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Introduction

As a data scientist working with Python, it's crucial to understand the importance of feature selection when building a machine learning model. In real-life data science problems, it's almost rare that all the variables in the dataset are useful for building a model. Adding redundant variables reduces the model's generalization capability and may also reduce the overall accuracy of a classifier. Furthermore, adding more variables to a model increases the overall complexity of the model.

As per the ***Law of Parsimony*** or '*Occam's Razor*', the best explanation of a problem is that which involves the fewest possible assumptions. Thus, feature selection becomes an indispensable part of building machine learning models.

Learning Objectives:

- Understanding the importance of feature selection.
- Familiarizing with different feature selection techniques.
- Applying feature selection techniques in practice and evaluating performance.

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What Is Feature Selection in Machine Learning?

The goal of feature selection techniques in machine learning is to find the best set of features that allows one to build optimized models of studied phenomena.

The techniques for feature selection in machine learning can be broadly classified into the following categories:

Supervised Techniques: These techniques can be used for labeled data and to identify the relevant features for increasing the efficiency of supervised models like classification and regression. For Example- linear regression, decision tree, SVM, etc.

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Now, let's discuss some of these popular machine learning feature selection methods in detail.

Types of Feature Selection Methods in ML

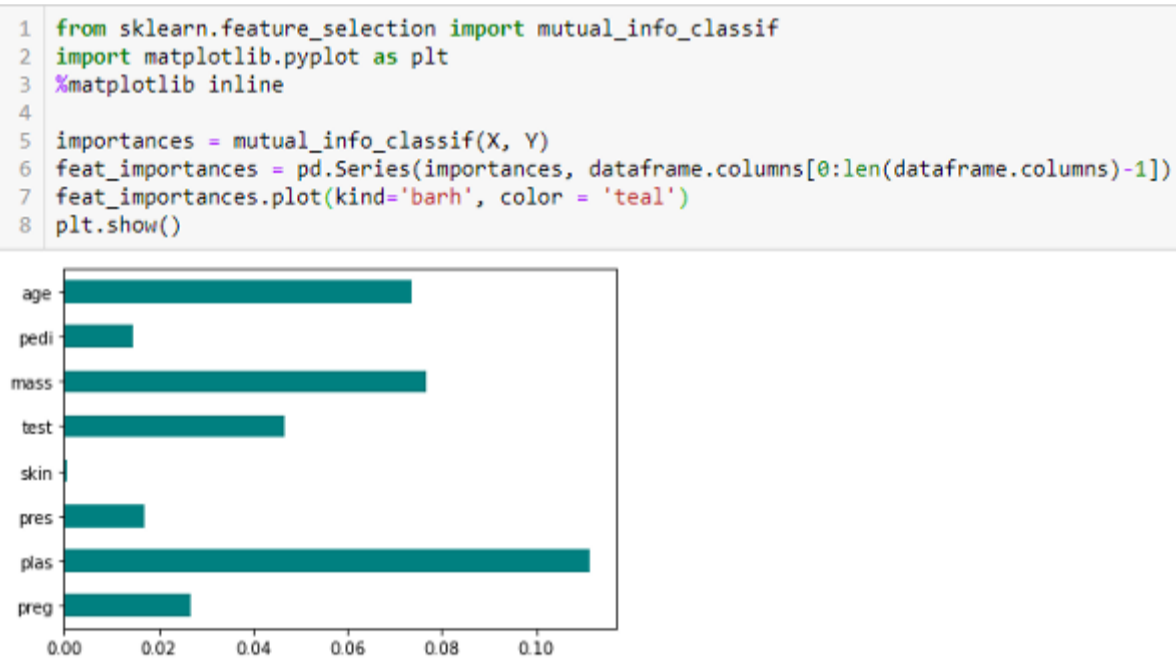
Filter Methods

Filter methods pick up the intrinsic properties of the features measured via univariate statistics instead of cross-validation performance. These methods are faster and less computationally expensive than wrapper methods. When dealing with high-dimensional data, it is computationally cheaper to use filter methods.

