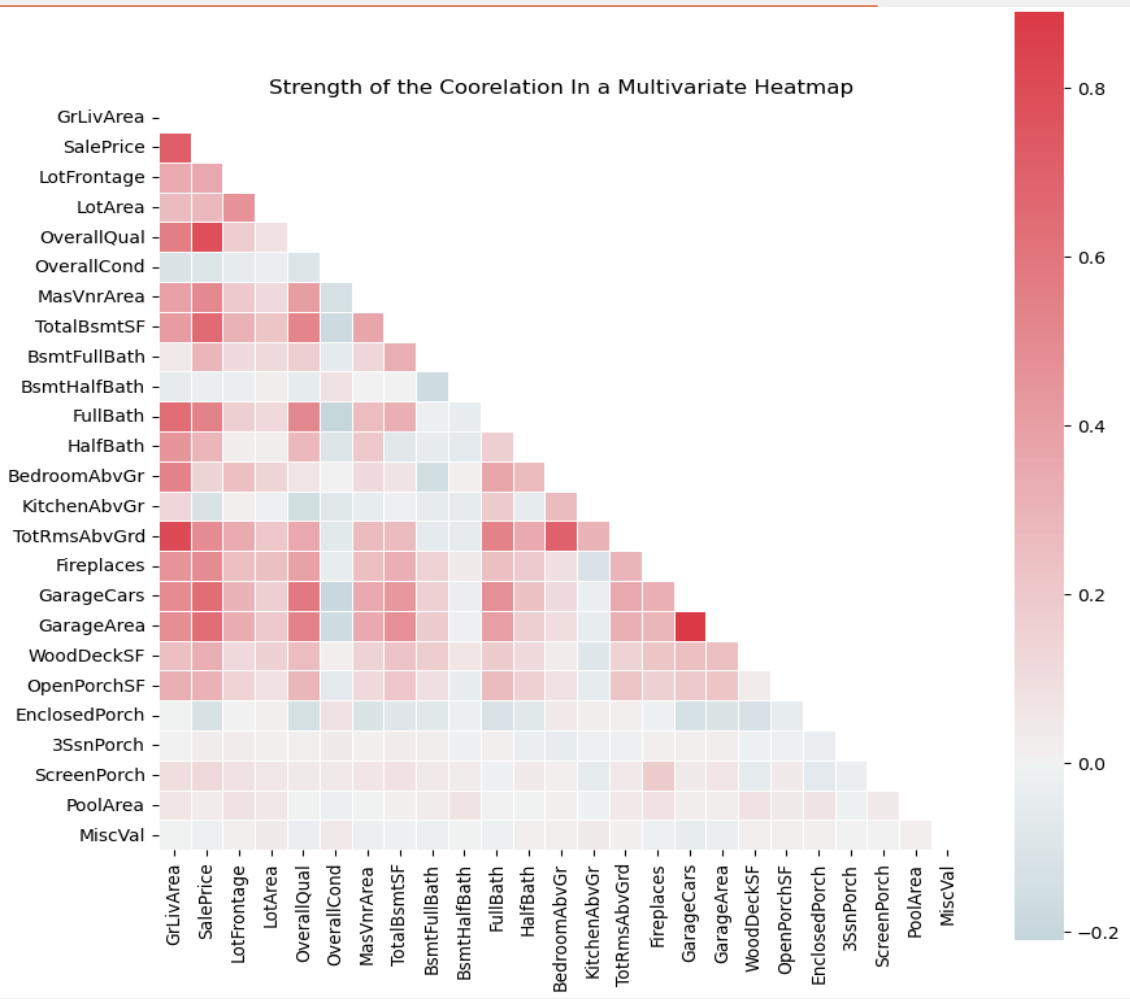


Overview: Numeric

- Here is every numeric variable tested against the target, which is SalePrice. We can see many of the variables have some sort of correlation just from eyeing them, and we can also tell which might satisfy the features of linear regression.
- We can also see which variables are ordinal judging by the spacing of the plot points.
- Correlations can be extracted from this.



Overview: Summarizing Coorelations



- To the left, we see the numeric and ordinal values (assuming it is ranked).
- We can see that overall quality, ground living area, garage area, and others have higher correlation, while values like pool area and the “MiscVal” feature are of weaker strength.
- This can be used to compare feature importance, as well as collinearity for a multiple linear regression model.



Multiple Linear Regression: As a Model

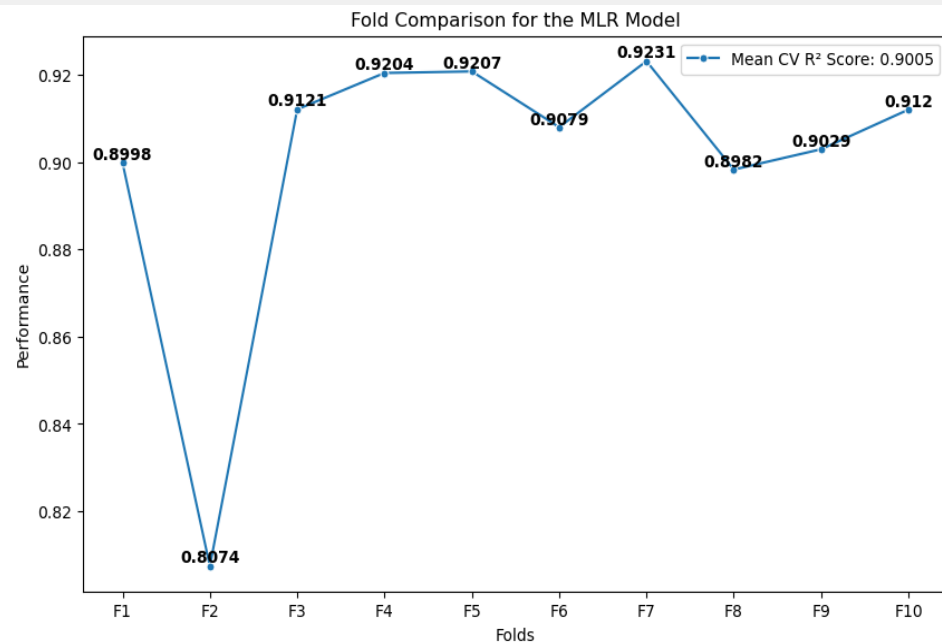
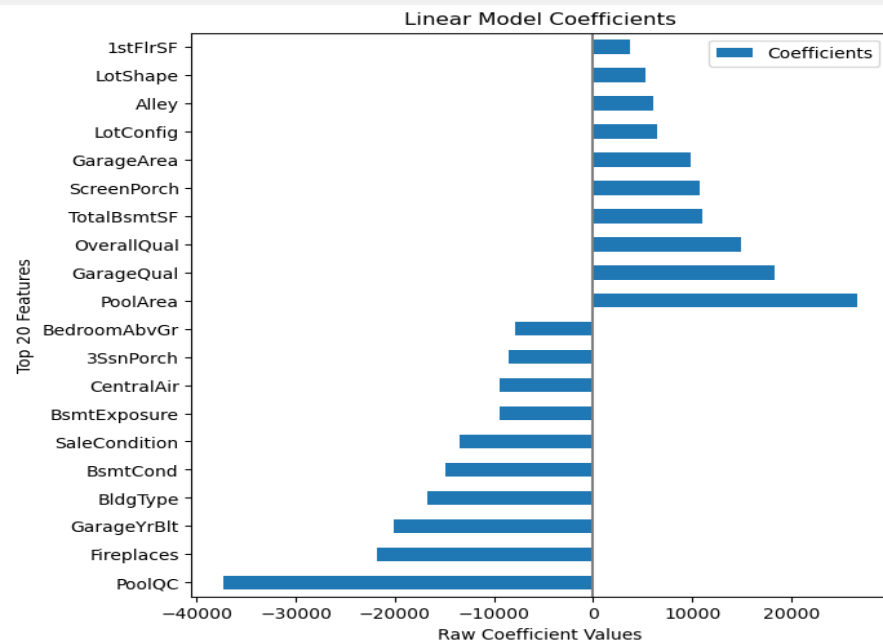
- A MLR is used to determine the strength of relationships, as well as statistical significance. When used in machine learning, it is using the more than one x value (independent variable) to predict the y value (dependent variable)
- This allows for the inclusion of several predictors, which an OLS (Ordinary Least Squares) model cannot do. This would, therefore, determine the significance of each predictor as well.
- One must ensure features are linear to the target, there is constant variance, normality of errors, there is independence of errors, and that there is as little multi-collinearity as possible.



