



# **Lecture Notes**

# Support Vector Machine

Support Vector Machine (SVM) is an advanced machine learning technique which has a unique way of solving complex problems such as image recognition, face detection, voice detection etc. As you will learn in this session, SVMs solves the Pioi;; 98oblem of nonlinearity through kernels.

For instance, if you have a data as shown in the figure below, SVMs can handle it easily and that's how SVM distinguishes from logistic regression.

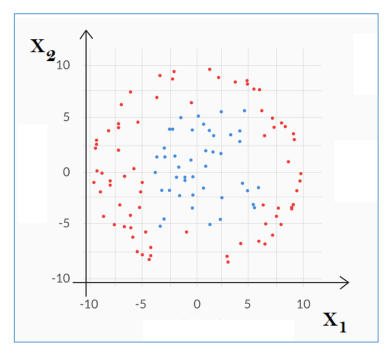


Figure 1: Nonlinear Data

It is important to remember that SVMs belong to the class of linear machine learning models (logistic regression is also a linear model).

A linear model uses a linear function (i.e. of the form y = ax + b) to model the relationship between the input x and output y. For example, in logistic regression, the log(odds) of an outcome (say, defaulting on a credit card) is linearly related to the attributes x1, x2, etc.

$$log(odds\ of\ default) = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \dots \beta_n X_n$$

Similarly, SVMs are also linear models and It needs attributes in the numeric form.





## Concept of Hyperplane in 2D

Before you move on to support vector machines, you need to understand the concept of hyperplanes. Essentially, it is a boundary which dassifies the data set (classifies Spam email from the ham ones). It could be lines, 2D planes, or even n-dimensional planes that are beyond our imagination.

A line that is used to classify one dass from another is called a hyperplane. In fact, it is the model you're trying to build as shown in the figure below:

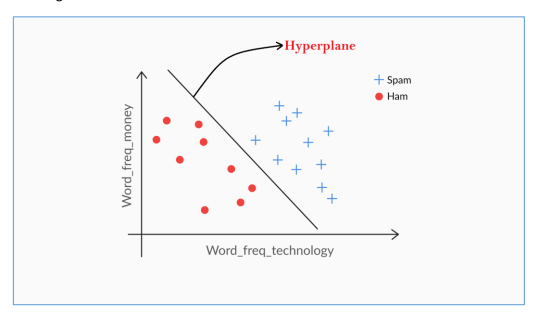


Figure 2: Hyperplane

The standard equation of a line is given by ax + by + c = 0. You could generalise it as W<sub>0</sub> + W<sub>1</sub>X<sub>1</sub> + W<sub>2</sub>X<sub>2</sub>=0, where X<sub>1</sub> and X<sub>2</sub> are the features — such as 'word\_freq\_technology' and 'word\_freq\_money' — and W<sub>1</sub> and W<sub>2</sub> are the coefficients.

For any line with W coefficients, substituting the value of features x1 and x2 in the equation of the line determined by its W coefficients, will return a value.

A **positive value** (**blue points** in the plot above) would mean that the set of values of the features is in one class; however, a **negative value** (**red points** in the plot above) would imply it belongs to the other class. A value of zero would imply that the point lies on the line (hyperplane) because any point on the line will satisfy the equation:  $W_0 + W_1X_1 + W_2X_2 = 0$ .

## Concept of Hyperplane in 3D

In 3 dimensions (refer to figure below), the hyperplane (light orange) will be a plane with an expression of **ax+by+cz+d = 0**. The plane divides the data set into two halves. Data points above the plane represent one class (red), while data points below the plane represent the other class (blue).





