**Broadly, there are 3 types of Machine Learning Algorithms..**

**1. Supervised Learning**

**How it works:** This algorithm consist of a target / outcome variable (or dependent variable) which is to be predicted from a given set of predictors (independent variables). Using these set of variables, we generate a function that map inputs to desired outputs. The training process continues until the model achieves a desired level of accuracy on the training data. Examples of Supervised Learning: Regression, Decision Tree, Random Forest, KNN, Logistic Regression etc.

**2. Unsupervised Learning**

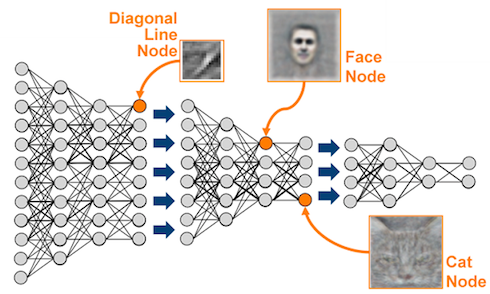
**How it works:**In this algorithm, we do not have any target or outcome variable to predict / estimate.  It is used for clustering population in different groups, which is widely used for segmenting customers in different groups for specific intervention. Examples of Unsupervised Learning: Apriori algorithm, K-means.

**3. Reinforcement Learning:**

**How it works:**  Using this algorithm, the machine is trained to make specific decisions. It works this way: the machine is exposed to an environment where it trains itself continually using trial and error. This machine learns from past experience and tries to capture the best possible knowledge to make accurate business decisions. Example of Reinforcement Learning: Markov Decision Process

**What is Deep Learning?**

Deep learning is probably one of the hottest topics in Machine learning today, and it has shown significant improvement over some of its counterparts. It falls under a class of unsupervised learning algorithms and uses multi-layered neural networks to achieve these remarkable outcomes. Here is a simple illustration from Analytic Store’s blog:



A large number of pixels are fed to the network as input, after which the network learns and evolves to recognize higher level features like faces and cats.

Here are a few achievements driving the attention to this area:

* Amazing accuracy on quite a few Kaggle competitions – Dogs vs. Cats image recognition (98.9% accuracy), Saving the whales problem (98% accuracy)
* Ability to learn and identify cats by using YouTube videos without any supervision

Following are a few events, which suggest the lookout for people with knowledge of deep learning:

* Google’s acquisition of Deepmind Technologies
* Google hiring Jeff Hinton, one of the thought leaders in this space (you can check out his course on neural networks on Coursera)
* Facebook hiring Yann LeCun, a student of Jeff Hinton to lead its AI lab
* Baidu hires Andrew Ng, another pioneer in the field (and  co-founder of Coursera).

**Applications of deep learning:**

If you have not figured it out already, deep learning finds its applications in following areas:

* Image recognition (e.g. Tagging faces in photos)
* Voice recognition (e.g. Voice based search, Siri)
* Pattern detection (e.g. Handwriting recognition)

**But, neural networks have been there for decades, what is re-kindling this interest now?**

Yes, neural networks have existed since ages. Interest in neural networks peaked in the 1980s and 90s and then died off because of the inherent problems with them and black box like approach.

There are a few reasons why this is happening now. The biggest one among them, being the drop in computational costs. Classification of cats through unsupervised learning of YouTube videos was achieved by deploying 16,000 computers in Google lab! The cost of deploying these algorithms is not small, even by today’s standards.

**List of Common Machine Learning Algorithms**

Here is the list of commonly used machine learning algorithms. These algorithms can be applied to almost any data problem:

1. Linear Regression
2. Logistic Regression
3. Decision Tree
4. SVM
5. Naive Bayes
6. kNN
7. K-Means
8. Random Forest
9. Dimensionality Reduction Algorithms
10. Gradient Boosting algorithms
    1. GBM
    2. XGBoost
    3. LightGBM
    4. CatBoost

**1. Linear Regression**

It is used to estimate real values (cost of houses, number of calls, total sales etc.) based on continuous variable(s). Here, we establish relationship between independent and dependent variables by fitting a best line. This best fit line is known as regression line and represented by a linear equation Y= a \*X + b.

The best way to understand linear regression is to relive this experience of childhood. Let us say, you ask a child in fifth grade to arrange people in his class by increasing order of weight, without asking them their weights! What do you think the child will do? He / she would likely look (visually analyze) at the height and build of people and arrange them using a combination of these visible parameters. This is linear regression in real life! The child has actually figured out that height and build would be correlated to the weight by a relationship, which looks like the equation above.

In this equation:

* Y – Dependent Variable
* a – Slope
* X – Independent variable
* b – Intercept

These coefficients a and b are derived based on minimizing the sum of squared difference of distance between data points and regression line.