MLR: The Transition From a Stats Model



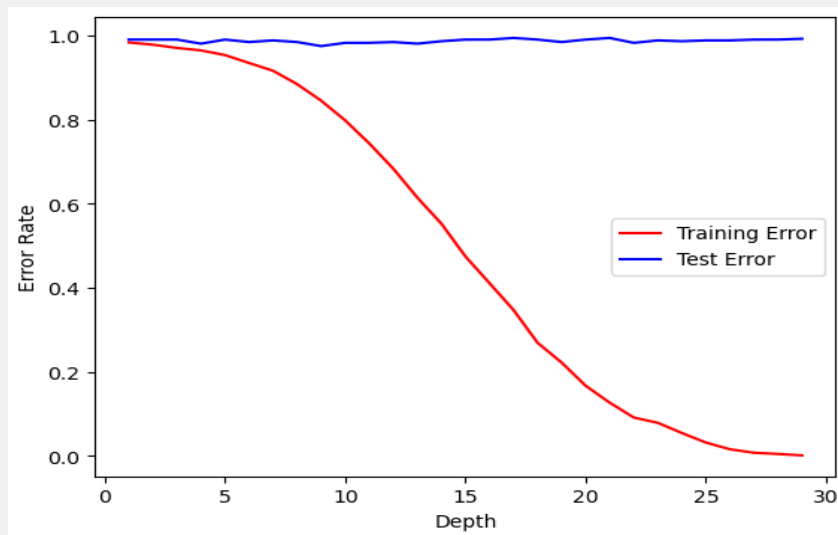
- An OLS model would typically give coefficients, standard error, and a P-value. The MLR mainly provides the intercepts and the coefficients, but is able to provide predictions at the cost of standard error.
- Cross-validation score is the performance of the model on unseen data. The data is split into groups, called folds, of hopefully equal size so that different models are being trained in order to improve accuracy and performance.



Regression Models: Lasso, Ridge, and Elastic Net

- Multiple-regression models are used as regularization techniques to deal with the problem of "overfitting", which occurs when the model is too closely adapted to the training data.
- The Lasso regression selects only the most relevant features, while the Ridge model reduces sensitivity to change by penalizing the "largest coefficients". Elastic Net is a hybrid of both.
- All have the goal of reducing model complexity, handling multicollinearity, and making predictions more explainable.

- Example of overfitting in a decision tree: High test error, but low training error. We want to prevent this.



