Support Vector Machines

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1 Introduction

This note is about the Support Vector Machines (SVM), a method that has exhibited excellent accuracy in classification problems. Complex mathematics are involved in various aspects of SVM, e.g., proof of its generalization bounds, its optimization, designing and proving the validity of various non-linear kernels, etc. We will, however, not focus on the mathematics. The main purpose of this note is to introduce how the ideas in SVM are formulated, why are they reasonable, and how various simplifications are useful in shaping the SVM primal form.

We will not touch any generalization bound of SVM, although that is an area which have attracted intensive research efforts. We will not talk about how the optimization problem in SVM can be solved or approximately solved (for efficiency and scalability). These choices enable us to focus on the key ideas that lead to SVM, and the strategies in SVM that may be helpful in other domains, and we encourage the readers to also pay attention to these aspects of SVM.

2 The key SVM idea

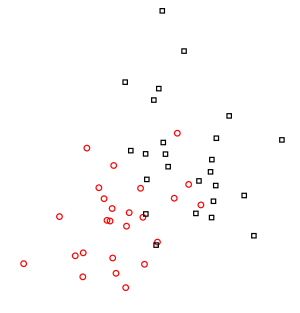
In SVM, we do not model either the generative or discriminative distributions of the training data. Instead, SVM seeks to directly find the best classification boundary which divides data points in different classes. The natural question is: which classification boundary is considered good or even best?

2.1 Simplify it, simplify it!

One key to answer this question is actually to simplify the problem. The caption says "simplify it" triple times—it is not only because it is "important" (so must be repeated three times), but also we are making three simplifying assumptions.

Given datasets (or classification problems) such as those appearing in Figure 1, it is difficult to figure out which classifier is good. In Figure 1a, the two classes (whose examples are denoted by squares and circles, respectively) cannot be classified by any linear boundary. Using a complex curve (i.e., non-linear boundary), we can separate these two classes with zero training error. However, it is not obvious either how to compare two complex non-linear classification boundaries. Hence, we are making the first two assumptions: linear boundary (or, linear classifier) and separable (i.e., linear classifiers can achieve 100% accuracy on the training set.) These two are usually combined to request a linearly separable problem.

The problem in Figure 1b is in fact linearly separable, in which the four classes can be divided into non-overlapping regions by linear boundaries, e.g., three lines that enclose the red class examples. However, it is not easy to



(a) Inseparable data are messing

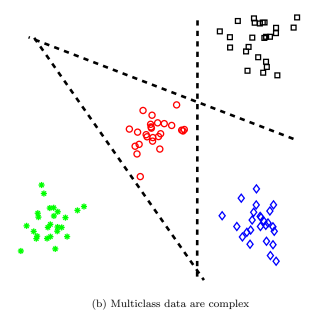


Figure 1: Illustration of complex classification problems.

judge whether one set of triple-lines are better than another triplet either. And, there are many other possible linear boundaries beyond triplets enclosing red examples. Hence, we make the third assumption: the problem is *binary*.

Putting all assumptions together, we start by only considering linearly separable binary classification problems. We want to emphasize that these simplifications are reasonable, i.e., they will not change the essence of the classification problem. As we will introduce in this note, all these assumptions will be relaxed and taken care of in SVM. Hence, reasonable simplifying assumptions help us find ideas and good solutions to a complex problem, and we have the option to re-consider these assumptions later. However, if an assumption changes some basic characteristics of our problem, it is better avoided.

2.2 Find a max (or large) margin classifier

Figure 2a shows a linearly separable binary problem. It is clear that any line between the cluster of red circles and black squares can separate them perfectly. In Figure 2a, three example boundaries are illustrated: the blue solid line, the red dashed line, and the black dotted line.

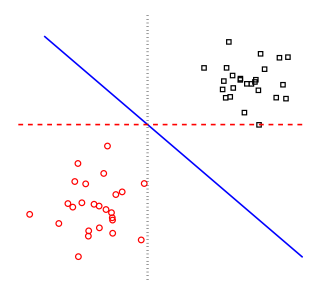
Furthermore, it becomes easier to determine which classifier is better (or the best). Most people will agree that the classification boundary determined by the blue solid line is better than the other two. The other two boundaries are too close to the examples, leaving very small (almost no) room for variations or noise. The examples in Figure 2a and 2b are generated using the same underlying distribution and contain the same number of training examples. However, because of the randomness, two red circles appear on the right side of the black dotted line, and one black square appears below the red dashed line in Figure 2b. These examples are errors for the two boundaries, respectively.

The blue solid line, however, correctly classified all examples in Figure 2b. The reason for this robustness is: this boundary is far away from all training examples; hence, when variations or noise are in effect, this classifier has the *margin* to accommodate for these changes since the scale of such changes are usually small. As shown in Figure 2, the margin of a classifier is the distance from it to the training examples which are the closest to it.

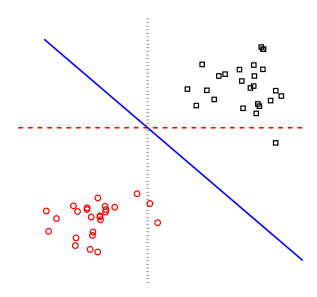
Hence, the margin of one example (with respect to a classification boundary) is the distance from that point to the boundary. Naturally, the margin of a dataset is the minimum of the margin of all its samples.

Because large margin is beneficial for classification, some classifiers directly maximize the margin, which are called *max-margin classifiers*. Some classifiers seek the compromise between large margin and other properties, and are called *large margin classifiers*. SVM is a max-margin classifier.

And, of course, the first task after having the max-margin idea is: how shall we change the max-margin idea into operational procedures? We need to first formalize the max-margin idea.



(a) Large vs. small margin boundaries



(b) Small margin can cause troubles

Figure 2: Illustration of large margin idea.

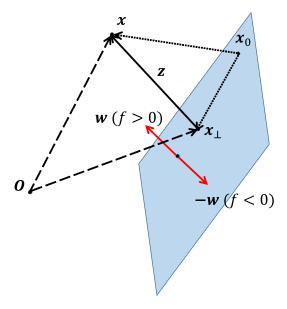


Figure 3: Illustration of projection, margin, and normal vector.

3 Visualize and calculate the margin

Given a linear classification boundary and a point x, how do we calculate the margin of x? We know in a d-dimensional space, a linear classification boundary is a hyperplane. An illustrated such as Figure 3 is helpful.

3.1 Visualizing the geometry

We have two ways to specify a vector \boldsymbol{x} in an illustration. One way is to draw an arrow from the origin (O) to the coordinates specified by the elements of \boldsymbol{x} . We use this way to draw \boldsymbol{x} , \boldsymbol{x}_{\perp} and \boldsymbol{x}_0 in Figure 3 (denoted by dashed arrows, but the arrow for \boldsymbol{x}_0 is omitted to keep the illustration's clarity). Another way is to use two end points to specify a vector, such that the vector is the subtraction of the two points. For example, $\boldsymbol{z} = \boldsymbol{x} - \boldsymbol{x}_{\perp}$ (denoted by the black solid arrow) and two vectors involving \boldsymbol{x}_0 (denoted by dotted arrows). \boldsymbol{w} and $-\boldsymbol{w}$ are also specified in this way.