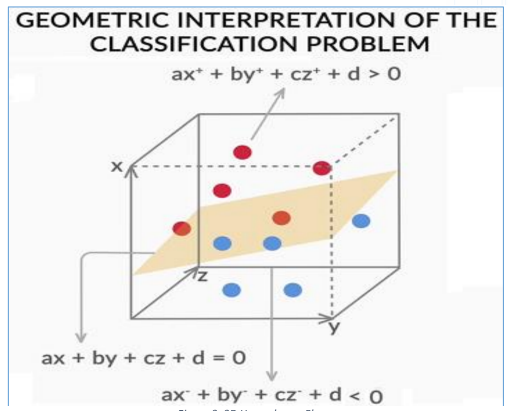


Figure 3: 3D Hyperplane - Plane

In general, if the hyperplane from d attributes in d-dimensional, the expression can be written as follows:

$$\sum_{i=1}^d (W_i.\,X_i+W_0)=0$$

The model denoted by the expression given above is called **a linear discriminator**. Similar to the 2D and 3D expressions, an n-dimensional hyperplane also follows the general rule: all points above the plane will yield a value **greater than 0**, and those below it will yield **lesser than 0** when plugged into the expression of the hyperplane.

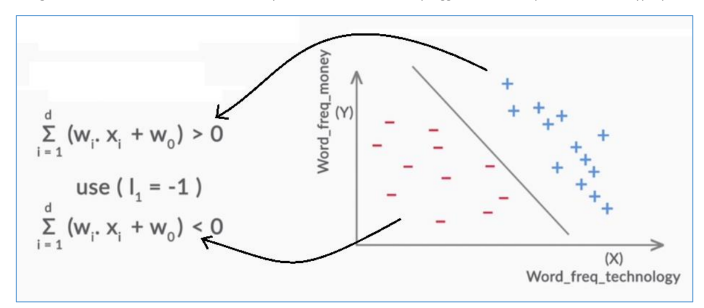


Figure 4: Linear Discriminator





# **Maximal Margin Classifier**

There could be multiple lines(Hyperplanes) possible which perfectly separate the two classes as shown in the figure below. But the best line, is the one which maintains the largest possible equal distance from the nearest points of both the classes so for the separator to be optimal, the margin or the distance of the nearest point to the separator should be maximum. This is called **Maximal Margin classifier**.

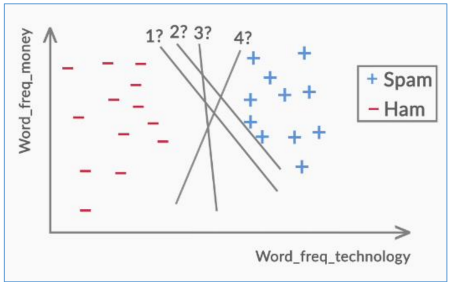


Figure 5: Maximal Margin Classifier

3rd line(Hyperplane) should be considered as a maximal margin dassifier in the above figure.

**Mathematical formulation** for maximal margin classifier requires two major constraints that need to be taken into account while maximising the margin. They are

• The standardisation of coefficients such that the summation of the square of the coefficients of all the attributes is equal to 1. For example, if you have 20 attributes, then the summation of the coefficients should be:

$$\sum_{i=0}^{20} (W_i^2) = 1$$

 Along with the first constraint, the maximal margin hyperplane should also follow the constraint given below:

$$(l_iX(W_i,Y_i))\geqslant M$$
 where,  $l_i$  = label (1, -1)  $W_i$  = coefficient of attributes  $Y_i$  = data points of all the attributes in each row





The maximal margin line (hyperplane), although it separates the two classes perfectly, is very sensitive to the **training data**. This means that **the Maximal Margin Classifier** will perform perfectly on the training data set. But on the unseen data, it may perform poorly. Also, there are cases where the classes cannot be perfectly separated.

Thus, the **soft margin classifier** helps in solving this problem.

## **Soft Margin Classifier**

The **Support Vector Classifier** essentially allows certain points to be deliberately misclassified. By doing this, it is able to classify most of the points correctly in the unseen data and is also more robust.

The Support Vector Classifier is also called the **Soft Margin Classifier** because instead of searching for the margin that exactly classifies each and every data point to the correct dass, the Soft Margin Classifier allows some observations to fall on the wrong side. The points which are close to the hyperplane are only considered for constructing the hyperplane and those points are called **support vectors**.

Support vector dassifier works well when the data is partially intermingled (i.e. the data can be classified by minimal misdassifications). But what if the distribution looks completely intermingled and follows some pattern, something like the circular distribution of labels (+ and -), as shown in figure below.