Regressions: “Default” and “Tuned” Models

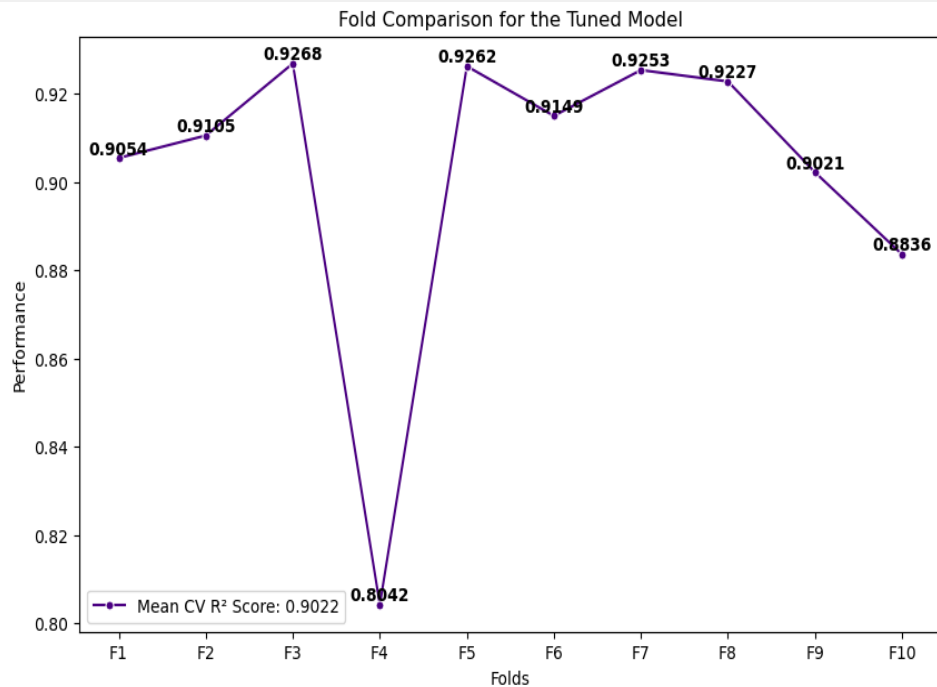
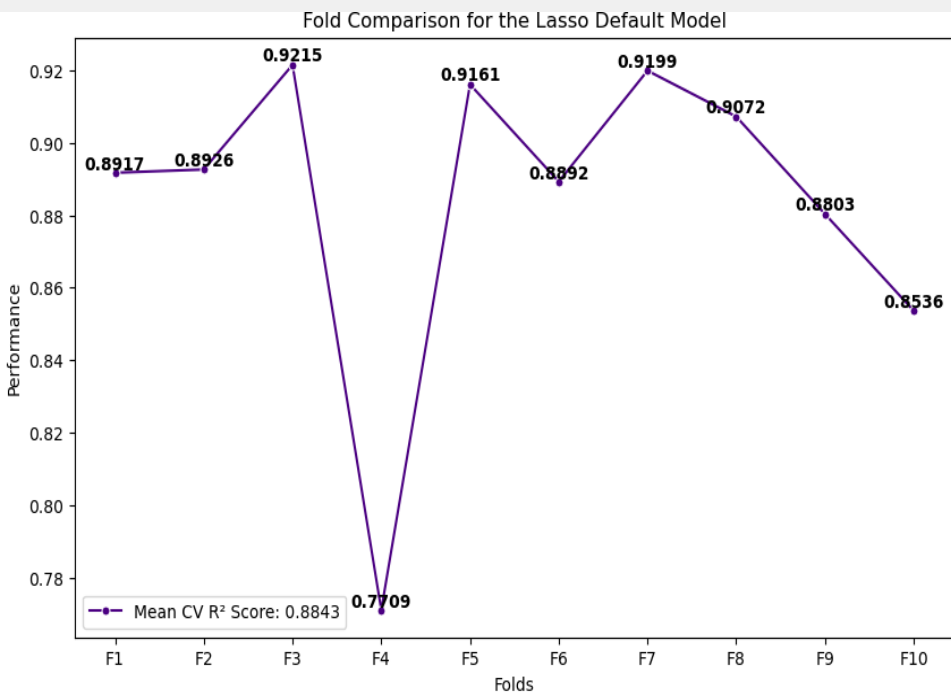
- The default model is the version of the model that has not been affected by parameter tuning, or anything related to it. It is the most basic cross-validation score one can get. In an ideal scenario, the default model is kept if it is superior in score to a tuned model.
- The tuned model is subject to parameter tuning, done through a grid search to find the best combination of "hyper-parameters" to find the highest accuracy. This is done manually with “ranges” of parameters, although for the regressors only the alphas were used.
- Below is an example of parameter tuning. Note the difference for elastic net, which corresponds directly to the lasso at 1, but then shrinks to the ridge as it approaches 0. Python will find the “best parameters” that lead to the “best score” out of these. In our case, it was 40 (and nothing below that), .1, and .9, respectively.

```
param_grids = {  
    'Lasso': {'regressor__alpha': [40, 50, 60]},  
    'Ridge': {'regressor__alpha': [40, 50, 60]},  
    'ElasticNet': {'regressor__alpha': [0.01, 0.1, 1, 10], 'regressor__l1_ratio': [0.8, .9, .95]}  
}
```

Regressions: The Lasso

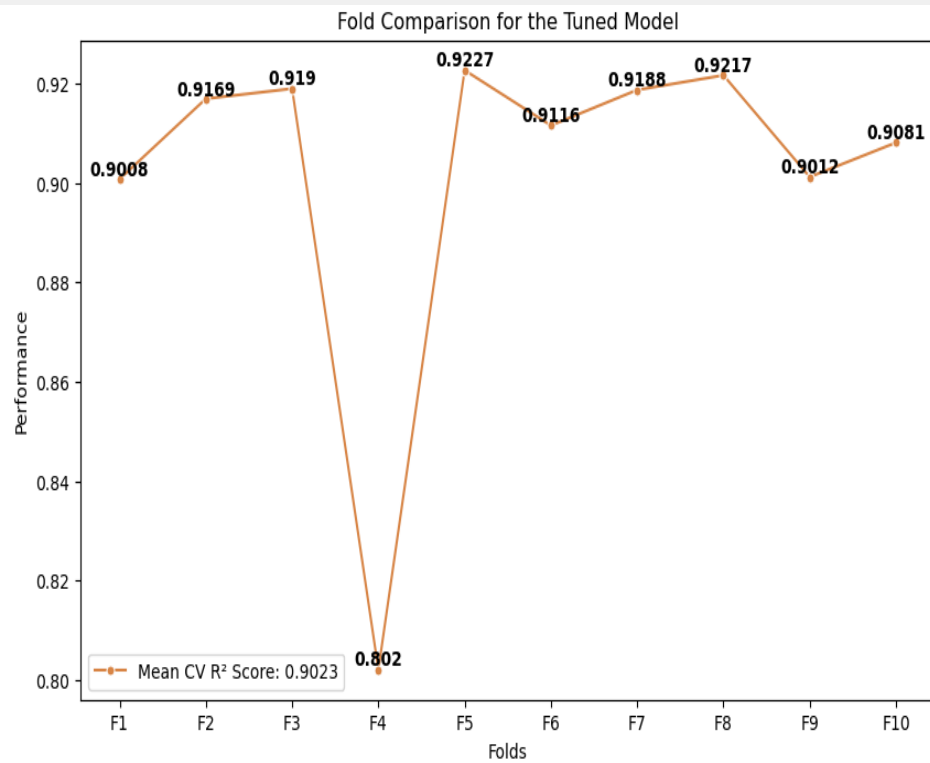
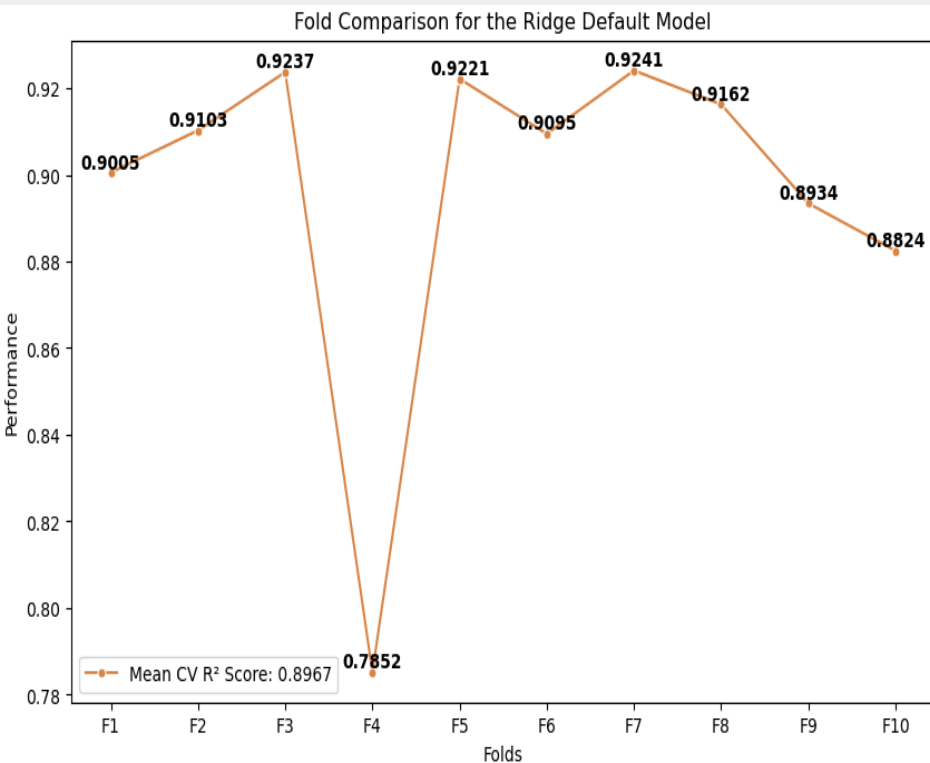