All points in a hyperplane in the d-dimensional space is specified by an equation $f(\boldsymbol{x}) = \boldsymbol{w}^T \boldsymbol{x} + b = 0$, where \boldsymbol{w} determines the direction of the hyperplane. The direction of \boldsymbol{w} (i.e., $\frac{\boldsymbol{w}}{\|\boldsymbol{w}\|}$) is called the normal vector of the hyperplane. A hyperplane is perpendicular to its normal vector. In other words, if \boldsymbol{x}_1 and \boldsymbol{x}_2 are two different points on the hyperplane (i.e., satisfying $f(\boldsymbol{x}_1) = f(\boldsymbol{x}_2) = 0$ and $\boldsymbol{x}_1 \neq \boldsymbol{x}_2$), we always have $\boldsymbol{w}^T(\boldsymbol{x}_1 - \boldsymbol{x}_2) = 0$. If \boldsymbol{x} is on the hyperplane, then $f(\boldsymbol{x}) = 0$; if \boldsymbol{x} is not on the hyperplane, but in the same direction of \boldsymbol{w} with respect to the hyperplane, then $f(\boldsymbol{x}) > 0$; and, $f(\boldsymbol{x}) < 0$ holds for points in the

opposite direction (cf. the red solid arrows).

In Figure 3, x_{\perp} is the projection of x onto the hyperplane. That is, x is decomposed as $x = x_{\perp} + z$. Given any point $x_0 \neq x_{\perp}$ on the hyperplane, we always have $z \perp (x_{\perp} - x_0)$ (However, x_{\perp} is not perpendicular to z if $b \neq 0$). This geometry tells us that $\|z\|$ is the distance we are looking for. Hence, a visualization helps us sort out the description "distance" into precise mathematics.

3.2 Calculate the margin as an optimization

What is even better, the visualization also hints us on how to calculate the distance. Because $z \perp (x_{\perp} - x_0)$, we know $||z|| \leq ||x - x_0||$ for any x_0 on the hyperplane, because of the Pythagorean theorem. That is, x_{\perp} is the solution of the following optimization problem:

$$\underset{\boldsymbol{y}}{\operatorname{arg\,min}} \quad \|\boldsymbol{x} - \boldsymbol{y}\|^{2} \tag{1}$$
 s.t. $f(\boldsymbol{y}) = 0$.

$$s.t. \quad f(\mathbf{y}) = 0. \tag{2}$$

This is an optimization problem with one equality constraint. The Lagrangian is $L(\boldsymbol{y}, \lambda) = \|\boldsymbol{x} - \boldsymbol{y}\|^2 - \lambda(\boldsymbol{w}^T \boldsymbol{y} + b)$. Setting $\frac{\partial L}{\partial \boldsymbol{y}} = 0$ gives $2(\boldsymbol{y} - \boldsymbol{x}) = \lambda \boldsymbol{w}$. That is, $\mathbf{y} = \mathbf{x} + \frac{\lambda}{2}\mathbf{w}$. Plugging it into $f(\mathbf{y}) = 0$, we get $\frac{\lambda}{2}\mathbf{w}^T\mathbf{w} + f(\mathbf{x}) = 0$, hence $\lambda = -\frac{2f(x)}{w^Tw}$. Then, the projection is $x_{\perp} = y^{\star} = x + \frac{\bar{\lambda}}{2}w = x - \frac{f(x)}{w^Tw}w$.

Now, $z = x - x_{\perp} = \frac{f(x)}{w^T w} w$, hence, the distance (and the margin of x) is

$$\left\| \frac{f(\boldsymbol{x})}{\boldsymbol{w}^T \boldsymbol{w}} \boldsymbol{w} \right\| = \left| \frac{f(\boldsymbol{x})}{\|\boldsymbol{w}\|^2} \right| \|\boldsymbol{w}\| = \frac{|f(\boldsymbol{x})|}{\|\boldsymbol{w}\|}.$$
 (3)

There are simpler ways to find out the margin than what is presented here. However, in this section we familiarized ourselves with the geometry useful for SVM, found the actual value of x_{\perp} , exercised the process of formalization and optimization, and applied the method of Lagrange multipliers. These are all fruitful outcomes.

4 Maximize the margin

SVM tries to maximize the margin of the dataset, i.e., the smallest margin of all training points.

4.1 The formalization

Let us denote the training set as $\{(\boldsymbol{x}_i, y_i)\}_{i=1}^n$, where $\boldsymbol{x}_i \in \mathbb{R}^d$ is an example, and $y \in \mathcal{Y}$. For a binary problem, $\mathcal{Y} = \{1, 2\}$. To maximize the margin of this data with respect to a linear boundary $f(x) = w^T x + b$ means to solve the following optimization problem:

$$\max_{\boldsymbol{w},b} \quad \min_{1 \le i \le n} \frac{|f(\boldsymbol{x}_i)|}{\|\boldsymbol{w}\|}, \tag{4}$$

with the constraint f(x) can classify all training examples correctly. Hence, we also need to mathematically describe "correctly classified."

That is quite simple if we change a little bit the definition of \mathcal{Y} . If we set $\mathcal{Y} = \{+1, -1\}$, the meaning of it will not change—the two numbers refers to the label of the two classes, and the specific value of them are not relevant (e.g., 2 vs. -1). However, with this slight change, the following statement holds:

$$f(\mathbf{x})$$
 correctly classifies all training examples, iff $y_i f(\mathbf{x}_i) > 0$ for all $1 \le i \le n$, (5)

in which "iff" means if and only if. When $y_i = 1$ (i.e., a positive example), $y_i f(\mathbf{x}_i) > 0$ means $f(\mathbf{x}_i) > 0$, thus the prediction is also positive; When $y_i = -1$, $f(x_i) < 0$ and it predicts x_i as negative.

SVM then optimizes the following problem for a binary linearly separable dataset:

$$\max_{\boldsymbol{w},b} \quad \min_{1 \le i \le n} \frac{|f(\boldsymbol{x}_i)|}{\|\boldsymbol{w}\|} \tag{6}$$

s.t.
$$y_i f(x_i) > 0, 1 \le i \le n.$$
 (7)

4.2 The simplifications

The constraints can be handled by a Lagrangian. However, the objective contains fraction, absolute value, vector norm and the maximization of a minimum, all of which are hostile to optimization. By looking at this objective more carefully, there exist ways to shun away from all these difficulties.

Our assumption ensures that $y_i f(\mathbf{x}_i) > 0$ for all $1 \le i \le n$, that is, $y_i f(\mathbf{x}_i) =$ $|f(x_i)|$ because $|y_i|=1$. Hence, the objective can be rewritten as

$$\min_{1 \le i \le n} \frac{|f(\boldsymbol{x}_i)|}{\|\boldsymbol{w}\|} = \min_{1 \le i \le n} \frac{y_i f(\boldsymbol{x}_i)}{\|\boldsymbol{w}\|} = \frac{1}{\|\boldsymbol{w}\|} \min_{1 \le i \le n} \left(y_i (\boldsymbol{w}^T \boldsymbol{x}_i + b) \right). \tag{8}$$

This type of objective (as a ratio) has appeared many times (e.g., in PCA). If (\mathbf{w}^*, b^*) is a maximizer of the above objective, so will be $(c\mathbf{w}^*, cb^*)$ for any nonzero constant c. However, since this is a constrained optimization problem, $c \leq 0$ will make all the constraints invalid. When c > 0, $(c\boldsymbol{w}^*, cb^*)$ will not change the objective and all constraints remain satisfied. That is, we have the freedom to choose any c > 0. In the past, we have chosen a c such that $\|\boldsymbol{w}\| = 1$. If we use this same assumption, the optimization becomes

$$\max_{\boldsymbol{w},b} \quad \min_{1 \le i \le n} y_i f(\boldsymbol{x}_i)$$
s.t. $y_i f(\boldsymbol{x}_i) > 0, \quad 1 \le i \le n$ (10)

s.t.
$$y_i f(x_i) > 0, 1 \le i \le n$$
 (10)

$$\boldsymbol{w}^T \boldsymbol{w} = 1. \tag{11}$$

The objective is still a maximization of a minimum.