Look at the below example. Here we have identified the best fit line having linear equation **y=0.2811x+13.9**. Now using this equation, we can find the weight, knowing the height of a person.



Linear Regression is of mainly two types: Simple Linear Regression and Multiple Linear Regression. Simple Linear Regression is characterized by one independent variable. And, Multiple Linear Regression(as the name suggests) is characterized by multiple (more than 1) independent variables. While finding best fit line, you can fit a polynomial or curvilinear regression. And these are known as polynomial or curvilinear regression.

**Python Code**

#Import Library

#Import other necessary libraries like pandas, numpy...

from sklearn import linear\_model

#Load Train and Test datasets

#Identify feature and response variable(s) and values must be numeric and numpy arrays

x\_train=input\_variables\_values\_training\_datasets

y\_train=target\_variables\_values\_training\_datasets

x\_test=input\_variables\_values\_test\_datasets

# Create linear regression object

linear = linear\_model.LinearRegression()

# Train the model using the training sets and check score

linear.fit(x\_train, y\_train)

linear.score(x\_train, y\_train)

#Equation coefficient and Intercept

print('Coefficient: \n', linear.coef\_)

print('Intercept: \n', linear.intercept\_)

#Predict Output

predicted= linear.predict(x\_test)

**R Code**

#Load Train and Test datasets

#Identify feature and response variable(s) and values must be numeric and numpy arrays

x\_train <- input\_variables\_values\_training\_datasets

y\_train <- target\_variables\_values\_training\_datasets

x\_test <- input\_variables\_values\_test\_datasets

x <- cbind(x\_train,y\_train)

# Train the model using the training sets and check score

linear <- lm(y\_train ~ ., data = x)

summary(linear)

#Predict Output

predicted= predict(linear,x\_test)

**2. Logistic Regression**

Don’t get confused by its name! It is a classification not a regression algorithm. It is used to estimate discrete values ( Binary values like 0/1, yes/no, true/false ) based on given set of independent variable(s). In simple words, it predicts the probability of occurrence of an event by fitting data to a logit function. Hence, it is also known as **logit regression**. Since, it predicts the probability, its output values lies between 0 and 1 (as expected).

Again, let us try and understand this through a simple example.

Let’s say your friend gives you a puzzle to solve. There are only 2 outcome scenarios – either you solve it or you don’t. Now imagine, that you are being given wide range of puzzles / quizzes in an attempt to understand which subjects you are good at. The outcome to this study would be something like this – if you are given a trignometry based tenth grade problem, you are 70% likely to solve it. On the other hand, if it is grade fifth history question, the probability of getting an answer is only 30%. This is what Logistic Regression provides you.

Coming to the math, the log odds of the outcome is modeled as a linear combination of the predictor variables.

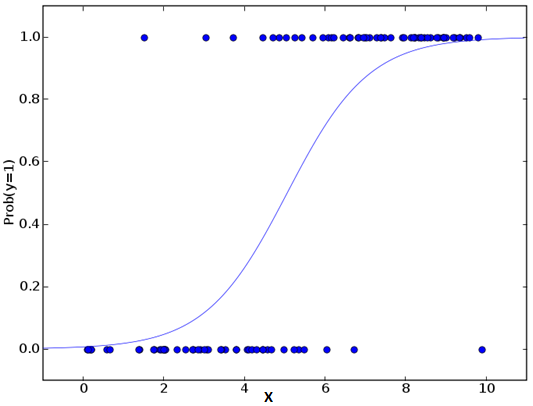
odds= p/ (1-p) = probability of event occurrence / probability of not event occurrence

ln(odds) = ln(p/(1-p))

logit(p) = ln(p/(1-p)) = b0+b1X1+b2X2+b3X3....+bkXk

Above, p is the probability of presence of the characteristic of interest. It chooses parameters that maximize the likelihood of observing the sample values rather than that minimize the sum of squared errors (like in ordinary regression).

Now, you may ask, why take a log? For the sake of simplicity, let’s just say that this is one of the best mathematical way to replicate a step function. I can go in more details, but that will beat the purpose of this article.

**Python Code**

#Import Library

from sklearn.linear\_model import LogisticRegression

#Assumed you have, X (predictor) and Y (target) for training data set and x\_test(predictor) of test\_dataset

# Create logistic regression object

model = LogisticRegression()

# Train the model using the training sets and check score

model.fit(X, y)

model.score(X, y)

#Equation coefficient and Intercept

print('Coefficient: \n', model.coef\_)

print('Intercept: \n', model.intercept\_)

#Predict Output

predicted= model.predict(x\_test)

**R Code**

x <- cbind(x\_train,y\_train)

# Train the model using the training sets and check score

logistic <- glm(y\_train ~ ., data = x,family='binomial')

summary(logistic)

#Predict Output

predicted= predict(logistic,x\_test)