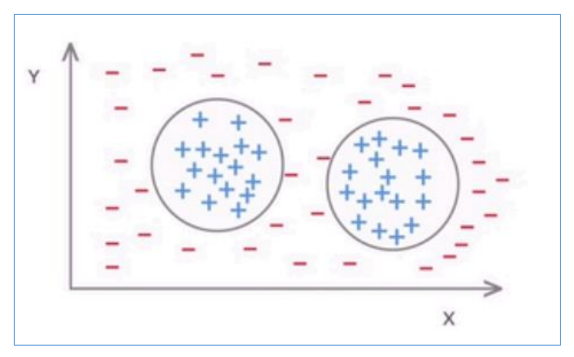


Figure 6: Intermingled Data

Obviously, the **Support Vector Classifier** can't classify the data above correctly, because it divides the data set into two halves, which misclassifies a lot of data points. But it doesn't mean that this problem cannot be solved. There is a way to solve such problems, which we will learn later.

Like the Maximal Margin Classifier, the **Support Vector Classifier** also maximises the margin; but the margin, here, will allow some points to be misdassified, as shown in figure below.





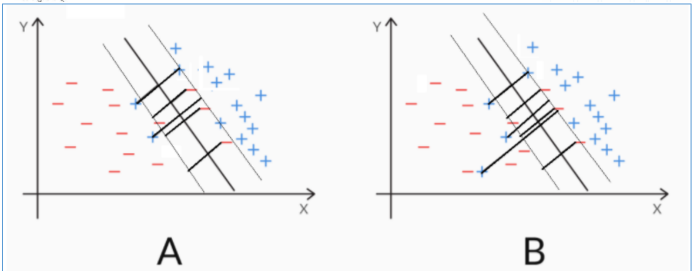


Figure 7: Soft Margin Classifier

So to select **the best-fit Support Vector Classifier**, the notion of slack variables (epsilons( $\epsilon$ )) can help in comparing the classifiers.

There is also a concept of the slack variable ( $\epsilon$ ). A slack variable is used to control misclassifications. It tells you where an observation is located relative to the margin and hyperplane.

There are three different conditions applied if any new data point comes into play. Suppose you draw a Support Vector Classifier in such a way that it doesn't allow any misdassification, i.e. Epsilon( $\epsilon$ ) = 0, then each observation is on the correct side of the margin as shown in figure below.

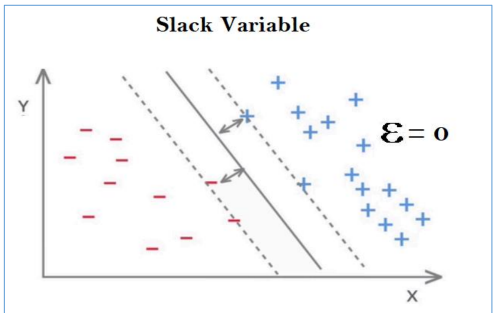


Figure 8: Slack Variable

But if you draw a Support Vector Classifier in such a way that it only violates the margin, i.e.  $0 < Epsilon(\epsilon) < 1$ , the observations classify correctly as shown in figure below.





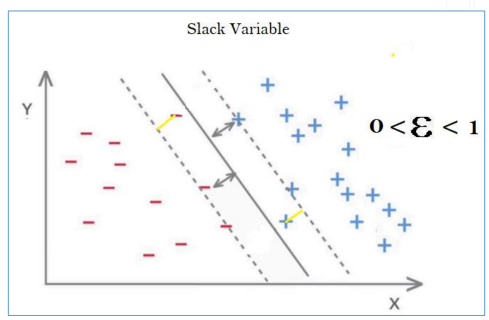


Figure 9: Slack Variables

But if the data points violate the hyperplane, i.e. Epsilon( $\epsilon$ ) > 1, then the observation is on the wrong side of the hyperplane, as shown in figure below.