Let's, discuss some of these techniques:

Information Gain

Information gain calculates the reduction in entropy from the transformation of a dataset. It can be used for feature selection by evaluating the Information gain of each variable in the context of the target variable.



Chi-square Test

The Chi-square test is used for categorical features in a dataset. We calculate Chi-square between each feature and the target and select the desired number of features with the best Chi-square scores. In order to correctly apply the chi-squared to test the relation between various features in the dataset and the target variable, the following conditions have to be met: the variables have to be *categorical*, sampled *independently*, and values should have an *expected frequency greater than 5*.

```
1 from sklearn.feature_selection import SelectKBest
2 from sklearn.feature_selection import chi2
3
4 # Convert to categorical data by converting data to integers
5 X_cat = X.astype(int)
6
7 # Three features with highest chi-squared statistics are selected
8 chi2_features = SelectKBest(chi2, k = 3)
9 X_kbest_features = chi2_features.fit_transform(X_cat, Y)
10
11 # Reduced features
12 print('Original feature number:', X_cat.shape[1])
13 print('Reduced feature number:', X_kbest_features.shape[1])
```

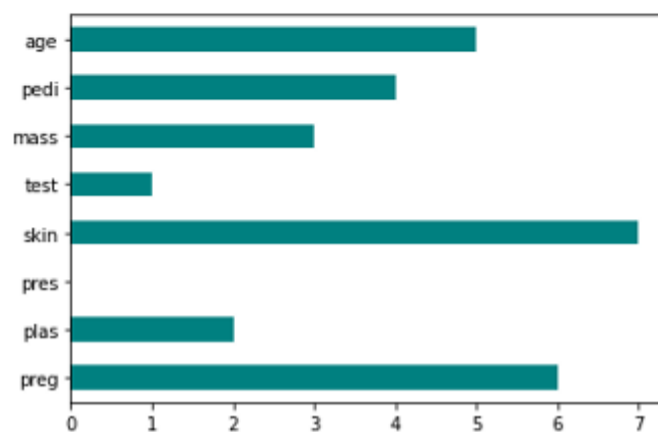
Original feature number: 8
Reduced feature number: 3

Fisher's Score

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of the variables based on the fisher's score in descending order. We can then select the variables as per the case.

```
1 from skfeature.function.similarity_based import fisher_score
2 import matplotlib.pyplot as plt
3 %matplotlib inline
4
5 # Calculating scores
6 ranks = fisher_score.fisher_score(X, Y)
7
8 # Plotting the ranks
9 feat_importances = pd.Series(ranks, dataframe.columns[0:len(dataframe.columns)-1])
10 feat_importances.plot(kind='barh', color = 'teal')
11 plt.show()
```



Correlation Coefficient

Correlation is a measure of the linear relationship between 2 or more variables. Through correlation, we can predict one variable from the other. The logic behind using correlation for feature selection is that good variables correlate highly with the target. Furthermore, variables should be correlated with the target but uncorrelated among themselves.

If two variables are correlated, we can predict one from the other. Therefore, if two features are correlated, the model only needs one, as the second does not add additional information. We will use the Pearson Correlation here.

```
1 import seaborn as sns
2 import matplotlib.pyplot as plt
3 %matplotlib inline
4
5 # Correlation matrix
6 cor = dataframe.corr()
7
8 # Plotting Heatmap
9 plt.figure(figsize = (10,6))
10 sns.heatmap(cor, annot = True)
```



We need to set an absolute value, say 0.5, as the threshold for selecting the variables. If we find that the predictor variables are correlated, we can drop the variable with a lower correlation coefficient value than the target variable. We can also compute multiple correlation coefficients to check whether more than two variables correlate. This phenomenon is known as multicollinearity.