- The Lasso regression selects only the important features in a data set, while also reducing complexity. It is one of the fastest models out of the three, if not the fastest, and can handle larger number of features.
- The problem with the Lasso regression is that it is attempting to shrink the coefficient estimates to zero. This means that, because it will only retain one variable, a highly correlated model will end up having lower accuracy and a loss of information.



Regressions: The Ridge

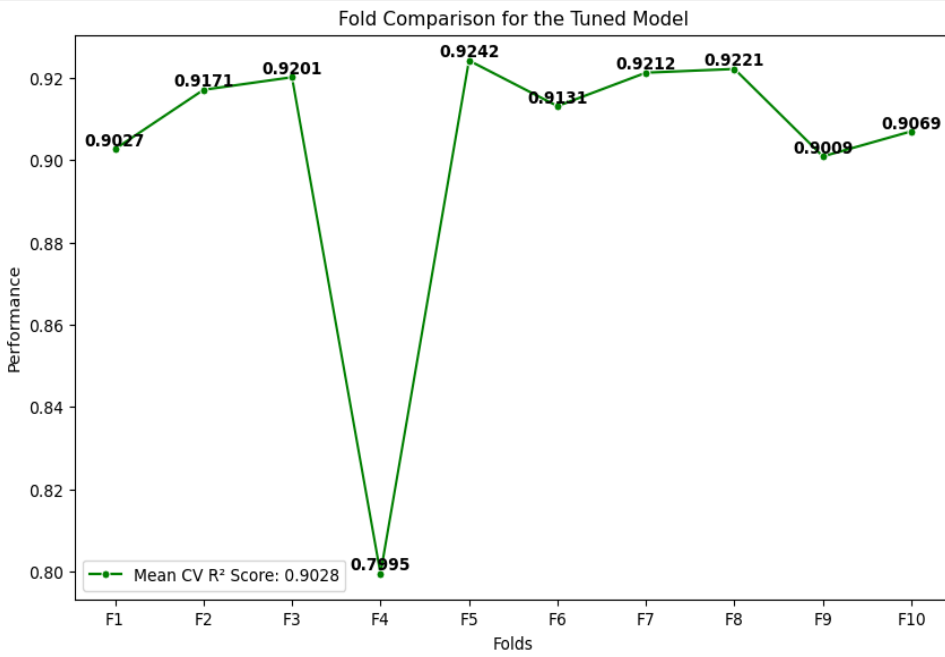
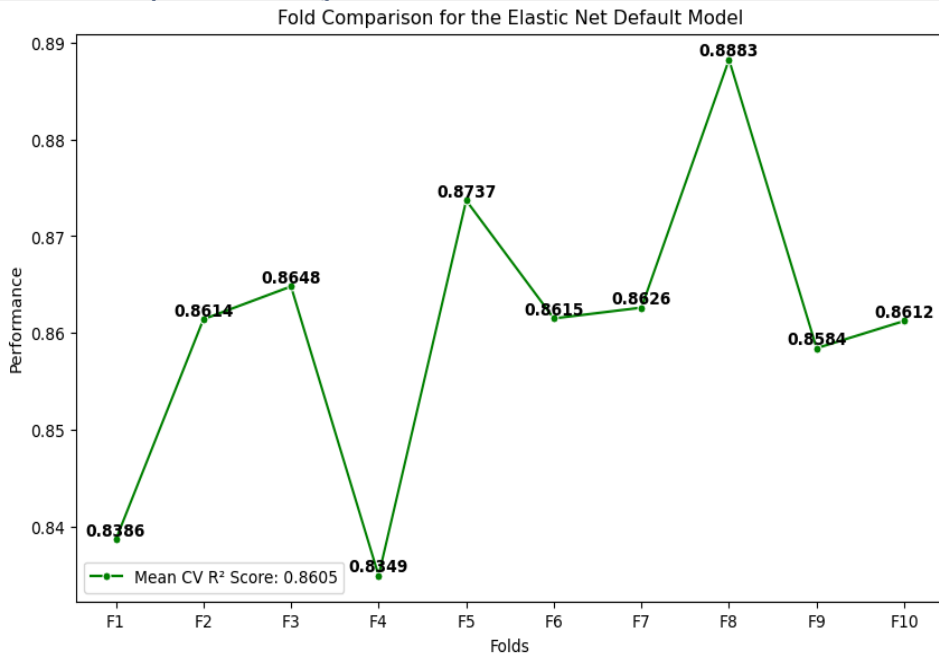
- Ridge regressions retain all the features in the model by only shrinking the coefficients, instead of setting them to zero like in the Lasso. This means there is no feature selection and less bias than the Lasso regression, at the cost of making the model more complex.



