Classification Tutorial

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Outline

- Introduction
- Decision Tree Classifiers
- Nearest Neighbors
- Support Vector Machines
- Bagging & Boosting
- **6** Contact Information

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Introduction

- What is Classification?
- Advantages & Disadvantages
- Classification & Time Series Data

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Decision Trees

Decision trees are a flow chart like structure that are created using decisions rules inferred from the features of the data.

Nodes

Nodes are features (attributes) of data. The top node of a tree is known as the "Root Node"

Branches

Decision rules

Leaves

Outcomes (labels)

Decision trees are easy to understand and easy to visualize.

Decision Trees

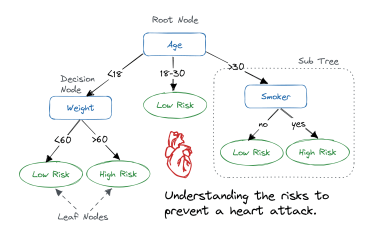


Figure: Decision Tree

Decision Algorithm

Select the best attribute

There several Attribute Selection Measures that can be used such as Information Gain, Gain Ration, or Gini Index.

- Make selected attribute a decision node
- Recursively build tree until one of the following conditions are met:

All tuples belong to the same attribute value

No attributes remain

No instances remain

Advantages

- Easy to understand & visualize (white box)
- Requires little data preperation
- Cost of using a tree is logarithmic
- Can handle numeric and categorical data
- Can handle multi-output problems
- Can be validated using statistical tests

Disdvantages

- Can create overly complex models that do not generalize well (overfitting)
- Requires little data preperation
- Cost of using a tree is logarithmic
- Can handle numeric and categorical data
- Can handle multi-output problems
- On be validated using statistical tests

Random Forest

Random Forest is a model composed of multiple decision trees. In machine learning, a model composed of multiple models is called an **ensemble**.

Each decision tree in a forest is created using a **random** subset of data and features. This technique is known as bagging.

A greedy search is used to select the values at which to split a feature.

Each tree classifies input independently, then the ensemble makes the final decision by voting.

Extra Trees

Extra Trees is an ensemble similar to Random Forest.

Unlike Random Forest, Extra Trees used the entire dataset to train the individual decision trees.

To ensure decision trees are sufficiently different, split values for each feature are randomly selected,

Extra Trees & Random Forest Parameters

n_estimators

The number of decision trees to use in the ensemble

More trees can improve performance, but increase computational time.

@ max_depth

The maximum depth of each decision tree in the forest

Too high of a depth can lead to overfitting while a too low depth can lead to underfitting.

min_samples_leaf

The minimum number of samples required for a leaf node

Higher values are less likely to overfit, while higher values create deeper and more specialized trees

Extra Trees & Random Forest Comparison

Extra Trees uses randomness to reduce variance

Extra Trees computational cost compared to Random Forest is much smaller

Extra Trees reduces bias compared to Random Forest because all the trees use the entire dataset, not just a random subset

Extra Trees excel with high dimension datasets

Extra Trees' randomness in tree building aids in preventing overfitting

Extra Trees' trains faster than Random Forests

Extra Trees can obtain high predictive accuracy, especially with large datasets

Extra Trees & Random Forest Comparison

Extra Trees and Random Forests have low interpertability

For both methods, small changes in Hyperparamters can create large changes in results

Random Forests are often more accurate than Extra Trees, except when features are reduced to the most essential, then they can perform about the same

Random Forests are less influenced by outliers than Extra Trees

Trees Tutorial Code

Build Pipes

```
decision_tree =
skc.pipeBuild_DecisionTreeClassifier(criterion=['gini','entropy'],
    max_depth=[5, 10])

xtree = skc.pipeBuild_ExtraTreesClassifier(criterion=['gini','entropy'],
    n_estimators=[10], max_depth=[3, 5, 10],max_features=[1])

random_forest =
skc.pipeBuild_RandomForestClassifier(criterion=['gini','entropy'],
    n_estimators=[10], max_depth=[3, 5, 10], max_features=[1])
```

Trees Tutorial Code

Set Up Grid Seach

```
names=['Decision Tree', 'Extra Trees', 'Random Forest']
pipes=[decision_tree, xtree, random_forest]
skc.gridsearch_classifier(names=names, pipes=pipes, X_train=X_train, X_test=X_test, y_train=y_train, y_test=y_test, plot_number=4)
```