Variance Threshold

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some threshold. By default, it removes all zero-variance features, i.e., features with the same value in all samples. We assume that features with a higher variance may contain more useful information, but note that we are not taking the relationship between feature variables or feature and target variables into account, which is one of the drawbacks of filter methods.

```
1 from sklearn.feature_selection import VarianceThreshold
2
3 # Resetting the value of X to make it non-categorical
4 X = array[:,0:8]
5
6 v_threshold = VarianceThreshold(threshold=0)
7 v_threshold.fit(X) # fit finds the features with zero variance
8 v_threshold.get_support()

array([ True,  True,  True,  True,  True,  True,  True,  True])
```

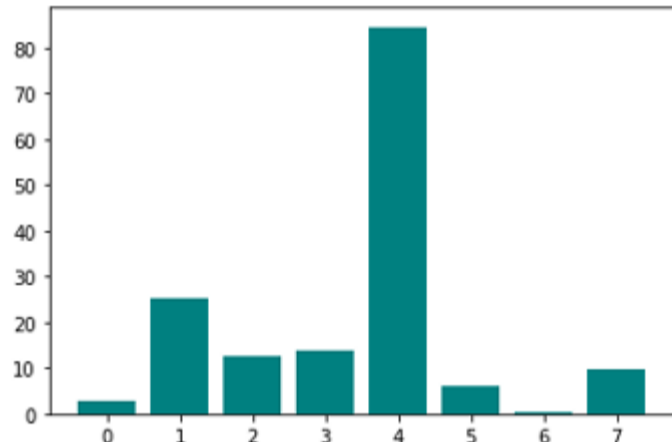
The get_support returns a Boolean vector where True means the variable does not have zero variance.

Mean Absolute Difference (MAD)

'The mean absolute difference (MAD) computes the absolute difference from the mean value. The main difference between the variance and MAD measures is the absence of the square in the latter. The MAD, like the variance, is also a scaled variant.' [1] This means that the higher the MAD, the higher the discriminatory power.

```
1 # Calculate MAD
2 mean_abs_diff = np.sum(np.abs(X - np.mean(X, axis = 0)), axis = 0)/X.shape[0]
3
4 # Plot the barchart
5 plt.bar(np.arange(X.shape[1]),mean_abs_diff, color = 'teal')

<BarContainer object of 8 artists>
```



Dispersion Ratio

'Another measure of dispersion applies the arithmetic mean (AM) and the geometric mean (GM). For a given (positive) feature X_i on n patterns, the AM and GM are given by

$$AM_i = \bar{X}_i = \frac{1}{n} \sum_{j=1}^n X_{ij}, \quad GM_i = \left(\prod_{j=1}^n X_{ij} \right)^{\frac{1}{n}},$$

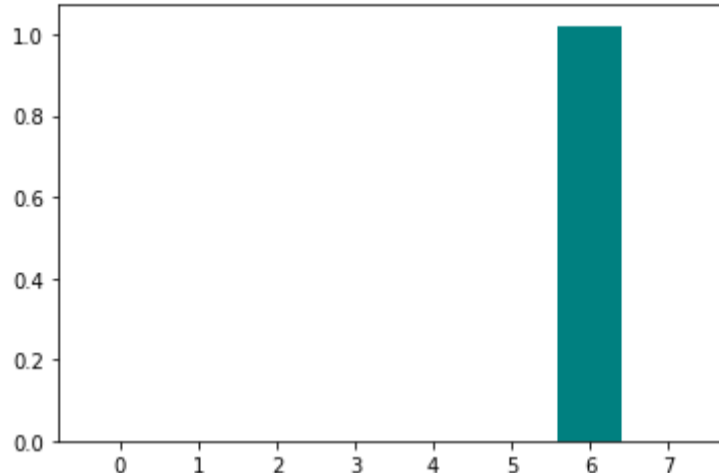
respectively; since $AM_i \geq GM_i$, with equality holding if and only if $X_{i1} = X_{i2} = \dots = X_{in}$, then the ratio

$$RM_i = \frac{AM_i}{GM_i} \in [1, +\infty),$$

can be used as a dispersion measure. Higher dispersion implies a higher value of R_i , thus a more relevant feature. Conversely, when all the feature samples have (roughly) the same value, R_i is close to 1, indicating a low relevance feature.' [1]