However, note that $y_i f(x_i)$ appears in both the objective and the constraints, there is a clever trick to further simplify it. If $(\boldsymbol{w}^{\star}, b^{\star})$ is a maximizer of the original objective $\min_{1 \leq i \leq n} \frac{|f(\boldsymbol{x}_i)|}{\|\boldsymbol{w}\|}$, we choose $c = \min_{1 \leq i \leq n} y_i((\boldsymbol{w}^*)^T \boldsymbol{x} + b^*)$. Obviously c > 0, hence $\frac{1}{c}(\boldsymbol{w}^*, b^*)$ is also a maximizer of the original problem. That is, we can add the following constraints without changing the optimal objective value:

$$\min_{1 \le i \le n} y_i(\boldsymbol{w}^T \boldsymbol{x}_i + b) = 1, \qquad (12)$$

or, equivalently, 1

$$y_i(\boldsymbol{w}^T\boldsymbol{x}_i + b) > 1(>0), \tag{13}$$

for all $1 \le i \le n$ —which means all constraints in the original problem are automatically satisfied! In other words, we can convert the original problem to the following equivalent one:

$$\max_{\boldsymbol{w},b} \quad \frac{\min_{1 \le i \le n} y_i(\boldsymbol{w}^T \boldsymbol{x} + b)}{\|\boldsymbol{w}\|} = \frac{1}{\|\boldsymbol{w}\|}$$
 (14)

s.t.
$$y_i f(x_i) \ge 1$$
, $1 \le i \le n$. (15)

A final step is to change max $\frac{1}{\|\boldsymbol{w}\|}$ to min $\|\boldsymbol{w}\|$ and furthermore (equivalently) $\boldsymbol{w}^T\boldsymbol{w}$ and $\frac{1}{2}\boldsymbol{w}^T\boldsymbol{w}$, which leads to

$$\min_{\boldsymbol{w},b} \quad \frac{1}{2} \boldsymbol{w}^T \boldsymbol{w} \tag{16}$$

s.t.
$$y_i f(\mathbf{x}_i) \ge 1$$
, $1 \le i \le n$. (17)

The additional term $\frac{1}{2}$ will make later derivations easier.

In this series of transformations and simplifications, we get a series of equivalent problems and hence will get the same optimal solutions as that of the original problem. The new constraints are very similar to the original ones (and are handled by the method of Lagrange multipliers in the same manner). However, the objective is now a quadratic form $(\boldsymbol{w}^T\boldsymbol{w})$, and very amenable to optimization.

We will work (a little bit) on the optimization of Equations (16–17), and they are called the primal form of the SVM formulation, or simply a primal SVM.

5 The Optimization and the solution

We use the method of Lagrange multipliers to handle the n inequality constraints and states the necessary and sufficient conditions for the minimizers of the primal SVM. The proof is beyond the scope of this note and is omitted. With these conditions, the primal form turns into an equivalent dual form.

To be precise, $\min_{1 \leq i \leq n} y_i(\boldsymbol{w}^T \boldsymbol{x}_i + b) = 1$ is not equivalent to $y_i(\boldsymbol{w}^T \boldsymbol{x}_i + b) \geq 1$ for all i. When $y_i(\boldsymbol{w}^T \boldsymbol{x}_i + b) \geq 1$, it is possible that $\min_{1 \leq i \leq n} y_i(\boldsymbol{w}^T \boldsymbol{x}_i + b) > 1$ (e.g., $\min_{1 \leq i \leq n} y_i(\boldsymbol{w}^T \boldsymbol{x}_i + b) = 2$. However, since in SVM the objective is to minimize $\frac{1}{2} \boldsymbol{w}^T \boldsymbol{w}$, $y_i(\boldsymbol{w}^T \boldsymbol{x}_i + b) \geq 1$ implies $\min_{1 \leq i \leq n} y_i(\boldsymbol{w}^T \boldsymbol{x}_i + b) = 1$ in the optimal solution.

5.1The Lagrangian and the KKT conditions

We still define one Lagrange multiplier α_i for each constraint, and the constraints $y_i f(x_i) \ge 1$ are rewritten as $y_i f(x_i) - 1 \ge 0$. The Lagrangian is

$$L(\boldsymbol{w}, b, \boldsymbol{\alpha}) = \frac{1}{2} \boldsymbol{w}^T \boldsymbol{w} - \sum_{i=1}^n \alpha_i \left(y_i(\boldsymbol{w}^T \boldsymbol{x}_i + b) - 1 \right)$$
(18)

s.t.
$$\alpha_i \ge 0$$
, $1 \le i \le n$, (19)

in which $\alpha = (\alpha_1, \alpha_2, \dots, \alpha_n)^T$ are the vector of Lagrange multipliers. For inequality constraints, however, the multipliers are not free any more, they are required to be non-negative. If one constraint is violated, i.e., if $y_i f(x_i) - 1 < 0$ for some i, we have $-\alpha_i (y_i(\mathbf{w}^T \mathbf{x}_i + b) - 1) > 0$, meaning a punishment term is added to $L(\boldsymbol{w}, b, \boldsymbol{\alpha})$. Hence, setting $\alpha_i \geq 0$ for all i is somehow enforcing the *i*-th constraint. Also, if

$$\alpha_i \left(y_i(\boldsymbol{w}^T \boldsymbol{x}_i + b) - 1 \right) = 0 \tag{20}$$

for all $1 \le i \le n$, then $L(\boldsymbol{w}, b, \boldsymbol{\alpha}) = \frac{1}{2} \boldsymbol{w}^T \boldsymbol{w}$ matches the primal objective. Later, we will show the optimality conditions indeed specifies $\alpha_i (y_i(\boldsymbol{w}^T\boldsymbol{x}_i+b)-1) =$ 0 for all i.

As usual, we compute the gradients and set them to $\mathbf{0}$ or 0, respectively:

$$\frac{\partial L}{\partial w} = \mathbf{0} \quad \Longrightarrow \quad \mathbf{w} = \sum_{i=1}^{n} \alpha_i y_i \mathbf{x}_i, \tag{21}$$

$$\frac{\partial L}{\partial b} = 0 \quad \Longrightarrow \quad \sum_{i=1}^{n} \alpha_i y_i = 0. \tag{22}$$

The above three equality conditions, the original inequality constraints, and the constraints on the Lagrange multipliers form the Karush-Kuhn-Tucker conditions for the primal SVM optimization problem:

$$\boldsymbol{w} = \sum_{i=1}^{n} \alpha_i y_i \boldsymbol{x}_i \tag{23}$$

$$\sum_{i=1}^{n} \alpha_i y_i = 0 \tag{24}$$

$$\alpha_i \left(y_i(\boldsymbol{w}^T \boldsymbol{x}_i + b) - 1 \right) = 0 \qquad i = 1, 2, \dots, n$$

$$\alpha_i \ge 0 \qquad i = 1, 2, \dots, n$$
(25)

$$\alpha_i \ge 0 \qquad i = 1, 2, \dots, n \tag{26}$$

$$y_i(\mathbf{w}^T \mathbf{x}_i + b) \ge 1$$
 $i = 1, 2, ..., n.$ (27)

The KKT conditions are not necessary and sufficient in general. However, for the primal SVM problem, these conditions are necessary and sufficient to specify optimal solutions.