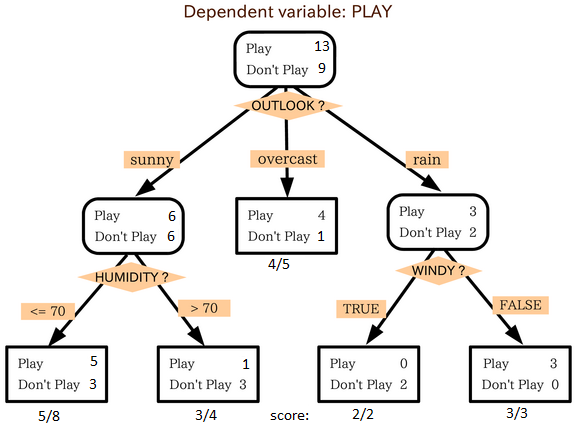
**Furthermore..**

There are many different steps that could be tried in order to improve the model:

* including interaction terms
* removing features
* regularization techniques
* using a non-linear model

**3. Decision Tree**

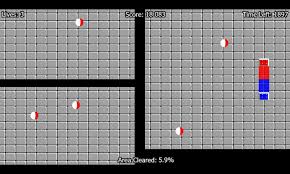
This is one of my favorite algorithm and I use it quite frequently. It is a type of supervised learning algorithm that is mostly used for classification problems. Surprisingly, it works for both categorical and continuous dependent variables. In this algorithm, we split the population into two or more homogeneous sets. This is done based on most significant attributes/ independent variables to make as distinct groups as possible. For more details, you can read: Decision Tree Simplified.



source: statsexchange

In the image above, you can see that population is classified into four different groups based on multiple attributes to identify ‘if they will play or not’. To split the population into different heterogeneous groups, it uses various techniques like Gini, Information Gain, Chi-square, entropy.

The best way to understand how decision tree works, is to play Jezzball – a classic game from Microsoft (image below). Essentially, you have a room with moving walls and you need to create walls such that maximum area gets cleared off with out the balls.



So, every time you split the room with a wall, you are trying to create 2 different populations with in the same room. Decision trees work in very similar fashion by dividing a population in as different groups as possible.

*More*: Simplified Version of Decision Tree Algorithms

**Python Code**

#Import Library

#Import other necessary libraries like pandas, numpy...

from sklearn import tree

#Assumed you have, X (predictor) and Y (target) for training data set and x\_test(predictor) of test\_dataset

# Create tree object

model = tree.DecisionTreeClassifier(criterion='gini') # for classification, here you can change the algorithm as gini or entropy (information gain) by default it is gini

# model = tree.DecisionTreeRegressor() for regression

# Train the model using the training sets and check score

model.fit(X, y)

model.score(X, y)

#Predict Output

predicted= model.predict(x\_test)

**R Code**

library(rpart)

x <- cbind(x\_train,y\_train)

# grow tree

fit <- rpart(y\_train ~ ., data = x,method="class")

summary(fit)

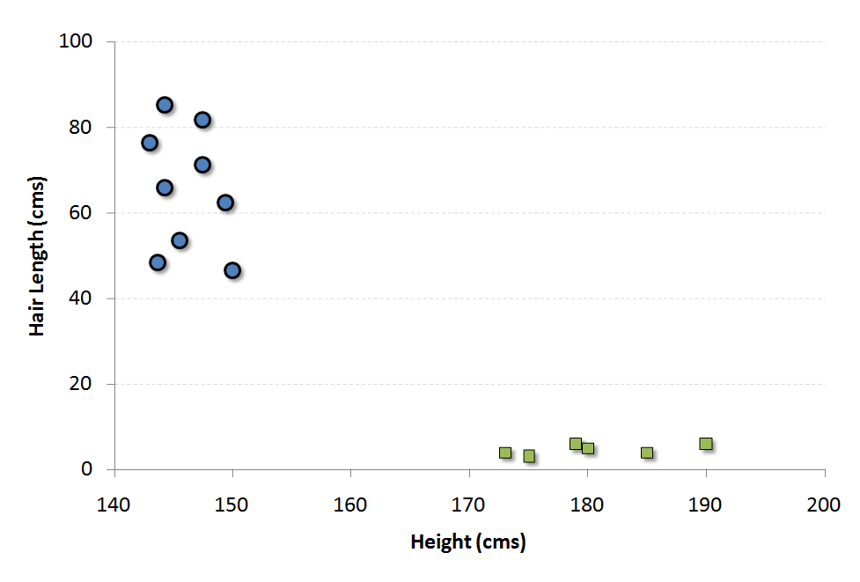
#Predict Output

predicted= predict(fit,x\_test)

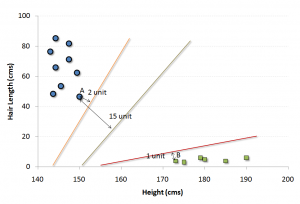
**4. SVM (Support Vector Machine)**

It is a classification method. In this algorithm, we plot each data item as a point in n-dimensional space (where n is number of features you have) with the value of each feature being the value of a particular coordinate.