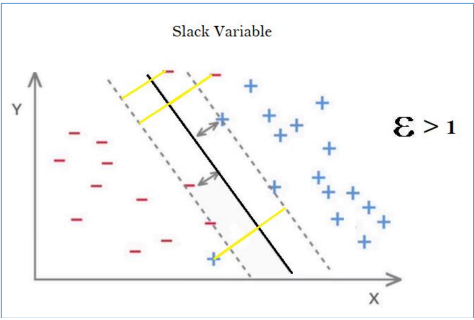


Figure 10: Slack Variable

### So you can see that:

- Each data point has a slack value associated to it, according to where the point is located.
- The value of slack lies between 0 and +infinity.

Lower values of slack are better than higher values (slack = 0 implies a correct dassification, but slack > 1 implies an incorrect classification, whereas slack within 0 and 1 classifies correctly but violates the margin)





#### **Cost of Misclassification**

Cost of misdassification is greater than or equal to the summation of all the epsilons of each data point, and is denoted by cost or 'C'.

$$\sum \epsilon_i \leq C$$
.

Once you understand the **notion of the slack variable**, you can easily compare the two **Support Vector Classifiers**. You can measure the summation of all the epsilons( $\epsilon$ ) of both the hyperplanes and choose the best one that gives you the least sum of epsilons( $\epsilon$ ). The summation of all the epsilons of each data point is denoted by cost or 'C', i.e.

When **C** is large, the slack variables can be large, i.e. you allow a larger number of data points to be misclassified or to violate the margin. So you get a hyperplane where the margin is wide and misclassifications are allowed. In this case, the model is flexible, more generalisable, and less likely to overfit. In other words, it has a high bias.

On the other hand, when **C** is small, you force the individual slack variables to be small, i.e. you do not allow many data points to fall on the wrong side of the margin or the hyperplane. So, the margin is narrow and there are few misclassifications. In this case, the model is less flexible, less generalisable, and more likely to overfit. In other words, it has a high variance.

#### **Kernels**

Kernels are one of the most interesting inventions in machine learning, partly because they were born through the creative imagination of mathematicians, and partly because of their utility in **dealing with non-linear datasets** 

So far, we have learnt about hyperplanes, the Maximal Margin Classifier, and the Support Vector Classifier. All of these are linear models (since they use linear hyperplanes to separate the classes). However, many real-world data sets **are not separable by linear boundaries**. For instance, what if the distribution of data points looks like the figure given below?

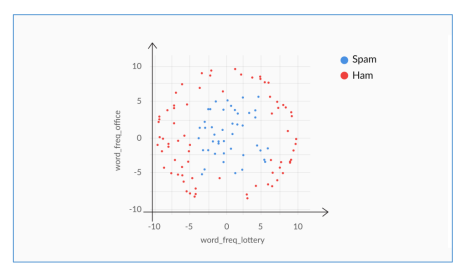


Figure-11: Intermingled Data





You'll agree that it is not possible to imagine a linear hyperplane (a line in 2D) that separates the red and blue points reasonably well. Thus, you need to tweak the linear SVM model and enable it to incorporate nonlinearity in some way. Kernels serve this purpose — they enable the linear SVM model to separate nonlinearly separable data points.

## **Mapping Nonlinear Data to Linear Data**

You can transform nonlinear boundaries to linear boundaries by applying certain functions to the original attributes. The original space (X, Y) is called the **attribute space**, and the transformed space (X', Y') is called the **feature space**.

Let's say you want to classify emails into 'spam' or 'ham' on the basis of two attributes — 'word\_freq\_office' (X) and 'word\_freq\_lottery' (Y). The following plot shows the data set below, which is dearly nonlinear

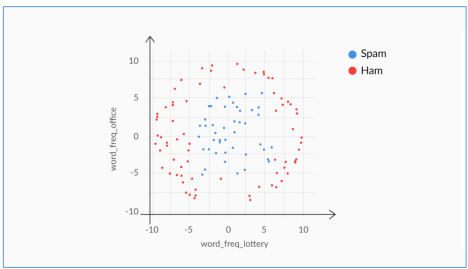


Figure 12: Nonlinear Distribution

To convert this data set into a linearly separable one, a simple transformation into a new feature space (X', Y') can be made. For now, don't worry about the math behind the transformation. You may almost never need to manually transform data sets. Just assume that some appropriate transformation from (X, Y) to (X', Y') can make the data linearly separable.