Equation 23 states that the optimal solution \boldsymbol{w} can be represented by a weighted combination of the training examples; the sign of the weights are determined by the labels of examples, and the magnitude of these weights are the Lagrange multipliers. In more general large margin learning, the *representer theorem* ensures this weighted average representation is still valid in many other situations, e.g., kernel SVM.

Equation 24 states that these weights are balanced. If we sum the Lagrange multipliers for all positive training examples, it must equal the sum of weights for all negative training examples. Equation 25 is called the *complementary slackness* property, which is important in SVM and we will soon discuss this property in more depth. The rest two conditions are the original constraints and the non-negative constraints of the Lagrange multipliers.

5.2 The dual SVM formulation

Equations 23 and 24 allows us to remove the original parameters (\boldsymbol{w},b) from the Lagrangian. Note that

$$\frac{1}{2}\boldsymbol{w}^{T}\boldsymbol{w} = \frac{1}{2}\sum_{i=1}^{n}\sum_{j=1}^{n}\alpha_{i}\alpha_{j}y_{i}y_{j}\boldsymbol{x}_{i}^{T}\boldsymbol{x}_{j},$$
(28)

$$\sum_{i=1}^{n} \alpha_i y_i \boldsymbol{w}^T \boldsymbol{x}_i = \boldsymbol{w}^T \boldsymbol{w} = \sum_{i=1}^{n} \sum_{j=1}^{n} \alpha_i \alpha_j y_i y_j \boldsymbol{x}_i^T \boldsymbol{x}_j,$$
 (29)

$$\sum_{i=1}^{n} \alpha_i y_i b = \left(\sum_{i=1}^{n} \alpha_i y_i\right) b = 0, \qquad (30)$$

when these conditions hold. Hence, the Lagrangian becomes

$$-\frac{1}{2}\sum_{i=1}^{n}\sum_{j=1}^{n}\alpha_{i}\alpha_{j}y_{i}y_{j}\boldsymbol{x}_{i}^{T}\boldsymbol{x}_{j}+\sum_{i=1}^{n}\alpha_{i},$$
(31)

which does not involve w or b anymore. Putting the constraints on α_i in, we get²

$$\max_{\alpha} \sum_{i=1}^{n} \alpha_i - \frac{1}{2} \sum_{i=1}^{n} \sum_{j=1}^{n} \alpha_i \alpha_j y_i y_j \boldsymbol{x}_i^T \boldsymbol{x}_j$$
 (32)

s.t.
$$\alpha_i \ge 0, \quad i = 1, 2, \dots, n,$$
 (33)

²In more details, the above choice for \boldsymbol{w} and b leads to $g(\boldsymbol{\alpha}) = \inf_{(\boldsymbol{w},b)} L(\boldsymbol{w},b,\boldsymbol{\alpha})$, in which inf means the infimum (i.e., greatest lower bound). g is called the Lagrange dual function, and it is always concave. The *maximization* of g with respect to $\boldsymbol{\alpha}$ is always smaller than or equal to the minimization of the original (primal) problem, and the difference between them is called the duality gap. When the duality gap is 0, we can maximize the dual instead of solve the original minimization problem.

$$\sum_{i=1}^{n} \alpha_i y_i = 0. \tag{34}$$

which is called the dual SVM formulation, or simply the dual. One notable difference between the dual and the primal is: in the dual the training data never appear alone (as in the primal), they always appear in a duo as a dot-product $\mathbf{x}_i^T \mathbf{x}_j$.

In general, the optimal objective value of the primal and dual form of an optimization problem does not equal (i.e., there exists a duality gap). In SVM, the duality gap is 0—the dual and the primal forms will lead to the same optimal value. We will not talk about the optimization techniques in this note, which is beyond the scope of this course. However, many high quality optimization toolboxes (including those specifically designed for SVM) are available to solve either the primal or the dual SVM optimization.

One can solve the SVM optimization using the primal form and obtain $(\boldsymbol{w}^{\star}, b^{\star})$ directly; one can also solve the dual form and obtain the optimal Lagrange multipliers $\boldsymbol{\alpha}^{\star}$, and Equation 23 will give us the optimal \boldsymbol{w}^{\star} .

5.3 The optimal b value and support vectors

To obtain the optimal value for b is a little bit more tricky, which hinges on the complementary slackness property:

$$\alpha_i (y_i(\mathbf{w}^T \mathbf{x}_i + b) - 1) = 0 \qquad i = 1, 2, ..., n.$$
 (35)

Based on the complementary slackness, if we find one constraint whose corresponding Lagrange multiplier is positive (i.e., $\alpha_i > 0$), then we must have $y_i(\boldsymbol{w}^T\boldsymbol{x}_i+b)-1=0$. Hence, $b^\star=y_i-(\boldsymbol{w}^\star)^T\boldsymbol{x}_i$ for this particular $i.^3$ The optimal b value obtained in this manner may not be very reliable because some numerical errors can reduce its accuracy. Hence, we can also find all those examples whose corresponding α_i is non-zero, compute b from them individually, and then use their average as b^\star .

Those examples that are useful for computing b^* are special in the training set, whose corresponding Lagrange multiplier is positive. They are called the *support vectors*, and consequently the classification method is called the Support Vector Machine.

When $\alpha_i > 0$, we know $y_i(\boldsymbol{w}^T\boldsymbol{x}_i + b) - 1 = 0$, which means the *i*-th constraint has to be active, i.e., the margin of the training example \boldsymbol{x}_i must be 1, as shown in Figure 4.

In Figure 4, examples in two classes are denoted as blue circles and orange squares, respectively; the black line is the classification boundary $f(\mathbf{x}) = 0$, while the red lines correspond to $f(\mathbf{x}) = \pm 1$, i.e., all such examples whose margin $y_i f(\mathbf{x}_i)$ is 1 (remember in the SVM formulation we assume $y_i = \pm 1$ and the smallest margin of all training examples is 1?) The three examples (two

³This solution is derived because $y_i b = 1 - y_i \boldsymbol{w}^T \boldsymbol{x}_i$ and note that $y_i^2 = 1$.

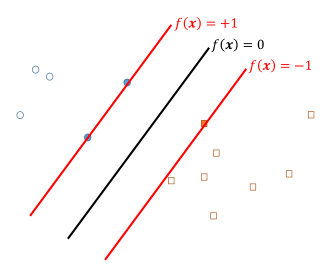


Figure 4: Illustration of the support vectors.

filled circles and one filled square) are the support vectors, whose corresponding $\alpha_i > 0$ and $y_i f(x_i) = 1$.

The classification boundary $f(x) = w^T x + b$ is a line in linear SVM. Hence, we can explicitly compute $w \in \mathbb{R}^d$, and it is very efficient to compute the decision value f(x), which only requires a dot-product computation. The linear SVM is hence both accurate and efficient. The computation of decision values for more complex SVM classifiers, e.g., non-linear SVM, requires a lot of computations.

The prediction (after the decision value f(x) is computed) is a simple task. As illustrated in Figure 4, the prediction is simply $\operatorname{sign}(f(x))$ in which sign is the sign function:

$$sign(x) = \begin{cases} +1 & \text{if } x > 0 \\ -1 & \text{if } x < 0 \\ 0 & \text{if } x = 0 \end{cases}$$
 (36)

When a test example x happens to have f(x) = 0, the above sign function return 0, which is not suitable for our classification problem. We can output a prediction of either +1 or -1 in an ad-hoc manner; we can also randomly assign a label for these special (and rare) testing examples.