For example, if we only had two features like Height and Hair length of an individual, we’d first plot these two variables in two dimensional space where each point has two co-ordinates (these co-ordinates are known as **Support Vectors**)



Now, we will find some *line* that splits the data between the two differently classified groups of data. This will be the line such that the distances from the closest point in each of the two groups will be farthest away.



In the example shown above, the line which splits the data into two differently classified groups is the *black* line, since the two closest points are the farthest apart from the line. This line is our classifier. Then, depending on where the testing data lands on either side of the line, that’s what class we can classify the new data as.

More: Simplified Version of Support Vector Machine

**Think of this algorithm as playing JezzBall in n-dimensional space. The tweaks in the game are:**

* You can draw lines / planes at any angles (rather than just horizontal or vertical as in classic game)
* The objective of the game is to segregate balls of different colors in different rooms.
* And the balls are not moving.

**Python Code**

#Import Library

from sklearn import svm

#Assumed you have, X (predictor) and Y (target) for training data set and x\_test(predictor) of test\_dataset

# Create SVM classification object

model = svm.svc() # there is various option associated with it, this is simple for classification. You can refer link, for mo# re detail.

# Train the model using the training sets and check score

model.fit(X, y)

model.score(X, y)

#Predict Output

predicted= model.predict(x\_test)

**R Code**

library(e1071)

x <- cbind(x\_train,y\_train)

# Fitting model

fit <-svm(y\_train ~ ., data = x)

summary(fit)

#Predict Output

predicted= predict(fit,x\_test)

**5. Naive Bayes**

It is a classification technique based on Bayes’ theorem with an assumption of independence between predictors. In simple terms, a Naive Bayes classifier assumes that the presence of a particular feature in a class is unrelated to the presence of any other feature. For example, a fruit may be considered to be an apple if it is red, round, and about 3 inches in diameter. Even if these features depend on each other or upon the existence of the other features, a naive Bayes classifier would consider all of these properties to independently contribute to the probability that this fruit is an apple.

Naive Bayesian model is easy to build and particularly useful for very large data sets. Along with simplicity, Naive Bayes is known to outperform even highly sophisticated classification methods.

Bayes theorem provides a way of calculating posterior probability P(c|x) from P(c), P(x) and P(x|c). Look at the equation below:  

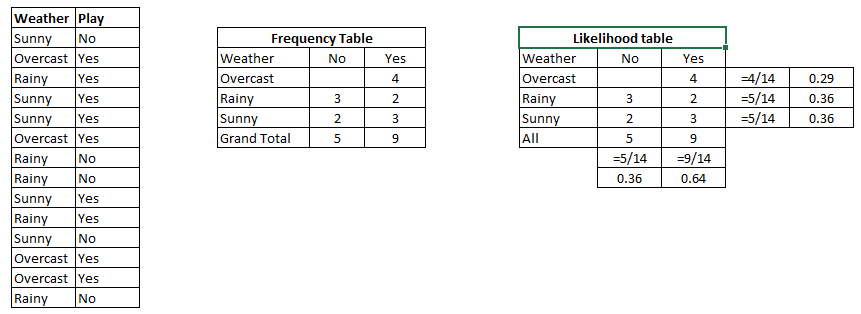

Here,

* *P*(*c|x*) is the posterior probability of *class* (*target*) given *predictor* (*attribute*).
* *P*(*c*) is the prior probability of *class*.
* *P*(*x|c*) is the likelihood which is the probability of *predictor* given *class*.
* *P*(*x*) is the prior probability of *predictor*.

**Example:**Let’s understand it using an example. Below I have a training data set of weather and corresponding target variable ‘Play’. Now, we need to classify whether players will play or not based on weather condition. Let’s follow the below steps to perform it.

Step 1: Convert the data set to frequency table

Step 2: Create Likelihood table by finding the probabilities like Overcast probability = 0.29 and probability of playing is 0.64.



Step 3: Now, use Naive Bayesian equation to calculate the posterior probability for each class. The class with the highest posterior probability is the outcome of prediction.

**Problem:**Players will pay if weather is sunny, is this statement is correct?

We can solve it using above discussed method, so P(Yes | Sunny) = P( Sunny | Yes) \* P(Yes) / P (Sunny)

Here we have P (Sunny |Yes) = 3/9 = 0.33, P(Sunny) = 5/14 = 0.36, P( Yes)= 9/14 = 0.64

Now, P (Yes | Sunny) = 0.33 \* 0.64 / 0.36 = 0.60, which has higher probability.