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```
3 am = np.mean(X, axis =0 )
4 #Geometric Mean
5 gm = np.power(np.prod(X, axis =0 ),1/X.shape[0])
6 # Ratio of Arithmetic Mean and Geometric Mean
7 disp_ratio = am/gm
8 # Plotting the bar chart
9 plt.bar(np.arange(X.shape[1]),disp_ratio, color = 'teal')
```



Wrapper Methods

Wrappers require some method to search the space of all possible subsets of features, assessing their quality by learning and evaluating a classifier with that feature subset. The feature selection process is based on a specific machine learning algorithm we are trying to fit on a given dataset. It follows a greedy search approach by evaluating all the possible combinations of features against the evaluation criterion. The wrapper methods usually result in better predictive accuracy than filter methods.

Let's, discuss some of these techniques:

Forward Feature Selection

This is an iterative method wherein we start with the performing features against the target features. Next, we select another variable that gives the best performance in combination with the first selected variable. This process continues until the preset criterion is achieved.

```
1 # Forward Feature Selection
2 from mlxtend.feature_selection import SequentialFeatureSelector
3 ffs = SequentialFeatureSelector(lr, k_features='best', forward = True, n_jobs=-1)
4 ffs.fit(X, Y)
5 features = list(ffs.k_feature_names_)
6 features = list(map(int, features))
7 lr.fit(x_train[features], y_train)
8 y_pred = lr.predict(x_train[features])
```

Backward Feature Elimination

This method works exactly opposite to the Forward Feature Selection method. Here, we start with all the features available and build a model. Next, we remove the variable from the model, which gives the best evaluation measure value. This process is continued until the preset criterion is achieved.

```
1 # Backward Feature Selection
2 from sklearn.linear_model import LogisticRegression
3 from mlxtend.feature_selection import SequentialFeatureSelector
4 lr = LogisticRegression(class_weight = 'balanced', solver = 'lbfgs', random_state=42, n_jobs=-1, max_iter=500)
5 lr.fit(X, Y)
6 bfs = SequentialFeatureSelector(lr, k_features='best', forward = False, n_jobs=-1)
7 bfs.fit(X, Y)
8 features = list(bfs.k_feature_names_)
9 features = list(map(int, features))
10 lr.fit(x_train[features], y_train)
11 y_pred = lr.predict(x_train[features])
```

This method, along with the one discussed above, is also known as the Sequential Feature Selection method.

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means it tries every possible combination of the variables and returns the best-performing subset.

```
1 # Exhaustive Feature Selection
2 from mlxtend.feature_selection import ExhaustiveFeatureSelector
3
4 # import the algorithm you want to evaluate on your features.
5 from sklearn.ensemble import RandomForestClassifier
6
7 # create the ExhaustiveFeatureSelector object.
8 efs = ExhaustiveFeatureSelector(RandomForestClassifier(),
9                                 min_features=4,
10                                max_features=8,
11                                scoring='roc_auc',
12                                cv=2)
13
14 # fit the object to the training data.
15 efs = efs.fit(X, Y)
16
17 # print the selected features.
18 selected_features = x_train.columns[list(efs.best_idx_)]
19 print(selected_features)
20
21 # print the final prediction score.
22 print(efs.best_score_)

Features: 163/163

Int64Index([0, 1, 2, 3, 4, 5, 6, 7], dtype='int64')
0.8252014925373135
```

Recursive Feature Elimination

'Given an external estimator that assigns weights to features (e.g., the coefficients of a linear model), the goal of recursive feature elimination (RFE) is to select features by recursively considering smaller and smaller sets of features. First, the estimator is trained on the initial set of features, and each feature's importance is obtained either through a `coef_` attribute or a `feature_importances_` attribute.

Then, the least important features are pruned from the current set of features. That procedure is recursively repeated on the pruned set until the desired number of features to select is eventually reached.'