5.4 Looking at the primal and dual simultaneously

Since the primal and dual give the same answer in SVM, we can look at them simultaneously. After the optimal primal and dual variables are obtained, they guarantee the margin of all training examples ≥ 1 , that is, the distance between any positive and negative training example is ≥ 2 . This observation provides a lower bound of the distance between examples from different classes.

The objective $\boldsymbol{w}^T\boldsymbol{w}$ is equal to $\sum_{i=1}^n\sum_{j=1}^n\alpha_i\alpha_jy_iy_j\boldsymbol{x}_i^T\boldsymbol{x}_j$. Let us define some more notations:

$$\alpha_{+} = \sum_{y_i = +1} \alpha_i \,, \tag{37}$$

$$\alpha_{-} = \sum_{y_i = -1} \alpha_i \,, \tag{38}$$

$$\mathbf{s}_{+} = \sum_{y_i = +1} \alpha_i \mathbf{x}_i \,, \tag{39}$$

$$\boldsymbol{s}_{-} = \sum_{y_i = -1} \alpha_i \boldsymbol{x}_i \,, \tag{40}$$

are the sum of Lagrange multipliers (weights) for all positive examples and negative examples, the weighted sum of all positive examples and all negative examples, respectively.

Equation 24 leads to $\alpha_+ = \alpha_-$, and Equation 23 shows $w = s_+ - s_-$. Hence, the primal objective is to minimize $||s_+ - s_-||^2$, which is also equivalent to

$$\left\| \frac{\mathbf{s}_+}{\alpha_+} - \frac{\mathbf{s}_-}{\alpha_-} \right\| . \tag{41}$$

 $\frac{s_+}{\alpha_+}$ ($\frac{s_-}{\alpha_-}$) is the weighted average of all positive (negative) examples. Hence, the distance between them must be ≥ 2 , and the equality can only happen if $\alpha_i = 0$ for all training examples whose margin is > 1. If any training example x_i has a margin $y_i f(x_i) > 1$ and $\alpha_i > 0$, then $\frac{s_+}{\alpha_+}$ or $\frac{s_-}{\alpha_-}$ will not reside on the two lines whose margin is 1 (cf. the red lines in Figure 4) and the distance between them will be > 2. Hence, in the separable case, all support vectors reside on these two lines and are sparse (i.e., the number of support vectors is only a small fraction of the number of all training examples, as shown in Figure 4).

6 Extensions for linearly inseparable and multiclass problems

We have made three assumptions in the above derivation: linear classifier, separable, and binary problems. These assumptions (and restrictions) have to be relaxed now that we already have a solution for the simplified problem. In this section, we deal with the non-separable case (still with linear classifiers), and the extension to multiclass problems. Nonlinear classifiers will be discussed in the next section.

6.1 Linear classifiers for non-separable problems

In some binary problems, the training set is not linearly separable, i.e., there is no single line that can perfectly separate examples in the two classes. However,

 $^{^4}$ The primal and dual parameters in this section refer to the optimal ones. However, we omit the * superscript to make the notations simpler.

as in the example of Fig. 1a, a linear classifier still seems the best model we can find using our eyes and brains.⁵ The linear SVM classifier can be extended to handle such inevitable classification errors for real-world problems, using a trick called the slack variables.

The current constraint, $y_i f(\mathbf{x}_i) \geq 1$, means that not only the *i*-th example is correctly classified, its distance to the linear boundary is also relatively far (> 1), leaving room for accommodating possible variations in test examples. However, shall we strictly maintain this constraint, if we can achieve a much larger margin of the dataset⁶ at the cost of violating this single constraint (e.g., $y_i f(\mathbf{x}_i) = 0.9 < 1$?) So long as $y_i f(\mathbf{x}_i) > 0$ (e.g., 0.9), this particular training example \mathbf{x}_i is still correctly classified, but larger margin are made possible for all other examples.

In another extreme, there may be one outlier x_i in the training set, e.g., $y_i = +1$ but it is in the cluster of negative examples. If we want to strictly maintain the constraint associated with x_i , we must pay the price that the problem is not solvable or the margin of other examples are greatly reduced. We need a mechanism to allow this outlier to be wrongly classified, i.e., to allow $y_i f(x_i) \leq 0$ in a few cases.

These exceptions (either $0 < y_i f(\boldsymbol{x}_i) < 1$ or $y_i f(\boldsymbol{x}_i) \leq 0$), however, cannot happen too frequently. We also need a mechanism to specify that the total prices we pay for such relatively rare or extreme cases are small. In SVM, a slack variable ξ_i is introduced as the price for \boldsymbol{x}_i , and its associated constraint is changed to two constraints:

$$y_i f(\mathbf{x}_i) \ge 1 - \xi_i \,, \tag{42}$$

$$\xi_i \ge 0. \tag{43}$$

When $\xi_i = 0$ for an \boldsymbol{x}_i , the original constraint is maintained, and no price is paid. When $0 < \xi_i \le 1$, this example is still correctly classified, but the margin is $1 - \xi_i < 1$, hence the price is ξ_i . When $\xi_i > 1$, this example is wrongly classified, and the price is still ξ_i . Hence, the total price is $\sum_{i=1}^n \xi_i$. To require the total price to be small, we just add this term to the (minimization) objective as a regularizer, and the primal linear SVM formulation becomes:

$$\min_{\boldsymbol{w},b} \quad \frac{1}{2} \boldsymbol{w}^T \boldsymbol{w} + C \sum_{i=1}^n \xi_i$$
 (44)

s.t.
$$y_i f(\mathbf{x}_i) \ge 1 - \xi_i$$
, $1 \le i \le n$, (45)

$$\xi_i \ge 0, \qquad 1 \le i \le n \,. \tag{46}$$

In the above SVM formulation, a new symbol C is introduced. C > 0 is a scalar that determines the relative importance between large margin (minimizing $\boldsymbol{w}^T\boldsymbol{w}$) and paying small total price $(\sum_{i=1}^n \xi_i)$. When C is large, the small

⁵In fact, the two classes in Fig. 1a are generated by two Gaussians with the same covariance matrix and equal prior, and the optimal classification boundary (in the minimum cost sense) is indeed a linear one.

⁶In the SVM formulation, this is equivalent to reducing $w^T w$ by a large amount while fixing the margin to 1.

total price part is more important; and when $C \to \infty$, the margin requirement is completely ignored. When C < 1, the large margin requirement is emphasized; and when $C \to 0$, no price is allowed at all. Hence, in problems showing different properties, different C value has to be tuned to seek best accuracy.

Using almost the same procedure as in the linearly separable case, we obtain the new dual form

$$\max_{\alpha} \sum_{i=1}^{n} \alpha_i - \frac{1}{2} \sum_{i=1}^{n} \sum_{j=1}^{n} \alpha_i \alpha_j y_i y_j \boldsymbol{x}_i^T \boldsymbol{x}_j$$
 (47)

s.t.
$$C \ge \alpha_i \ge 0, \quad i = 1, 2, \dots, n,$$
 (48)

$$\sum_{i=1}^{n} \alpha_i y_i = 0. \tag{49}$$

Note that the only difference is in the constraints on dual variables: $\alpha_i \geq 0$ is changed to $0 \leq \alpha_i \leq C$, which almost does not increase the difficulty of solving the dual problem, but the capacity of our SVM formulation has been highly extended.