Naive Bayes uses a similar method to predict the probability of different class based on various attributes. This algorithm is mostly used in text classification and with problems having multiple classes.

**Python Code**

#Import Library

from sklearn.naive\_bayes import GaussianNB

#Assumed you have, X (predictor) and Y (target) for training data set and x\_test(predictor) of test\_dataset

# Create SVM classification object model = GaussianNB() # there is other distribution for multinomial classes like Bernoulli Naive Bayes, Refer link

# Train the model using the training sets and check score

model.fit(X, y)

#Predict Output

predicted= model.predict(x\_test)

**R Code**

library(e1071)

x <- cbind(x\_train,y\_train)

# Fitting model

fit <-naiveBayes(y\_train ~ ., data = x)

summary(fit)

#Predict Output

predicted= predict(fit,x\_test)

**6. kNN (k- Nearest Neighbors)**

It can be used for both classification and regression problems. However, it is more widely used in classification problems in the industry. K nearest neighbors is a simple algorithm that stores all available cases and classifies new cases by a majority vote of its k neighbors. The case being assigned to the class is most common amongst its K nearest neighbors measured by a distance function.

These distance functions can be Euclidean, Manhattan, Minkowski and Hamming distance. First three functions are used for continuous function and fourth one (Hamming) for categorical variables. If K = 1, then the case is simply assigned to the class of its nearest neighbor. At times, choosing K turns out to be a challenge while performing kNN modeling.

More: Introduction to k-nearest neighbors : Simplified.



KNN can easily be mapped to our real lives. If you want to learn about a person, of whom you have no information, you might like to find out about his close friends and the circles he moves in and gain access to his/her information!

**Things to consider before selecting kNN:**

* KNN is computationally expensive
* Variables should be normalized else higher range variables can bias it
* Works on pre-processing stage more before going for kNN like outlier, noise removal

**Python Code**

#Import Library

from sklearn.neighbors import KNeighborsClassifier

#Assumed you have, X (predictor) and Y (target) for training data set and x\_test(predictor) of test\_dataset

# Create KNeighbors classifier object model

KNeighborsClassifier(n\_neighbors=6) # default value for n\_neighbors is 5

# Train the model using the training sets and check score

model.fit(X, y)

#Predict Output

predicted= model.predict(x\_test)

**R Code**

library(knn)

x <- cbind(x\_train,y\_train)

# Fitting model

fit <-knn(y\_train ~ ., data = x,k=5)

summary(fit)

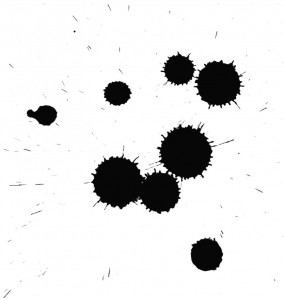
#Predict Output

predicted= predict(fit,x\_test)

**7. K-Means**

It is a type of unsupervised algorithm which  solves the clustering problem. Its procedure follows a simple and easy  way to classify a given data set through a certain number of  clusters (assume k clusters). Data points inside a cluster are homogeneous and heterogeneous to peer groups.

Remember figuring out shapes from ink blots? k means is somewhat similar this activity. You look at the shape and spread to decipher how many different clusters / population are present!



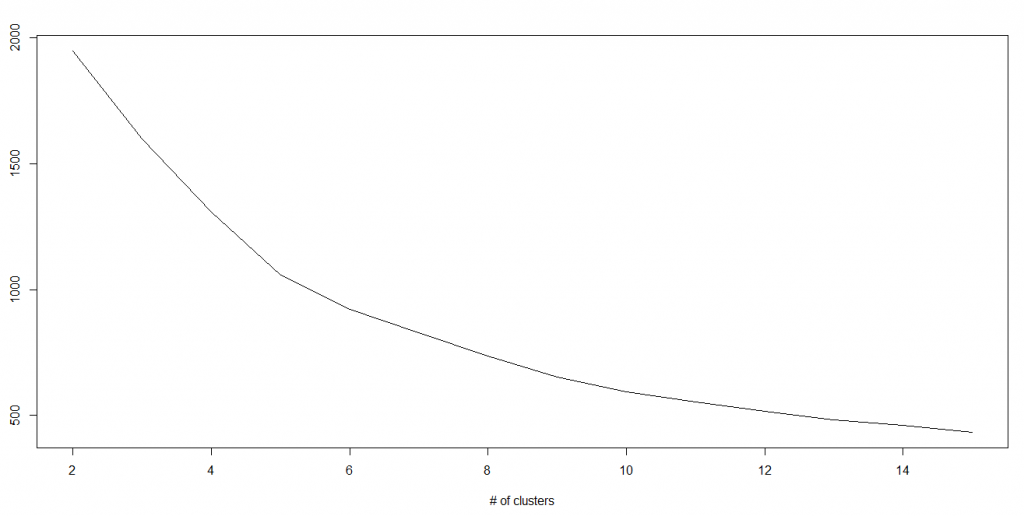
**How K-means forms cluster:**

1. K-means picks k number of points for each cluster known as centroids.
2. Each data point forms a cluster with the closest centroids i.e. k clusters.
3. Finds the centroid of each cluster based on existing cluster members. Here we have new centroids.
4. As we have new centroids, repeat step 2 and 3. Find the closest distance for each data point from new centroids and get associated with new k-clusters. Repeat this process until convergence occurs i.e. centroids does not change.