In the original attribute space, notice that the observations are distributed in a circular fashion. This gives you a hint that the transformation should convert the circular distribution to a linear distribution.

- word\_freq\_office(X')=(word\_freq\_office(X)-a)<sup>2</sup>
- word\_freq\_office'(Y')=(word\_freq\_office(Y)-b)<sup>2</sup>

#### **Feature Transformation**

The process of transforming the original attributes into a new feature space is called 'feature transformation'. However as the number of attributes increases, there is an exponential increase in the number of dimensions in the transformed feature space. Suppose you have four variables in your data set, then considering only a polynomial transformation with degree 2, you end up making **15 features** in the new feature space, as shown in the figure below.





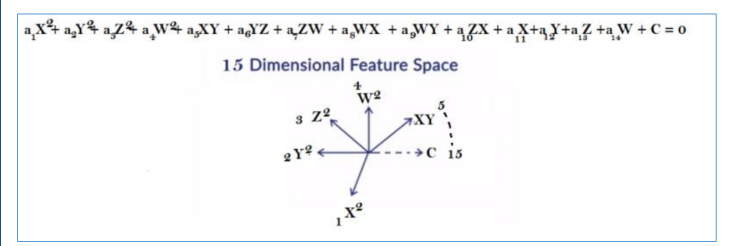


Figure 13: 4D Feature Space

### **Kernel Trick**

As feature transformation results in large number of features, it makes the modelling (i.e. the learning process) computationally expensive. The use of kernel resolves this issue. The key fact that makes the kernel trick possible is that to find a best fit model, the learning algorithm only needs the inner products of the observations  $(X_i^T X_j)$ . It never uses the individual data points X1, X2 etc. in silo.

Think of a kernel as a **black box**, as shown in the figure below. The attributes are passed on the black box, and it returns the linear boundaries for the classification of the nonlinear data implicitly.

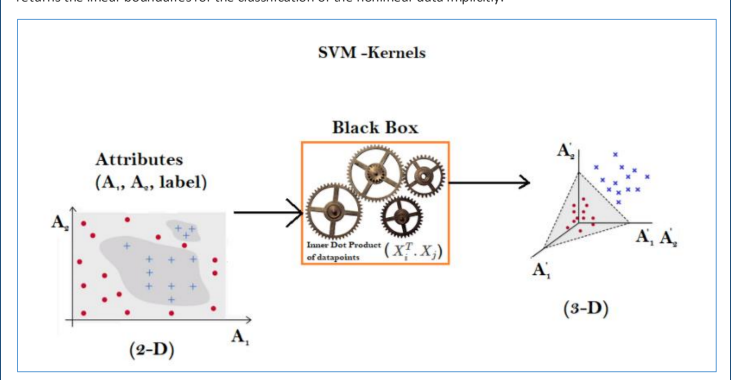


Figure 14: Black Box





Kernel functions use this fact to bypass the explicit transformation process from the attribute space to the feature space, and rather do it implicitly. The benefit of implicit transformation is that now you do not need to:

- Manually find the mathematical transformation needed to convert a nonlinear to a linear feature space
- Perform computationally heavy transformations

In practice, you only need to know that kernels are functions which help you transform non-linear datasets. Given a dataset, you can try various kernels, and choose the one that produces the best model. The **three most popular** types of kernel functions are:

- **The linear kernel**: This is the same as the support vector classifier, or the hyperplane, without any transformation at all
- The polynomial kernel: It is capable of creating nonlinear, polynomial decision boundaries
- The radial basis function (RBF) kernel: This is the most complex one, which is capable of transforming highly nonlinear feature spaces to linear ones. It is even capable of creating elliptical (i.e. endosed) decision boundaries

The three types of kernel functions shown in the figures below represent the typical decision boundaries they are capable of creating. Notice that the linear kernel is same as the vanilla hyperplane, the polynomial kernel can produce polynomial shaped nonlinear boundaries, and the RBF kernel can produce highly nonlinear, ellipsoid shaped boundaries.