We can also view the new objective from a different perspective. The total cost $\sum_{i=1}^{n} \xi_i$ can be seen as minimizing the cost over the training set, i.e., the empirical cost (or empirical loss). The term $\boldsymbol{w}^T\boldsymbol{w}$ can alternatively been viewed as a regularization term, which encourages linear classification boundaries that are less complex, because large components in \boldsymbol{w} allows larger variations in it and is hence more complex. Since $\xi_i \geq 0$ and $\xi_i \geq 1 - y_i f(\boldsymbol{x}_i)$ (based on Equations 42 and 43), we can combine them as

$$\xi_i = (1 - y_i f(\boldsymbol{x}_i))_+ , \qquad (50)$$

in which x_+ is called the *hinge loss* for x; $x_+ = x$ if $x \ge 0$, and $x_+ = 0$ if x < 0. It is obvious that $x_+ = \max(0, x)$. The hinge loss incurs no penalty if $y_i f(\boldsymbol{x})_i \ge 1$, i.e., when the margin for \boldsymbol{x}_i is larger than 1. Only when the margin is small $(0 < y_i f(\boldsymbol{x}_i) < 1)$ or \boldsymbol{x}_i is wrongly classified $(y_i f(\boldsymbol{x}_i) < 0)$, the loss $1 - y_i f(\boldsymbol{x}_i)$ is incurred. Using the hinge loss, we can rewrite the primal SVM as

$$\min_{\boldsymbol{w},b} \quad \frac{1}{2} \boldsymbol{w}^T \boldsymbol{w} + C \sum_{i=1}^n \xi_i \,, \quad \text{or}$$
 (51)

$$\min_{\boldsymbol{w},b} \quad \frac{1}{2} \boldsymbol{w}^T \boldsymbol{w} + C \sum_{i=1}^{n} (1 - y_i f(\boldsymbol{x}_i))_+ . \tag{52}$$

One practical question is: how to find the optimal C value for this tradeoff parameter? We will leave this to later sections.

6.2 Multiclass SVM

There are two main strategies to handle multiclass problems in SVM: one is to combine multiple binary SVM to solve the multiclass one; the other is to

extend the max-margin idea and its mathematics into multiclass using similar techniques, then solve the multiclass problem using a single optimization. We will introduce the first strategy.

The One-vs.-Rest (or one-versus-all, or OvA) method trains m binary SVM classifiers for an m-class problem. In training the i-th binary SVM, examples belonging to the i-th class are used as the positive class examples in the binary problem, and all other training examples are used as the negative class examples. Let us denote this binary SVM classifier as $f_i(x)$. In the binary case, when an example is far away from the classification boundary (i.e., with a large margin), there is good reason to believe it will be more tolerable to variations in the distribution of examples. In other words, we are more confident about our prediction if it has large margin.

In the multiclass case, if $f_i(\boldsymbol{x})$ is large (and positive), we can also say with high confidence that \boldsymbol{x} belongs to the *i*-th class. Hence, the m binary classifiers $f_i(\boldsymbol{x})$ ($1 \leq i \leq m$) gives us m confidence scores, and our prediction for \boldsymbol{x} is simply the class having the highest confidence. That is, the predicted class for a test example \boldsymbol{x} is

$$\underset{1 \le i \le m}{\arg\max} f_i(\boldsymbol{x}) \tag{53}$$

in the OvA method.

The One-vs-one (or one-versus-one, or OvO) method trains $\binom{m}{2} = \frac{m(m-1)}{2}$ binary SVM classifiers. For any pair (i,j) satisfying $1 \le i < j \le m$, we can use the examples in the i-th class as positive ones, and those in the j-th class as negative ones; all other training examples are ignored for this pair. The trained classifier $f_{i,j}(\boldsymbol{x})$ (after the sign function) determines whether \boldsymbol{x} should be assigned to the i- or j-th class, although it is possible that \boldsymbol{x} is neither in the i- nor j-th class; and we say the i- or j-th class received one vote. For any test example \boldsymbol{x} , it will receive $\binom{m}{2}$ votes, distributed among the m classes. If the real class of \boldsymbol{x} is k ($1 \le k \le m$), we expect the k-th class will receive the largest amount of votes. Hence, in the OvO method, the prediction for any example \boldsymbol{x} is the class who receives the largest amount of votes.

There have been empirical comparisons between the OvA and OvO methods. In terms of classification accuracy, there is not an obvious winner. Hence, both methods are applicable when m is not large. However, for problems with a large number of classes, e.g., when m=1000, the number of binary SVM classifiers to be trained is $\binom{m}{2}$ (499,500 if m=1000). To train a large number of classifiers requires excessively long time, and to store and apply all these trained models are also prohibitively expensive. Hence, the OvA method is more popular when m is large.

7 Kernel SVM

The final major missing block is a nonlinear SVM in which the classification boundary is not lines, but complex curves such as those in Figure 5. Figure 5a shows a problem with 200 training examples while Figure 5b has 2000. The blue

hyperbolic curves are the groundtruth decision boundary which separates the two classes, and the green curves are decision boundaries constructed by SVM using the nonlinear RBF kernel, which we will explain in this section. As shown in these figures, the kernel (or nonlinear) SVM can approximate the groundtruth decision boundary well, especially in the regions where many training examples are available.

Kernel methods are a class of methods that are popular in nonlinear learning, and SVM is a typical example of kernel methods.

7.1 The kernel trick

First, Let us consider and reexamine the second term in the SVM dual objective, $\sum_{i=1}^{n} \sum_{j=1}^{n} \alpha_i \alpha_j y_i y_j \boldsymbol{x}_i^T \boldsymbol{x}_j$. Inside the summations are an inner product between $\alpha_i y_i \boldsymbol{x}_i$ and $\alpha_j y_j \boldsymbol{x}_j$, i.e., inner product between two training examples whose sign is determined by the labels and the weights by the dual variables.

It is well known that the dot-product between two vectors can be treated as a measure of their similarity. If both vectors are unit ones, the dot-product equals the cosine of their angles. A small dot-product means the angle is close to 90° and the two vectors are far away from each other; a large positive dot-product means the angle is close to 0 and the two vectors are close to each other; and, a large negative dot-product means the angle is close to 180° and the two vectors are almost the farthest on a unit hypersphere.

Hence, it is natural to guess: if we replace the dot-product with some other nonlinear similarity metrics, can we get a nonlinear boundary better than a line? This guess seems viable because in the dual SVM formulation, as we have observed, the training examples always appear as dot-products!

Some theory (unfortunately beyond the scope of this note) endorses the above guess, but the nonlinear similarity measure must satisfy some condition (called Mercer's condition). Let κ be a nonlinear function satisfying the Mercer's condition, then the nonlinear dual SVM formulation is

$$\max_{\alpha} \sum_{i=1}^{n} \alpha_i - \frac{1}{2} \sum_{i=1}^{n} \sum_{j=1}^{n} \alpha_i \alpha_j y_i y_j \kappa(\boldsymbol{x}_i, \boldsymbol{x}_j)$$
 (54)

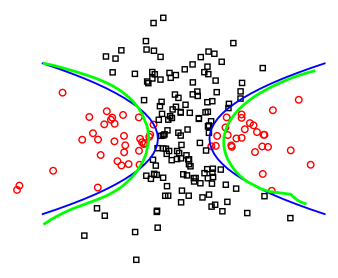
s.t.
$$C \ge \alpha_i \ge 0, \quad i = 1, 2, \dots, n,$$
 (55)

$$\sum_{i=1}^{n} \alpha_i y_i = 0. \tag{56}$$

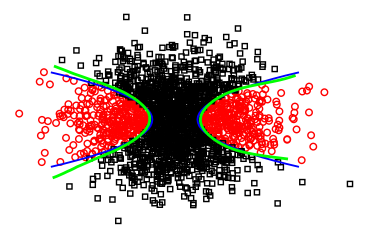
The optimization is the same as in the linear case, just replace the (constant) values $\mathbf{x}_i^T \mathbf{x}_i$ by $\kappa(\mathbf{x}_i, \mathbf{x}_i)$.