**How to determine value of K:**

In K-means, we have clusters and each cluster has its own centroid. Sum of square of difference between centroid and the data points within a cluster constitutes within sum of square value for that cluster. Also, when the sum of square values for all the clusters are added, it becomes total within sum of square value for the cluster solution.

We know that as the number of cluster increases, this value keeps on decreasing but if you plot the result you may see that the sum of squared distance decreases sharply up to some value of k, and then much more slowly after that. Here, we can find the optimum number of cluster.



**Python Code**

#Import Library

from sklearn.cluster import KMeans

#Assumed you have, X (attributes) for training data set and x\_test(attributes) of test\_dataset

# Create KNeighbors classifier object model

k\_means = KMeans(n\_clusters=3, random\_state=0)

# Train the model using the training sets and check score

model.fit(X)

#Predict Output

predicted= model.predict(x\_test)

**R Code**

library(cluster)

fit <- kmeans(X, 3) # 5 cluster solution

**8. Random Forest**

Random Forest is a trademark term for an ensemble of decision trees. In Random Forest, we’ve collection of decision trees (so known as “Forest”). To classify a new object based on attributes, each tree gives a classification and we say the tree “votes” for that class. The forest chooses the classification having the most votes (over all the trees in the forest).

Each tree is planted & grown as follows:

1. If the number of cases in the training set is N, then sample of N cases is taken at random but *with replacement*. This sample will be the training set for growing the tree.
2. If there are M input variables, a number m<<M is specified such that at each node, m variables are selected at random out of the M and the best split on these m is used to split the node. The value of m is held constant during the forest growing.
3. Each tree is grown to the largest extent possible. There is no pruning.

For more details on this algorithm, comparing with decision tree and tuning model parameters, I would suggest you to read these articles:

1. Introduction to Random forest – Simplified
2. Comparing a CART model to Random Forest (Part 1)
3. Comparing a Random Forest to a CART model (Part 2)
4. Tuning the parameters of your Random Forest model

**Python**

#Import Library

from sklearn.ensemble import RandomForestClassifier

#Assumed you have, X (predictor) and Y (target) for training data set and x\_test(predictor) of test\_dataset

# Create Random Forest object

model= RandomForestClassifier()

# Train the model using the training sets and check score

model.fit(X, y)

#Predict Output

predicted= model.predict(x\_test)

**R Code**

library(randomForest)

x <- cbind(x\_train,y\_train)

# Fitting model

fit <- randomForest(Species ~ ., x,ntree=500)

summary(fit)

#Predict Output

predicted= predict(fit,x\_test)

**9. Dimensionality Reduction Algorithms**

In the last 4-5 years, there has been an exponential increase in data capturing at every possible stages. Corporates/ Government Agencies/ Research organisations are not only coming with new sources but also they are capturing data in great detail.

For example: E-commerce companies are capturing more details about customer like their demographics, web crawling history, what they like or dislike, purchase history, feedback and many others to give them personalized attention more than your nearest grocery shopkeeper.

As a data scientist, the data we are offered also consist of many features, this sounds good for building good robust model but there is a challenge. How’d you identify highly significant variable(s) out 1000 or 2000? In such cases, dimensionality reduction algorithm helps us along with various other algorithms like Decision Tree, Random Forest, PCA, Factor Analysis, Identify based on correlation matrix, missing value ratio and others.

To know more about this algorithms, you can read “Beginners Guide To Learn Dimension Reduction Techniques“.

**Python  Code**

#Import Library

from sklearn import decomposition

#Assumed you have training and test data set as train and test

# Create PCA obeject pca= decomposition.PCA(n\_components=k) #default value of k =min(n\_sample, n\_features)

# For Factor analysis

#fa= decomposition.FactorAnalysis()

# Reduced the dimension of training dataset using PCA

train\_reduced = pca.fit\_transform(train)

#Reduced the dimension of test dataset

test\_reduced = pca.transform(test)

#For more detail on this, please refer  this link.