Regressions: Elastic Net



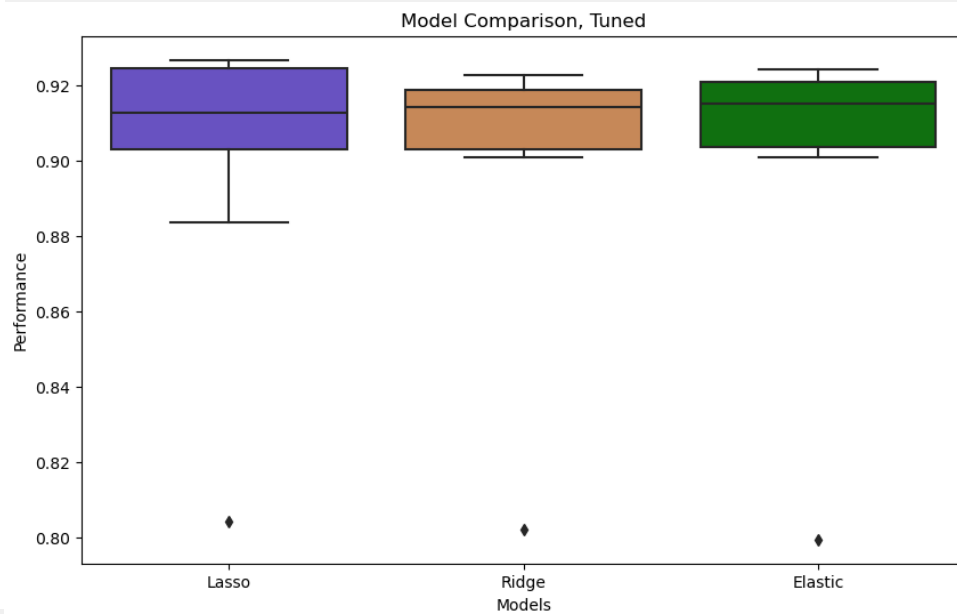
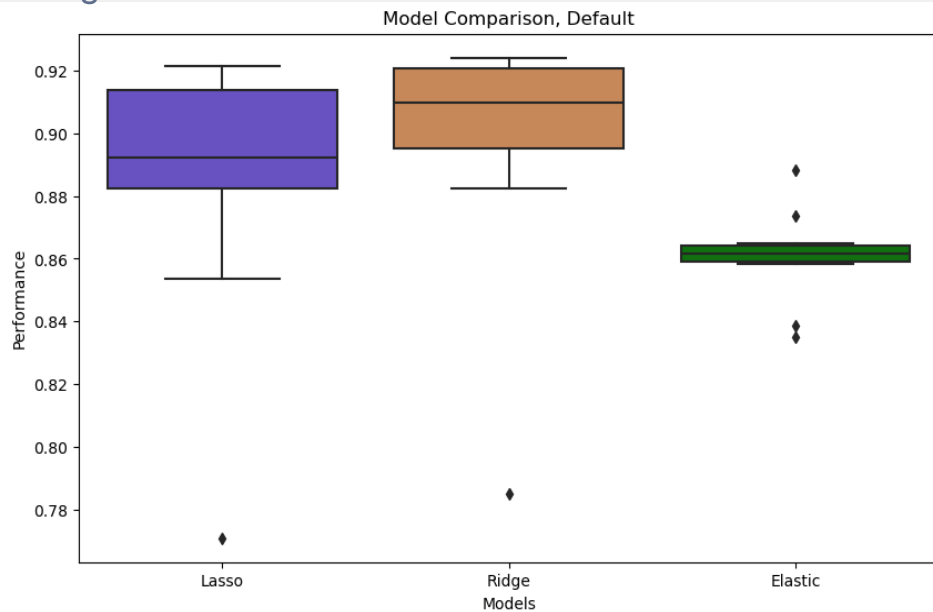
- Elastic Net is a hybrid model, adding a penalty equivalent to the sum of the absolute values (L1-norm) of the coefficients and the squares (L2-norm) of the coefficients. It is used when the Lasso becomes too dependent on a particular variable.
- The problem, being that it is a hybrid, is that the Elastic Net is extremely computationally intensive.



Regressions: Model Summary



- When it comes to the default, the Elastic model had the lowest mean CV score, while the regression model had the highest. The tuned model, on the other hand, has the elastic having the highest mean CV score, but the lasso, overall, has some of the highest fold scores out of the three.
- It should also be noted that the tuned Elastic model has one of the lowest fold score of the three, while the Lasso seems to have the greatest variation.





Random Forest: As a Model

- A Random Forest model combines multiple decision trees into a single model. In this case, it would determine feature importance rather than, for example, a MLR's correlations. Feature importance is a degree of dependency on the variables being compared. The higher feature importance, the larger the effect it has on the model.
- A decision tree is a hierarchical structure where each "node" represents branches based on different conditions being satisfied. As seen before, decision trees are subject to "overfitting", which can lead to "poor predictive performance".

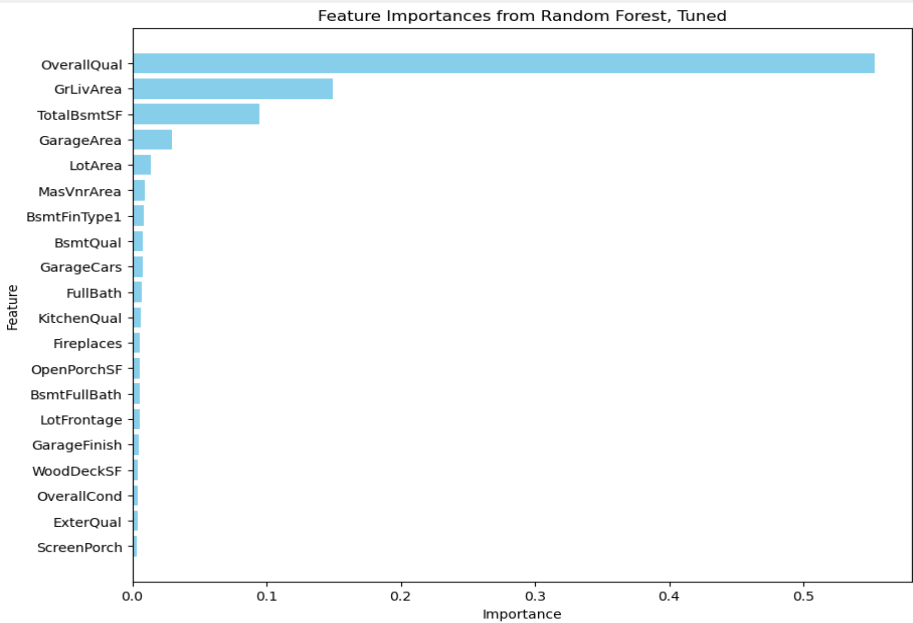
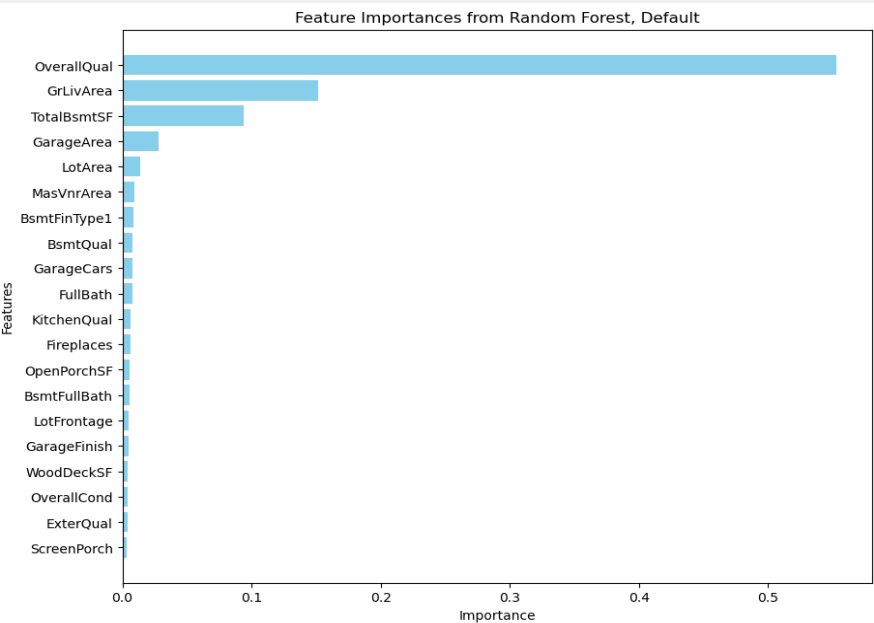