Classifiying Waveforms with Trees

Data: 1st sample of each waveform type

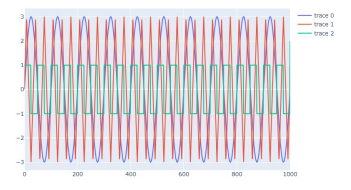


Figure: Sine, Square, & Triangle Waves

Classifiying Waveforms with Trees

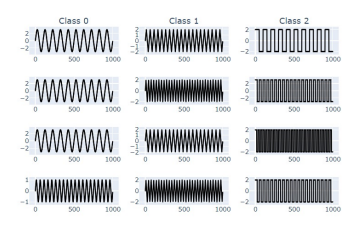


Figure: Waveforms Classified

Classifying Waveforms with Trees

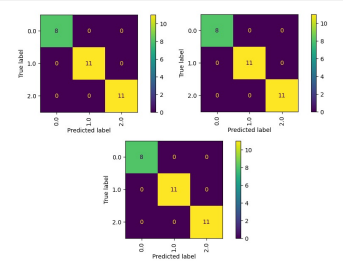


Figure: Confusion Matrices

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Nearest Neighbors

Nearest Neighbors

Nearest Neighbors algorithms classify a data point by determining which class a majority of the points neighbors belong to

There are several methods for determining neighbors

K Nearest Neighbors

KNN

Assumption: "Similar things exist in close proximity"

Estimates the likelihood of a data point being a member of a group or another depending on the data points nearest to it are in

Note: There is a "Time Series" version of this algorithm avaliable in the TS Learn package. It is the same as the standard KNN, but with distance metrics more suited for times series data, such as Dynamic Time Warping.

K Nearest Neighbors

KNN Algorithm

- 1. Load data
- **2.** Initialize **K** to your chosen # of neighbors
- **3.** For each datapoint: Calculate the distance between querry example & all current examples in the dataset
- 4. Then add the distance & index of the example to a ordered collection
- **5.** Sort collection in descending order
- **6.** Select the first **K** entries
- **7.** If regression, return the mean of the **K** labels
- 7. If classification, return the mode of the K labels

Nearest Centroid

Nearest Centroid is one of the simplest machine learning algorithms for classification.

Algorithm

- 1. Load Training Data
- 2. Compute the centroid of each class in the training data
- **3.** For each new data point, calculate its distance from the centroid of each class
- **4.** Select the class with the smallest distance from the data point and assign it that class's label

Radius Nearest Neighbors

Radius NN works the same as the KNN algrothim, except it compares a data sample to all neighbors within a specified radius instead of a fixed number.

Algorithm

- 1. Load data
- 2. Select the length of your radius
- **3.** For each data point: Calculate the distance between query example & all current examples in the dataset
- **4.** Then add all points with a distance equal to or less than the specified radius to a collection
- **5.** If regression, return the mean of the collection's labels
- 5. If classification, return the mode of the collection's labels

Nearest Neighbors Pros & Cons

Pros:

Fast. In particular, KNN is a Lazy Learner, which has no training period.

Easy to implement

Easy to understand

Useful for both Classification & Regression

Cons:

Does not work well with large datasets

Does not work well with high dimensions

Requires feature scaling

Sensitive to noise

Nearest Neighbors Tutorial Code

Build Pipes

```
knn = skc.pipeBuild_KNeighborsClassifier(n_neighbors=[3,5],
weights=['uniform'], algorithm=['auto'])

ncent = skc.pipeBuild_NearestCentroid()

rad = skc.pipeBuild_RadiusNeighborsClassifier(radius=[50])

tsknn = skc.pipeBuild_KNeighborsTimeSeriesClassifier(n_neighbors=[3,5],
weights=['uniform'], metric=['dtw','softdtw'])
```

Nearest Neighbors Tutorial Code

Set Up Grid Search

```
names=['K Nearest Neighbors', 'Nearest Centroid', 'Radius Neighbors', 'Time Series KNN']
```

```
pipes=[knn, ncent, rad, tsknn]
```

```
skc.gridsearch\_classifier(names=names, pipes=pipes, X\_train=X\_train, X\_test=X\_test, y\_train=y\_train, y\_test=y\_test, plot\_number=3)
```

Nearest Neighbors Confusion Matrix

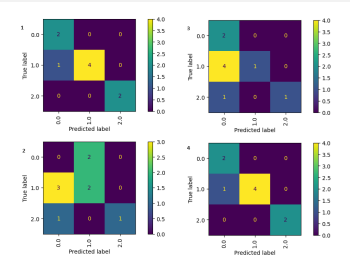


Figure: Confusion Matrices

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Filters

Support Vector Machines (SVM) can be used for classification and regression.