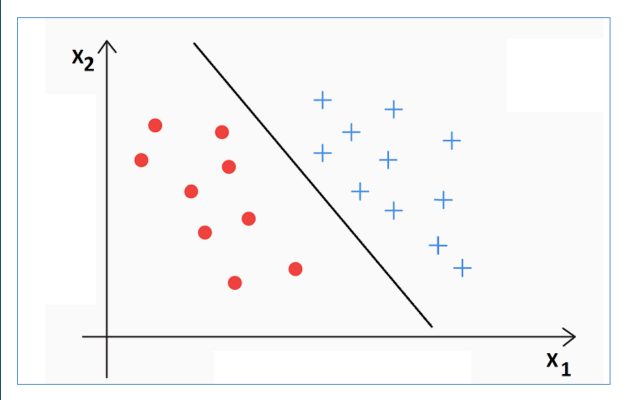


Figure 15:Linear Kernel (Hyperplane)





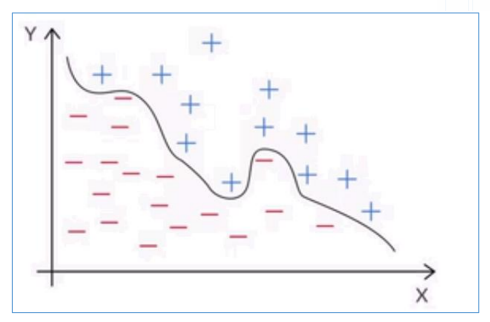


Figure 16:Polynomial Kernel

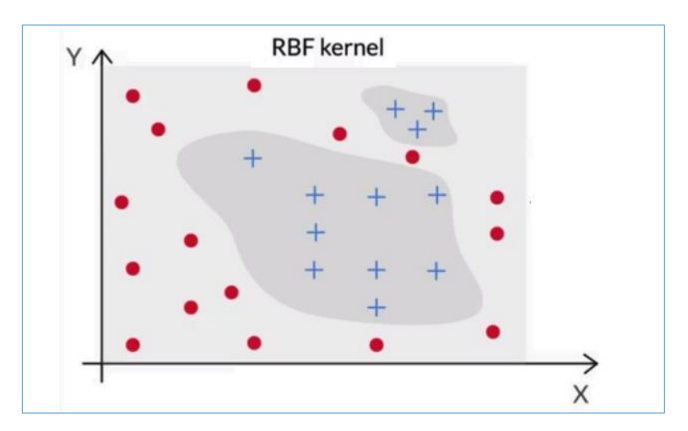


Figure 17: RBF Kernel

# **Kernel Parameter**

In nonlinear kernels such as the RBF, you use the parameter gamma/sigma to control the amount of nonlinearity in the model. The higher the value of sigma, the more is the nonlinearity introduced; the lower the value of gamma/sigma, the lesser is the nonlinearity. It is also denoted as gamma in some texts and packages. Apart from sigma, you also have the hyperparameter C, or the cost (with all types of kernels).





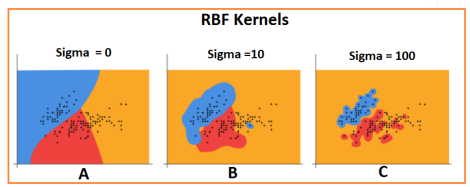


Figure 18: RBF Kernels

The plot above shows three RBF kernels with different values of sigma. When sigma is high, more nonlinearity is added. When you increase the sigma from 10 to 100, the nonlinearity is further increased, and all the training points are correctly mapped, as is visible in the highly complex decision boundaries in the figure 1(C). However, this results in a highly biased model that may overfit the training set. Like most other hyperparameters, it is advisable to tune the value of sigma using cross-validation.

Choosing the appropriate kernel is important for building a model of optimum complexity. If the kernel is highly nonlinear, the model is likely to overfit. On the other hand, if the kernel is too simple, then it may not fit the training data well.

Usually, it is difficult to choose the appropriate kernel by visualising the data or using exploratory analysis. Thus, cross-validation (or hit-and-trial, if you are only choosing from 2-3 types of kernels) is often a good strategy.