(Image Created Via Stable Diffusion)

RF: The Feature Importance

- With visually indistinct feature importances, the random forest models tell us not only a ranking of the features, but a model score. However, the model score, which is .886, is so close between both models it is almost not worth adjusting the default parameters.
- To the right, one can see just how close the importances are.

	Feature	Importance		Feature	Importance
4	OverallQual	5.526790e-01	4	OverallQual	5.528583e-01
1	GrLivArea	1.514606e-01	1	GrLivArea	1.492397e-01
7	TotalBsmtSF	9.358243e-02	7	TotalBsmtSF	9.456192e-02
17	GarageArea	2.779386e-02	17	GarageArea	2.934008e-02
3	LotArea	1.365511e-02	3	LotArea	1.409694e-02



Gradient Boosting

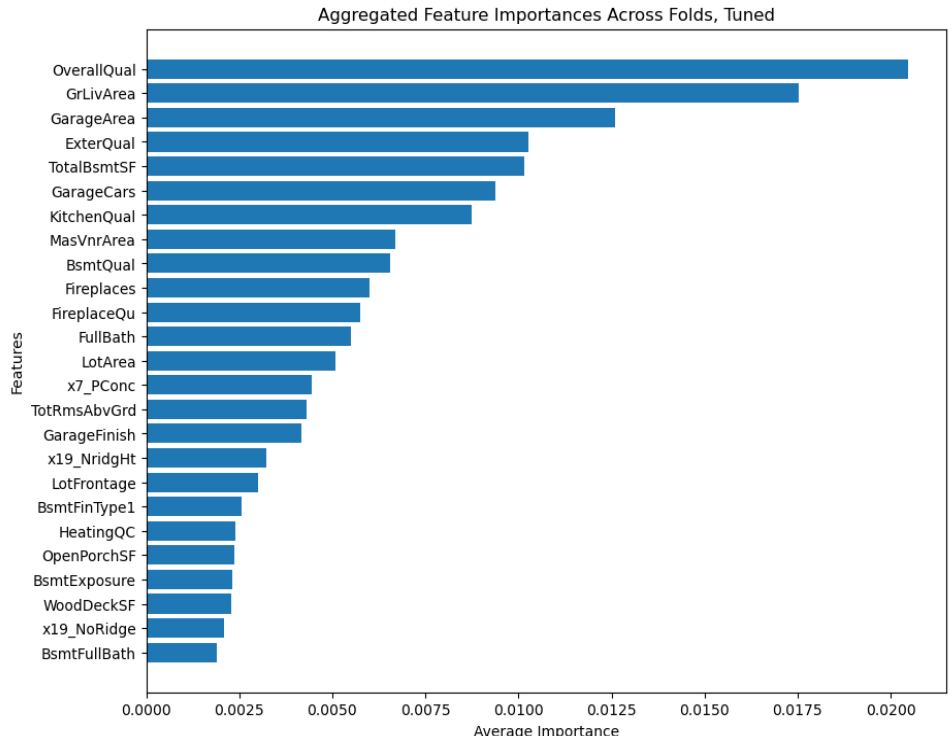
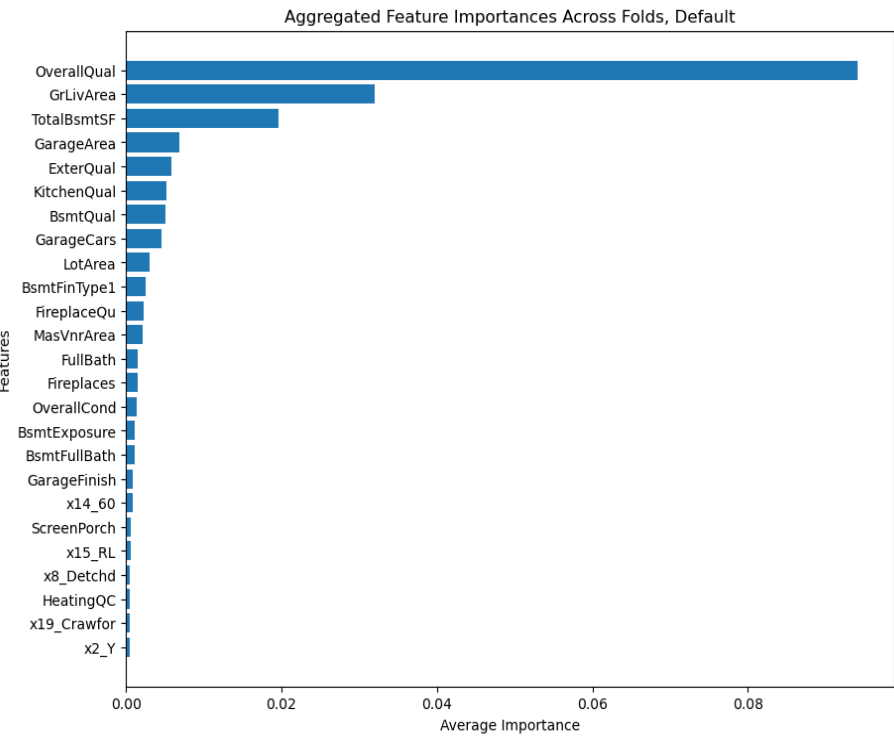
- Gradient boosting combines weaker models (usually decision trees), and adds said models sequentially, where each "addition" tries to make up for the errors made by the previous ones. The final prediction is an aggregate sum of the individual predictions.
- Because of the sequencing, the gradient boosting model reduces bias with the addition of each additional learner. This further allows for greater accuracy, especially on larger data sets.