```
1 # Recursive Feature Selection
2 from sklearn.feature_selection import RFE
3 rfe = RFE(lr, n_features_to_select=7)
4 rfe.fit(x_train, y_train)
5 y_pred = rfe.predict(x_train)
```

Embedded Methods

These methods encompass the benefits of both the wrapper and filter methods by including interactions of features but also maintaining reasonable computational costs. Embedded methods are iterative in the sense that takes care of each iteration of the model training process and carefully extract those features which contribute the most to the training for a particular iteration.

Let's discuss some of these techniques here:

LASSO Regularization (L1)

Regularization consists of adding a penalty to the different parameters of the machine learning model to reduce the freedom of the model, i.e., to avoid over-fitting. In linear model regularization, the penalty is applied over the coefficients that multiply each predictor. From the different types of regularization, Lasso or L1 has the property that can shrink some of the coefficients to zero. Therefore, that feature can be removed from the model.

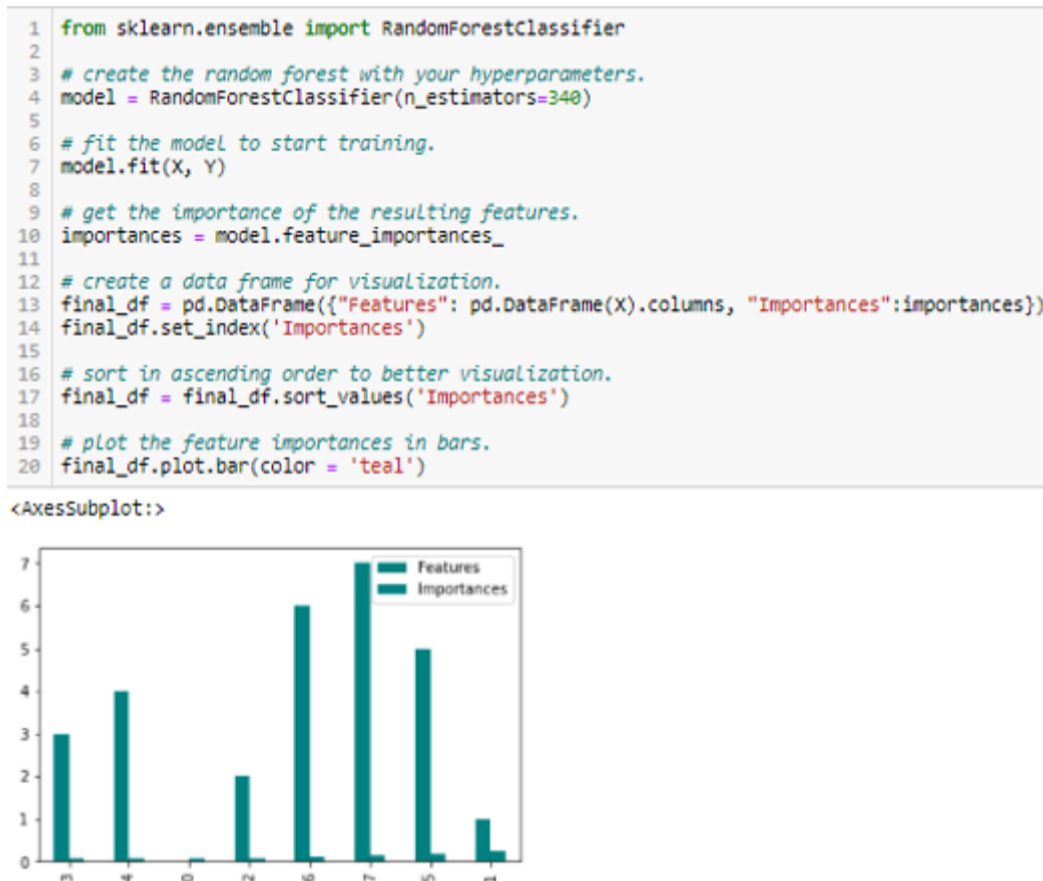
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```
4 # Set the regularization parameter C=1
5 logistic = LogisticRegression(C=1, penalty="l1", solver='liblinear', random_state=7).fit(X, Y)
6 model = SelectFromModel(logistic, prefit=True)
7
8 X_new = model.transform(X)
9
10 # Dropped columns have values of all 0s, keep other columns
11 selected_columns = selected_features.columns[selected_features.var() != 0]
12 selected_columns
```

Int64Index([0, 1, 2, 3, 4, 5, 6, 7], dtype='int64')

Random Forest Importance

Random Forests is a kind of Bagging Algorithm that aggregates a specified number of decision trees. The tree-based strategies used by random forests naturally rank by how well they improve the purity of the node, or in other words, a decrease in the impurity (**Gini impurity**) over all trees. Nodes with the greatest decrease in impurity happen at the start of the trees, while nodes with the least decrease in impurity occur at the end of the trees. Thus, by pruning trees below a particular node, we can create a subset of the most important features.