7.2 Mercer's condition and feature mapping

To introduce the Mercer's condition, we need to define the quadratically integrable (or square integrable) function concept. A function $g: \mathbb{R}^d \to \mathbb{R}$ is square



(a) 200 training examples



(b) 2000 training examples

Figure 5: Illustration of non-linear classifiers.

integrable if

$$\int_{-\infty}^{\infty} g^2(\mathbf{x}) \, \mathrm{d}\mathbf{x} < \infty \,, \tag{57}$$

 $\kappa(\cdot,\cdot): \mathbb{R}^d \times \mathbb{R}^d \mapsto \mathbb{R}$ satisfies the Mercer's condition if for any square integrable function $g(\boldsymbol{x})$, the following inequality is always true:

$$\iint \kappa(\boldsymbol{x}, \boldsymbol{y}) g(\boldsymbol{x}) g(\boldsymbol{y}) \, \mathrm{d}\boldsymbol{x} \, \mathrm{d}\boldsymbol{y} \ge 0, \qquad (58)$$

and it is symmetric, i.e., $\kappa(x, y) = \kappa(y, x)$.

In the context of SVM, Mercer's condition translates to another way to check whether κ is a valid kernel (i.e., meets Mercer's condition). For a $\kappa(\cdot, \cdot)$ and a set of examples $\boldsymbol{x}_1, \boldsymbol{x}_2, \ldots, \boldsymbol{x}_n \in \mathbb{R}^d$, one can define a matrix $K \in \mathbb{R}^n \times \mathbb{R}^n$, with $[K]_{ij} = \kappa(\boldsymbol{x}_i, \boldsymbol{x}_j)$ for $1 \leq i, j \leq n$. If the matrix K is positive semi-definite for arbitrary integer n > 0 and arbitrary set of examples $\boldsymbol{x}_1, \boldsymbol{x}_2, \ldots, \boldsymbol{x}_n$, then κ is a valid kernel function. When κ is a valid kernel, K is called the kernel matrix (induced by κ).

When κ is a valid kernel, then there exists a mapping $\phi : \mathbb{R}^d \mapsto \mathcal{X}$, which maps the input \boldsymbol{x} to a *feature space* \mathcal{X} , and satisfies

$$\kappa(\boldsymbol{x}, \boldsymbol{y}) = \phi(\boldsymbol{x})^T \phi(\boldsymbol{y}) \tag{59}$$

for any $x, y \in \mathbb{R}^d$. Although in many cases we do not know how to compute this mapping associated with κ , its existence has been rigorously proved. When we cannot explicitly spell out ϕ , we call it an implicit mapping.

The feature space \mathcal{X} is usually high dimensional. Its dimensionality is usually much larger than d, and for many kernels is infinite! Although we skip the advanced concepts (e.g., reproducing kernel Hilbert space), we want to assure the readers that so long as an infinite dimensional space satisfy certain conditions (such as the mappings in kernel SVM), the inner product between any two vectors in it are well-defined.

Conceptually, the kernel method has turned a nonlinear classification problem (in the input space) into an equivalent linear classification problem in the (usually much higher dimensional) feature space, as shown by the dual formulation for kernel SVM. In the feature space, the linear boundary shall enjoy the same max-margin benefit. Fortunately, because of the kernel trick, we do not need to compute the dot-product explicitly: $\kappa(\mathbf{x}, \mathbf{y})$ replaces $\phi(\mathbf{x})^T \phi(\mathbf{y})$.

The representer theorem also gives us $\mathbf{w} = \sum_{i=1}^{n} \alpha_i y_i \phi(\mathbf{x}_i)$. However, \mathbf{w} can be infinite dimensional, too. Hence, the prediction is performed through

$$f(\mathbf{x}) = \mathbf{w}^T \phi(\mathbf{x}) + b \tag{60}$$

$$= \sum_{i=1}^{n} \alpha_i y_i \phi(\mathbf{x}_i)^T \phi(\mathbf{x}) + b$$
 (61)

$$= \sum_{i=1}^{n} \alpha_i y_i \kappa(\boldsymbol{x}_i, \boldsymbol{x}) + b.$$
 (62)

This computation, however, is much more expensive than that in linear SVM. Assuming the complexity of $\kappa(\boldsymbol{x}_i, \boldsymbol{x})$ is O(d), predicting one example may take O(nd) steps. When the number of training examples is large (e.g., n = 1000000), kernel SVM prediction is very slow and we need to store all training examples in the SVM model.

The actual SVM prediction cost, however, is lower than O(nd). If a training example x_i is not a support vector, then its Lagrange multiplier α_i is 0, and is useless in the above summation. Hence, only support vectors are used in prediction and stored in the SVM model. The Lagrange multipliers are sparse, i.e., many α_i are zero. Let n' denote the number of support vectors, usually $n' \ll n$, and the kernel SVM prediction complexity O(n'd) is much lower than nd.

7.3 Popular kernels and the hyper-parameters

Commonly used kernel functions include

Linear:
$$\kappa(\boldsymbol{x}, \boldsymbol{y}) = \boldsymbol{x}^T \boldsymbol{y}$$
, (63)

RBF:
$$\kappa(\boldsymbol{x}, \boldsymbol{y}) = \exp(-\gamma \|\boldsymbol{x} - \boldsymbol{y}\|^2),$$
 (64)

Polynomial:
$$\kappa(\boldsymbol{x}, \boldsymbol{y}) = (\gamma \boldsymbol{x}^T \boldsymbol{y} + c)^D$$
. (65)

The RBF (radial basic function) kernel is sometimes called the Gaussian kernel. Some of these kernels are parameterized, e.g., the RBF kernel has a parameter $\gamma > 0$, and the polynomial kernel has three parameters γ , c and D (D is a positive integer, called the degree of the polynomial).

We can examine a special case of the polynomial kernel, when $\gamma = 1$, c = 1 and D = 2. With these parameters, the polynomial kernel is $\kappa(\boldsymbol{x}, \boldsymbol{y}) = (1 + \boldsymbol{x}^T \boldsymbol{y})^2$. If we assume the examples are in the two dimensional space, i.e., $\boldsymbol{x} = (x_1, x_2)^T$, $\boldsymbol{y} = (y_1, y_2)^T$, then

$$\kappa(\mathbf{x}, \mathbf{y}) = (1 + x_1 y_1 + x_2 y_2)^2 \tag{66}$$

$$= 1 + 2x_1y_1 + 2x_2y_2 + x_1^2y_1^2 + x_2^2y_2^2 + 2x_1x_2y_1y_2$$
 (67)

$$= \begin{bmatrix} 1\\ \sqrt{2}x_1\\ \sqrt{2}x_2\\ x_1^2\\ x_2^2\\ \sqrt{2}x_1x_2 \end{bmatrix}^T \begin{bmatrix} 1\\ \sqrt{2}y_1\\ \sqrt{2}y_2\\ y_1^2\\ y_2^2\\ \sqrt{2}y_1y_2 \end{bmatrix}$$
(68)

$$= \phi(\mathbf{x})^T \phi(\mathbf{y}). \tag{69}$$

Hence, for the degree 2 polynomial kernel, we can explicitly write down its mapping from the input space to the feature space. The feature space is 6 dimensional, three times that of the input space.