(Image Created Via Stable Diffusion)

GB: The Booster

- The Gradient Boosting model is an interest one as its tuned version is had the best model performance out of all models, but the feature importances had a stark contrasts to each other. “Rank” should be looked at instead of the “strength”, as the GB will start taking into account One Hot Encoded values, which were a part of those weaker models.



Summary: Results

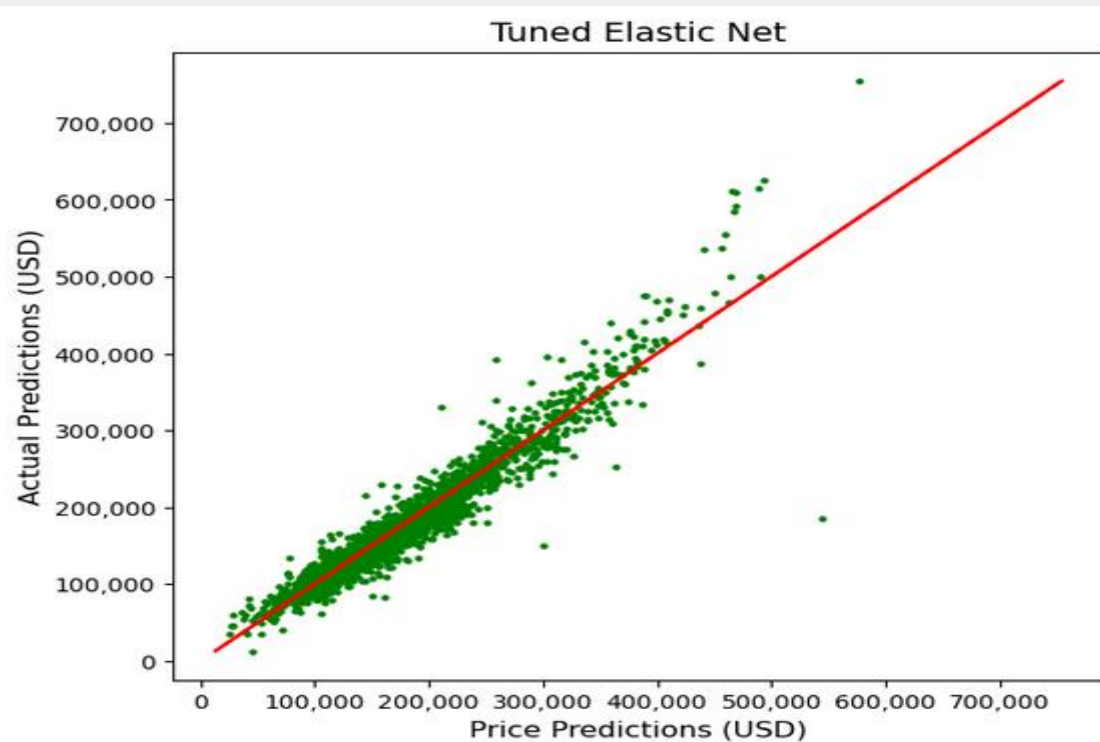
- The models themselves all have their strengths and weaknesses, but the results bring out which models may be preferred over the other. Of the regression models, the Elastic Net has the highest average CV score, although not by much of a margin.
- On the other hand, the Gradient Boosting model is the strongest model of the six, while the Random Forest model is the weakest. The MLR model was the only model that actually performed worse when subject to a tuning, so it was left untouched.

MLR Default Score	Lasso Default Score	Ridge Default Score	EN Default Score	RF Default Score	GB Default Score
0.9005	0.8844	0.8968	0.8605	0.8863	0.8947

MLR Tuned Score	Lasso Tuned Score	Ridge Tuned Score	EN Tuned Score	RF Tune Score	GB Tuned Score
N/A	0.9022	0.9023	0.9028	0.8864	0.9056

Final Thoughts: Linear Models

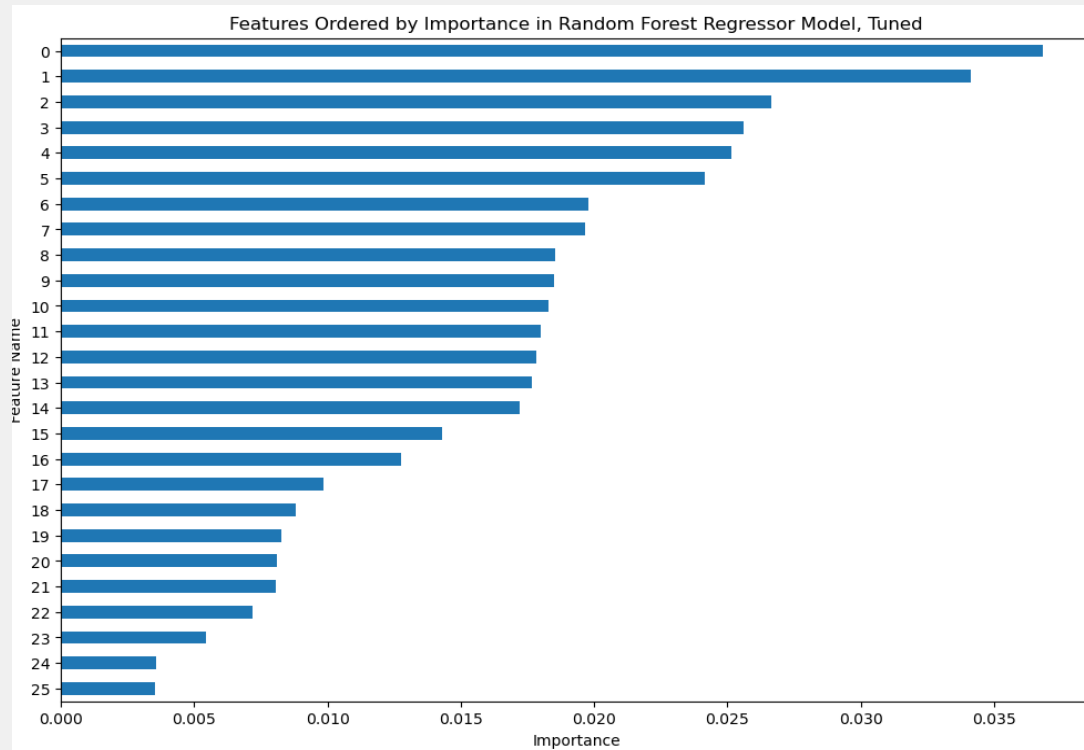
- The regression models are an interesting case in that they all not only are varied, but all have their advantages and disadvantages. Some of them are specifically used in certain situations, but after using four of them, there's no one that actually stands out well-above the rest.
- Therefore, it ends up turning into a “pick-and-choose” situation, where one would select regression models that suites the situation to compare them alongside other regression models. This is fine because, in our experience, these models are: compact, easy to use, and similar enough to each other that they can be compared with ease.
- Take this prediction for example from the Elastic Net. This is nearly identical visually across the regressions, so it's pointless to show a different one.