SVMs construct a **hyperplane** in multidimensional space to separate different classes.

The **hyperplane** is iteratively constructed using error minimization to find the optimal hyperplane.

Support Vector Machine Terminology

Support Vectors

A set of data points closest to the hyperplane.

4 Hyperplane

A decision plane which separates objects by class.

Margin

The gap between a hyperplane and support vectors. The distance is measured perpendicular from the plane to the vector.

Maximum Marginal Hyperplane

The hyperplane that gives the best division of classes. The larger the margin, the better.

Kernel

The kernel is the method by which an SVM transforms lower dimensional data into higher dimensional data. Hyperplanes are created in the higher dimensional space. There are multiple kernels available.

SVMs Illustrated

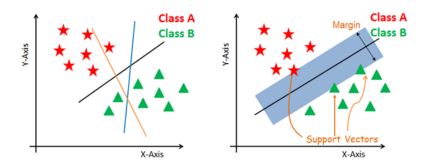


Figure: Simple SVM Illustration

SVMs Illustrated

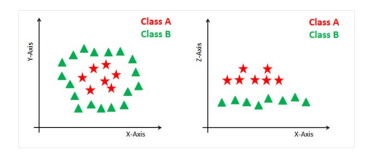


Figure: Simple Kernel Illustration

SVM Kernels

Linear

$$k(x,y) = x^T y$$

Polynomial

$$k(x,y) = (\gamma x^T y + c_0)^d$$

Gaussian Radial Basis Function (RBF)

$$k(x,y) = exp\left(-\gamma \|x - y\|^2\right)$$

SVM Kernels

Sigmoid

$$k(x,y) = tanh(\gamma x^T y + c_0)$$

Global Alignment Kernal (GAK)

$$k(x,y) = \sum_{\pi \in A(x,y)} \prod_{i=1}^{|\pi|} exp\left(-\frac{\|x_{1\pi_1(i)} - y_{\pi 2j}\|^2}{2\sigma^2}\right)$$

GAK Kernel

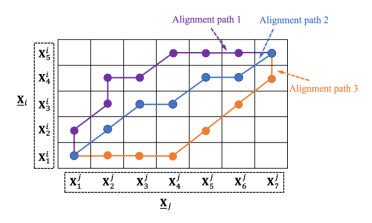


Figure: GAK Kernel Illustration

SVM Pros & Cons

Pros

Work well when there is a clear margin of seperation

More effective in higher dimensional space

Effective in cases where there are more dimensions than the number of samples

Relatively memory efficient

Cons

Doesn't work well with large data sets

Doesn't perform well with a lot of noise

Under-performs when the number of features exceeds the number of samples

Difficult to explain decision process

SVM Tutorial Code

Build Pipes

```
\label{eq:svc} \begin{split} & \textbf{svc} = \text{skc.pipeBuild\_SVC}(C=[1.0], \text{ kernel}=['linear'], \\ & \text{degree}=[3], \text{gamma}=['\text{scale'}], \text{ tol}=[1.0\text{e-}3], \text{ random\_state}=None) \\ & \textbf{nusvc} = \text{skc.pipeBuild\_NuSVC}(\text{nu}=[0.5], \text{ kernel}=['\text{rbf'}], \text{ degree}=[3], \\ & \text{gamma}=['\text{scale'}], \text{ tol}=[1.0\text{e-}3], \text{random\_state}=None) \\ & \textbf{tssvc} = \text{skc.pipeBuild\_TimeSeriesSVC}(\text{kernel}=['\text{gak'}]) \end{split}
```

SVM Tutorial Code

Setup Grid Search

```
names = ['SVClassifier', 'Nu-SVClassifier']
pipes = [svc, nusvc]
```

 $skc.gridsearch_classifier(names=names, pipes=pipes, X_train=X_train, X_test=X_test, y_train=y_train, y_test=y_test, plot_number=4)$

SVM Confusion Matrices

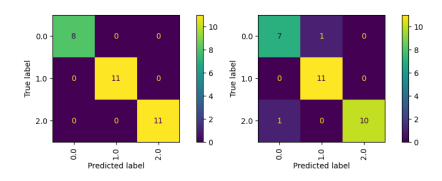


Figure: SVM & NuSVM Results

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Overview

Weak Learners

An algorithm that predicts slightly better than random.