**R Code**

library(stats)

pca <- princomp(train, cor = TRUE)

train\_reduced <- predict(pca,train)

test\_reduced <- predict(pca,test)

**10. Gradient Boosting Algorithms**

**10.1. GBM**

GBM is a boosting algorithm used when we deal with plenty of data to make a prediction with high prediction power. Boosting is actually an ensemble of learning algorithms which combines the prediction of several base estimators in order to improve robustness over a single estimator. It combines multiple weak or average predictors to a build strong predictor. These boosting algorithms always work well in data science competitions like Kaggle, AV Hackathon, CrowdAnalytix.

More: Know about Boosting algorithms in detail

**Python Code**

#Import Library

from sklearn.ensemble import GradientBoostingClassifier

#Assumed you have, X (predictor) and Y (target) for training data set and x\_test(predictor) of test\_dataset

# Create Gradient Boosting Classifier object

model= GradientBoostingClassifier(n\_estimators=100, learning\_rate=1.0, max\_depth=1, random\_state=0)

# Train the model using the training sets and check score

model.fit(X, y)

#Predict Output

predicted= model.predict(x\_test)

**R Code**

library(caret)

x <- cbind(x\_train,y\_train)

# Fitting model

fitControl <- trainControl( method = "repeatedcv", number = 4, repeats = 4)

fit <- train(y ~ ., data = x, method = "gbm", trControl = fitControl,verbose = FALSE)

predicted= predict(fit,x\_test,type= "prob")[,2]

GradientBoostingClassifier and Random Forest are two different boosting tree classifier and often people ask about the difference between these two algorithms.

**10.2. XGBoost**

Another classic gradient boosting algorithm that’s known to be the decisive choice between winning and losing in some Kaggle competitions.

The XGBoost has an immensely high predictive power which makes it the best choice for accuracy in events as it possesses both linear model and the tree learning algorithm, making the algorithm almost 10x faster than existing gradient booster techniques.

The support includes various objective functions, including regression, classification and ranking.

One of the most interesting things about the XGBoost is that it is also called a regularized boosting technique. This helps to reduce overfit modelling and has a massive support for a range of languages such as Scala, Java, R, Python, Julia and C++.

Supports distributed and widespread training on many machines that encompass GCE, AWS, Azure and Yarn clusters. XGBoost can also be integrated with Spark, Flink and other cloud dataflow systems with a built in cross validation at each iteration of the boosting process.

To learn more about XGBoost and parameter tuning, visit https://www.analyticsvidhya.com/blog/2016/03/complete-guide-parameter-tuning-xgboost-with-codes-python/.

Python Code:

from xgboost import XGBClassifier

from sklearn.model\_selection import train\_test\_split

from sklearn.metrics import accuracy\_score

X = dataset[:,0:10]

Y = dataset[:,10:]

seed = 1

X\_train, X\_test, y\_train, y\_test = train\_test\_split(X, Y, test\_size=0.33, random\_state=seed)

model = XGBClassifier()

model.fit(X\_train, y\_train)

#Make predictions for test data

y\_pred = model.predict(X\_test)

R Code:

require(caret)

x <- cbind(x\_train,y\_train)

# Fitting model

TrainControl <- trainControl( method = "repeatedcv", number = 10, repeats = 4)

model<- train(y ~ ., data = x, method = "xgbLinear", trControl = TrainControl,verbose = FALSE)

OR

model<- train(y ~ ., data = x, method = "xgbTree", trControl = TrainControl,verbose = FALSE)

predicted <- predict(model, x\_test)

**10.3. LightGBM**

LightGBM is a gradient boosting framework that uses tree based learning algorithms. It is designed to be distributed and efficient with the following advantages:

* Faster training speed and higher efficiency
* Lower memory usage
* Better accuracy
* Parallel and GPU learning supported
* Capable of handling large-scale data

The framework is a fast and high-performance gradient boosting one based on decision tree algorithms, used for ranking, classification and many other machine learning tasks. It was developed under the Distributed Machine Learning Toolkit Project of Microsoft.

Since the LightGBM is based on decision tree algorithms, it splits the tree leaf wise with the best fit whereas other boosting algorithms split the tree depth wise or level wise rather than leaf-wise. So when growing on the same leaf in Light GBM, the leaf-wise algorithm can reduce more loss than the level-wise algorithm and hence results in much better accuracy which can rarely be achieved by any of the existing boosting algorithms.

Also, it is surprisingly very fast, hence the word ‘Light’.