Final Thoughts: Random Forest



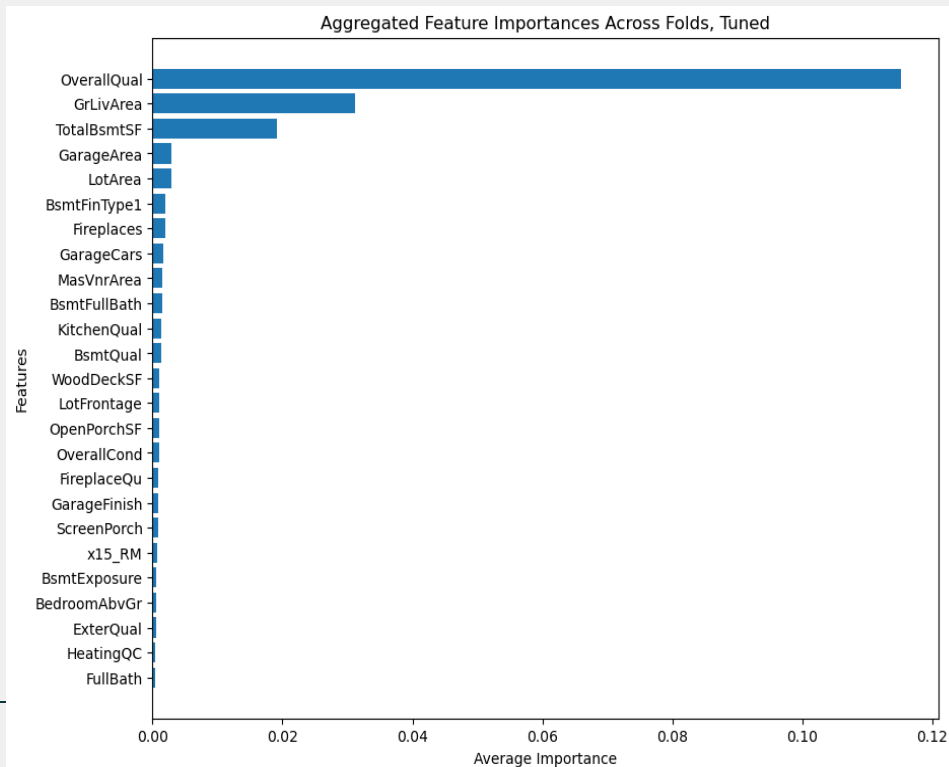
- The Random Forest is an interesting case as we found it to be the most unique out of the models presented. Although not shown, its construction method is visually distinct and that allows for greater parameter tuning.
- Yet even with that ability, it happened to be the weakest of all models. Worst of all, it was the most computationally expansive of all models. This is due to the parameter of “feature”, a parameter that has the chance to completely change the results. For example, to the right are the RF results if “feature” was square root instead of none. It would have a lower score, but everything would have higher importance. Note its similarity in appearance to the GB model.



Final Thoughts: Gradient Boosting



- A similar model to the Random Forest, the Gradient Boosting model was the best model in performance out of the six. It was computationally impressive (even with a “none” parameter, it was still faster than the RF), and allows one to extract similar results from a RF model.
- Included to the right is if square root was not used for the feature parameter, which shows just how similar the importance would be to a RF model. Its average importance is still weaker, but the spread is almost identical. However, this version of the model had a much weaker score than the version that was chosen.



Areas For Improvement

Modeling

1

Predictive Reliability: The score for all models was lower than desired. Ideally, they should all be above .90.

2

Beyond Score: The goal of the project was to find the model with the best score. While we did dabble in some other results, ideally one would use the models to truly find out the limits of what they're capable of.

3

Other Models: Other models can be used to address various issues. This includes classification modeling (regressors were used), so-called XGboosting and AdaBoosting, and bootstrap aggregating. Additionally the parameters can be adjusted.

Data

1

Feature Selection: We never actually used the MLR model for feature selection, which can be used to further tune the model.

2

Pipeline Changes: The data was edited in pipelines, done in different ways throughout the models to match their specifications. Obviously, the feature engineering was not perfect and can be changed to better adapt to certain models. Just for an example, the mean was used to fill null values for the numeric features. Sometimes this isn't ideal.

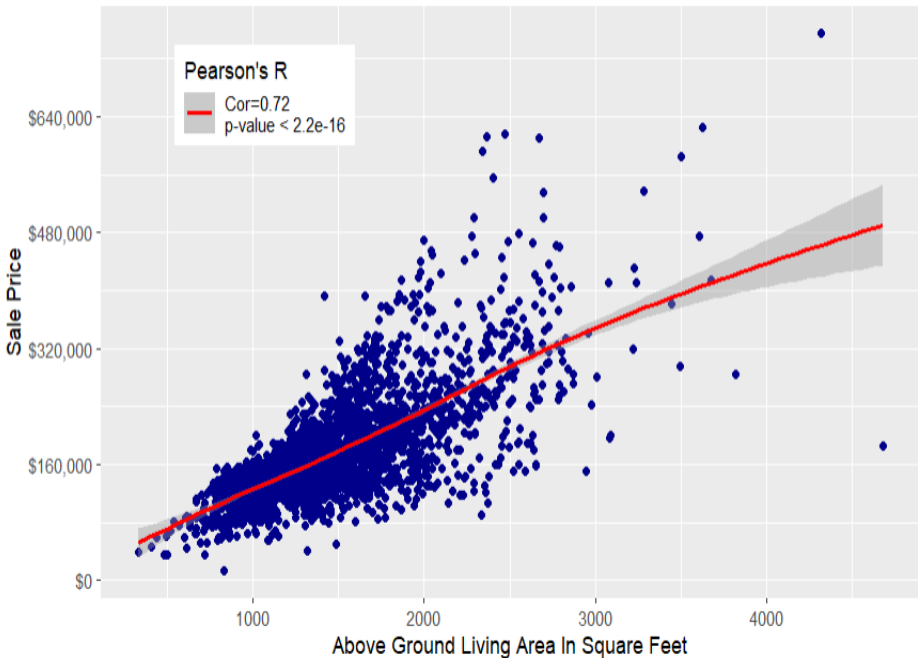
3

Feature Engineering : Some features were changed to account for their uselessness in the grand scheme of the data. This can be done for other features, such as with total baths.

Questions? Comments? Suggestions?

- Massive thank you to my mentor Vinod, who pushed me well beyond what I knew to complete this project.

Sale Price Regression on Above Ground Living Area



Sale Price Regression on Overall Quality