Conclusion

We have discussed a few techniques for feature selection. We have purposely left the feature extraction techniques like Principal Component Analysis, Singular Value Decomposition, Linear Discriminant Analysis, etc. These methods help to reduce the dimensionality of the data or reduce the number of variables while preserving the variance of the data.

Apart from the methods discussed above, there are many other feature selection methods. There are hybrid methods, too, that use both filtering and wrapping techniques. If you wish to explore more about feature selection techniques, great comprehensive reading material, in my opinion, would be '*Feature Selection for Data and Pattern Recognition*' by Urszula Stańczyk and Lakhmi C. Jain.

Key Takeaways

- Understanding the importance of feature selection and feature engineering in building a machine learning model.
- Familiarizing with different feature selection techniques, including supervised techniques (Information Gain, Chi-square Test, Fisher's Score, Correlation Coefficient), unsupervised techniques (Variance Threshold, Mean Absolute Difference, Dispersion Ratio) and their classifications (Filter methods, Wrapper methods, Embedded methods, Hybrid methods).

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Q1. What are the different types of feature selection techniques?

A. Here are some ways of selecting the best features out of all the features to increase the model performance as the irrelevant features decrease the model performance of the machine learning or deep learning model.

- Filter Methods: Select features based on statistical measures such as correlation or chi-squared test. For example- Correlation-based Feature Selection, chi2 test, SelectKBest, and ANOVA F-value.
- Wrapper Methods: Select features by evaluating their combinations using a predictive model. For example- Recursive Feature Elimination, Backward Feature Elimination, Forward Feature Selection
- Embedded Methods: Select features by learning their importance during model training. For example- Lasso Regression, Ridge Regression, and Random Forest.
- Hybrid Methods: Combine the strengths of filter and wrapper methods. For Example- SelectFromModel
- Dimensionality Reduction Techniques: Reduce the dimensionality of the dataset and select the most important features. For Example- pca, Ida, and ica.

Q2. What are the three steps in feature selection?

A. The three steps of feature selection can be summarized as follows:

1. Data Preprocessing: Clean and prepare the data for feature selection.
2. Feature Scoring: Compute scores for each feature to reflect its importance to the target variable.
3. Selection: Select a subset of the most important features based on their scores, and use them for training the predictive model.

Q3. What is the difference between a feature selection technique and a classification algorithm?

A. Feature selection is a process in machine learning to identify important features in a dataset to improve the performance and interpretability of the model. Classification algorithms, on the other hand, are used to predict a categorical label based on the input features, such as logistic regression, decision trees, and neural networks.

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About the Author

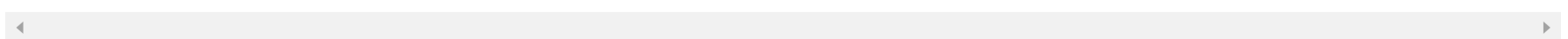


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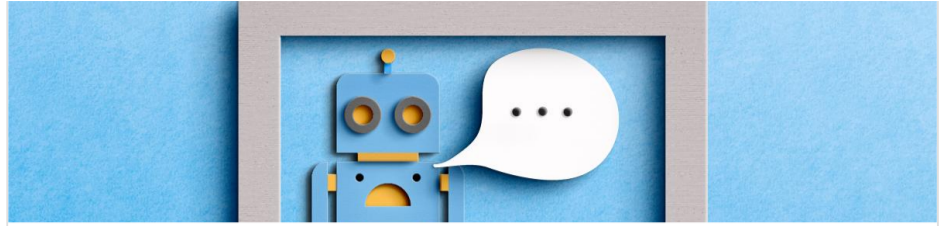
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