Refer to the article to know more about LightGBM: https://www.analyticsvidhya.com/blog/2017/06/which-algorithm-takes-the-crown-light-gbm-vs-xgboost/

Python Code:

data = np.random.rand(500, 10) # 500 entities, each contains 10 features

label = np.random.randint(2, size=500) # binary target

train\_data = lgb.Dataset(data, label=label)

test\_data = train\_data.create\_valid('test.svm')

param = {'num\_leaves':31, 'num\_trees':100, 'objective':'binary'}

param['metric'] = 'auc'

num\_round = 10

bst = lgb.train(param, train\_data, num\_round, valid\_sets=[test\_data])

bst.save\_model('model.txt')

# 7 entities, each contains 10 features

data = np.random.rand(7, 10)

ypred = bst.predict(data)

R Code:

library(RLightGBM)

data(example.binary)

#Parameters

num\_iterations <- 100

config <- list(objective = "binary",  metric="binary\_logloss,auc", learning\_rate = 0.1, num\_leaves = 63, tree\_learner = "serial", feature\_fraction = 0.8, bagging\_freq = 5, bagging\_fraction = 0.8, min\_data\_in\_leaf = 50, min\_sum\_hessian\_in\_leaf = 5.0)

#Create data handle and booster

handle.data <- lgbm.data.create(x)

lgbm.data.setField(handle.data, "label", y)

handle.booster <- lgbm.booster.create(handle.data, lapply(config, as.character))

#Train for num\_iterations iterations and eval every 5 steps

lgbm.booster.train(handle.booster, num\_iterations, 5)

#Predict

pred <- lgbm.booster.predict(handle.booster, x.test)

#Test accuracy

sum(y.test == (y.pred > 0.5)) / length(y.test)

#Save model (can be loaded again via lgbm.booster.load(filename))

lgbm.booster.save(handle.booster, filename = "/tmp/model.txt")

If you’re familiar with the Caret package in R, this is another way of implementing the LightGBM.

require(caret)

require(RLightGBM)

data(iris)

model <-caretModel.LGBM()

fit <- train(Species ~ ., data = iris, method=model, verbosity = 0)

print(fit)

y.pred <- predict(fit, iris[,1:4])

library(Matrix)

model.sparse <- caretModel.LGBM.sparse()

#Generate a sparse matrix

mat <- Matrix(as.matrix(iris[,1:4]), sparse = T)

fit <- train(data.frame(idx = 1:nrow(iris)), iris$Species, method = model.sparse, matrix = mat, verbosity = 0)

print(fit)

**10.4. Catboost**

CatBoost is a recently open-sourced machine learning algorithm from Yandex. It can easily integrate with deep learning frameworks like Google’s TensorFlow and Apple’s Core ML.

The best part about CatBoost is that it does not require extensive data training like other ML models, and can work on a variety of data formats; not undermining how robust it can be.

Make sure you handle missing data well before you proceed with the implementation.

Catboost can automatically deal with categorical variables without showing the type conversion error, which helps you to focus on tuning your model better rather than sorting out trivial errors.

Learn more about Catboost from this article: https://www.analyticsvidhya.com/blog/2017/08/catboost-automated-categorical-data/

Python Code:

import pandas as pd

import numpy as np

from catboost import CatBoostRegressor

#Read training and testing files

train = pd.read\_csv("train.csv")

test = pd.read\_csv("test.csv")

#Imputing missing values for both train and test

train.fillna(-999, inplace=True)

test.fillna(-999,inplace=True)

#Creating a training set for modeling and validation set to check model performance

X = train.drop(['Item\_Outlet\_Sales'], axis=1)

y = train.Item\_Outlet\_Sales

from sklearn.model\_selection import train\_test\_split

X\_train, X\_validation, y\_train, y\_validation = train\_test\_split(X, y, train\_size=0.7, random\_state=1234)

categorical\_features\_indices = np.where(X.dtypes != np.float)[0]

#importing library and building model

from catboost import CatBoostRegressormodel=CatBoostRegressor(iterations=50, depth=3, learning\_rate=0.1, loss\_function='RMSE')

model.fit(X\_train, y\_train,cat\_features=categorical\_features\_indices,eval\_set=(X\_validation, y\_validation),plot=True)

submission = pd.DataFrame()

submission['Item\_Identifier'] = test['Item\_Identifier']

submission['Outlet\_Identifier'] = test['Outlet\_Identifier']

submission['Item\_Outlet\_Sales'] = model.predict(test)

R Code:

set.seed(1)

require(titanic)

require(caret)

require(catboost)

tt <- titanic::titanic\_train[complete.cases(titanic::titanic\_train),]

data <- as.data.frame(as.matrix(tt), stringsAsFactors = TRUE)

drop\_columns = c("PassengerId", "Survived", "Name", "Ticket", "Cabin")

x <- data[,!(names(data) %in% drop\_columns)]y <- data[,c("Survived")]

fit\_control <- trainControl(method = "cv", number = 4,classProbs = TRUE)

grid <- expand.grid(depth = c(4, 6, 8),learning\_rate = 0.1,iterations = 100, l2\_leaf\_reg = 1e-3,            rsm = 0.95, border\_count = 64)

report <- train(x, as.factor(make.names(y)),method = catboost.caret,verbose = TRUE, preProc = NULL,tuneGrid = grid, trControl = fit\_control)

print(report)

importance <- varImp(report, scale = FALSE)

print(importance)