Boosting

An ensemble of **sequential** weak learners combined for and accurate algorithm.

Predictions are made iteratively for "T" rounds on the entire training set and each iteration improves performance using information from the previous iteration.

Overview

Bagging

An ensemble of non-sequential weak learners combined for and accurate algorithm. This method is also known as **bootstrapping**

Random subsets of training data are drawn for "T" rounds. Each draw is independent. Each draw is used to create a weak learner.

The weak learners run in parallel after training, and a vote is taken to determine the class of new input. Voting may or may not be weighted.

AdaBoost

AdaBoost is composed of sequentially grown trees.

It "punishes" incorrectly predicted samples by assigning a larger weight to them after each round during of prediction.

In **SKLearn**, AdaBoost defaults to using decision trees, but other classification algorithms may be used with it.

AdaBoost Algorithm

For t in T rounds

- 1. Normalize the weight vector \mathbf{w} to calculate distribution \mathbf{p} . \mathbf{w} initial weight is $\mathbf{1/N}$, where \mathbf{N} is the number of labeled examples
- **2.** Grow a decision tree using the distribution ${\bf p}$ then return hypothesis ${\bf h}$ with prediction values for each example
- 3. Calculate error term epsilon of h
- **4.** Assign $\beta = \epsilon \frac{\epsilon}{1-\epsilon}$
- **5.** Update weights to $\mathbf{w} = \mathbf{w} * \mathbf{B}$ such that poor predictions will have higher weights and good predictions have lower weights

AdaBoost Pros & Cons

Pros

Relatively robust to over-fitting on low noise data sets Few hyper-parameters for easy tuning

Easy to understand & visualize

Cons

Debate among experts as to whether or not it generalizes will with noisy data

When irrelevant features are in the model, AdaBoost performs worse than Random Forests & Extra Trees

Not optimized for speed

Other Bagging & Boosting Models

Gradient Boosting

Gradient Boosting is essentially AdaBoost with Decisions Trees. It's an optimized version specifically for trees.

SKLearn Bagging Classifier

This is a generic baggling classifier. With it, you may use the boot strapping approach with any model

Bagging Pros & Cons

Pros

Robust against noise.

Easy & effective way to convert weak learners into strong learners

Reduces variance & over-fitting for more accurate models

Cons

Requires many comparable classifiers

Can result in under-fitting if not trained properly

Computationally more expensive as the number of parallel models increases

Bagging & Boosting Tutorial Code

Build Pipes

```
decision_tree = skc.pipeBuild_DecisionTreeClassifier(criterion=['gini',
'entropy'], max_depth=[5, 10])
xtree = skc.pipeBuild_ExtraTreesClassifier(n_estimators=[50,100])
random_forest =
skc.pipeBuild_RandomForestClassifier(criterion=['gini', 'entropy'],
n_{estimators}=[50,100], max_depth=[3, 5, 10], max_features=[1])
ada =
skc.pipeBuild_AdaBoostClassifier(n_estimators=[50,100],learning_rate=[1.0])
bag = skc.pipeBuild_BaggingClassifier(n_estimators=[50,100])
\mathbf{gb} = \text{skc.pipeBuild\_GradientBoostingClassifier}(n_\text{estimators} = [50,100])
```

Bagging & Boosting Tutorial Code

Set Up Grid Search

```
names = ['Decision Tree', 'Extra Trees', 'Random Forest', 'Adaboost',
'Bagging', 'Gradient Boosting']
pipes = [decision_tree, xtree, random_forest, ada, bag, gb]
skc.gridsearch_classifier(names=names, pipes=pipes, X_train=X_train,
X_test=X_test, y_train=y_train, y_test=y_test, plot_number=3)
```

Bagging & Boosting Confusion Matrices

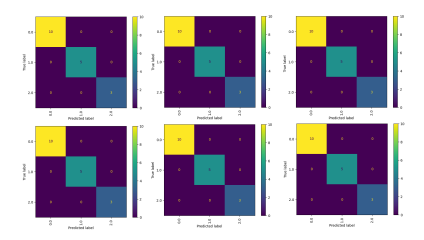


Figure: Tree Comparisons

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Thank you!

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