The tradeoff parameter C in the SVM formulation and the kernel parameters $\gamma,\ c$ and D are called hyper-parameters. Different hyper-parameters lead to

different kernels and SVM solutions. However, there is very few theoretical work that guide us on how to choose hyper-parameters. In practice, the most commonly used method is to use cross-validation to choose them. For example, we can try a few candidate values for C in the set

$$\{2^{-10}, 2^{-8}, 2^{-6}, 2^{-4}, 2^{-2}, 2^{0}, 2^{2}, 2^{4}, 2^{6}, 2^{8}, 2^{10}\}$$
.

Using the training set, we can obtain the cross-validation error rate on the $training\ set$ using each of these values. The C value that achieves the smallest cross-validation error rate is chosen.

This strategy, however, is very time consuming. Assuming an m class problem and a K-fold cross validation, $K \times \binom{m}{2}$ binary SVM classifiers are required to obtain a single cross-validation error rate if the OvO method is used. If there are two hyper-parameters (e.g., C and γ if the RBF kernel is used), and K_1 and K_2 candidate values exist for them, respectively, then the total number of binary SVM training is $KK_1K_2\binom{m}{2}$, which is very large.

7.4 SVM complexity, tradeoff, and more

In the early days, the popular SVM solver is the SMO (sequential minimal optimization) method, whose complexity is rather difficult to precisely estimate. Empirical evaluations usually show complexity higher than $O(n^{2.5}d)$, where n and d are the number and dimensionality of training examples, respectively.

For linear SVM, however, many fast methods have been proposed, which can finish the learning in O(nd) steps, using techniques such as column generation, stochastic gradient descent, and coordinate descent. These new methods enable the training of linear SVM for large problems, e.g., with millions of examples and/or millions of dimensions.

Although there are also acceleration techniques for nonlinear SVM learning, its training complexity is still much higher than that of linear SVM. The same also applies to the testing complexity. Linear SVM testing only requires a single dot-product in a binary linear SVM, while nonlinear SVM is much slower.

Nonlinear SVM, if can be trained in time for a problem, can usually achieve higher (and significantly higher in some problems) accuracy than the linear one. Hence, there is a tradeoff between the accuracy and complexity of SVM classifiers. Nonlinear SVM is a good choice when the dataset is small or medium sized, and linear SVM is a good option for large problems.

For some special kernels, such as the power mean kernels, algorithms specifically designed for them can obtain both fast speed and highly accurate models. High quality SVM solvers for both general nonlinear and fast linear kernels are available. Examples are the LIBSVM package (for general nonlinear SVM) and LIBLINEAR (for fast linear SVM). We will discuss more details about these topics in the exercises.

SVM can also be extended to handle regression problems, called the support vector regression (SVR), which is also implemented in many software packages, such as LIBSVM and LIBLINEAR.

Exercises

- 1. LIBSVM is a widely used software package for learning kernel SVM classifiers and regressors. In this problem, we will experiment with this software. LIBSVM also has an accompanying page that collects datasets.
 - (a) Download the LIBSVM software package from http://www.csie.ntu.edu.tw/~cjlin/libsvm/. Read the instructions and compile the software from source code.
 - (b) Download the svmguide1 dataset from the LIBSVM datasets page (https://www.csie.ntu.edu.tw/~cjlin/libsvmtools/datasets/).

For each of the following setup, learn SVM model(s) using the training set, and compare the accuracy on the test set under different setups.

- i. Use the default parameters (e.g., C = 1 and using the RBF kernel);
- ii. Use the svm-scale utility to normalize the features. Make sure your scaling operation is *proper*: use the parameters from the training set to normalize the test set data. Read the instructions for this utility program carefully;
- iii. Use the linear kernel instead of the RBF kernel;
- iv. Use C = 1000 and the RBF kernel:
- v. Use the <code>easy.py</code> utility to determine hyper-parameters C and gamma in the RBF kernel.

What have you learned from these experiments?

- (c) Play with the datasets in the LIBSVM datasets page. Can you find an imbalanced dataset? The LIBSVM parameter -wi is useful in handling imbalanced data. Try this parameter for the imbalanced dataset you have found. Is it helpful?
- 2. (Additive kernels) A kernel is called additive if it has the following form:

$$\kappa(\boldsymbol{x},\boldsymbol{y}) = \sum_{i=1}^d \kappa(x_i,y_i)$$

for two d-dimensional vectors \boldsymbol{x} and \boldsymbol{y} . Note that we used the same symbol κ to denote the kernel for comparing two vectors and two scalars.

- (a) The histogram intersection kernel is a widely used additive kernel, which is defined for vectors with only non-negative values, as $\kappa_{\rm HI}(\boldsymbol{x},\boldsymbol{y}) = \sum_{i=1}^d \min(x_i,y_i)$. Show that if $\kappa_{\rm HI}(x,y) = \min(x,y)$ for non-negative scalars x and y is a valid kernel, then $\kappa_{\rm HI}$ is a valid kernel for non-negative vectors \boldsymbol{x} and \boldsymbol{y} .
- (b) Now we consider κ_{HI} for scalars. For any set of non-negative scalar values x_1, x_2, \ldots, x_n , the kernel matrix formed by these values is $[X]_{ij} = \min(x_i, x_j)$ for $1 \le i, j \le n$, in which n is an arbitrary positive integer.

Let y_1, y_2, \ldots, y_n be a permutation of x_1, x_2, \ldots, x_n such that $y_1 \leq y_2 \leq \cdots \leq y_n$, and similarly define an $n \times n$ matrix Y with $y_{ij} = \min(y_i, y_j) = y_{\min(i,j)}$. Prove that X is positive (semi-) definite if and only if Y is positive (semi-)definite.

- (c) For any set of values $0 \le x_1 \le x_2 \le \cdots \le x_n$, prove that the kernel matrix X with $x_{ij} = \min(x_i, x_j) = x_{\min(i,j)}$ is positive semi-definite. (Hint: what is the LDL factorization of X?) Note that combining the above results have proved that $\kappa_{\rm HI}$ is a valid kernel for non-negative vectors.
- (d) The χ^2 kernel is another widely used additive kernel defined for positive data, which is defined as $\kappa_{\chi^2}(\boldsymbol{x},\boldsymbol{y}) = \sum_{i=1}^d \frac{2x_iy_i}{x_i+y_i}$. For positive data, this kernel is positive definite (see the next problem for a proof). Prove that $\kappa_{\rm HI}(\boldsymbol{x},\boldsymbol{y}) \leq \kappa_{\chi^2}(\boldsymbol{x},\boldsymbol{y})$.
- (e) The *Hellinger's kernel* is defined for non-negative data, as $\kappa_{\text{HE}}(\boldsymbol{x}, \boldsymbol{y}) = \sum_{i=1}^{d} \sqrt{x_i y_i}$. Prove that this is a valid kernel and $\kappa_{\text{HE}}(\boldsymbol{x}, \boldsymbol{y}) \geq \kappa_{\chi^2}(\boldsymbol{x}, \boldsymbol{y})$ for positive data.
- (f) Additive kernels are particularly effective when the feature vector is a *histogram*. A histogram (which is not normalized) contains natural numbers (i.e., zero or positive integers). Write out an explicit mapping for the histogram intersection kernel when the features are all natural numbers.
- 3. (Power mean kernels) In this problem, we introduce the *power mean kernels*, which are a family of additive kernels and are closely related to the *generalized mean* in mathematics.
 - (a) Read the information in the page https://en.wikipedia.org/wiki/Generalized_mean. When we only consider the power mean of two positive numbers x and y, what is $M_0(x,y)$, $M_{-1}(x,y)$ and $M_{-\infty}(x,y)$?
 - (b) The power mean kernels are a family of additive kernels, each indexed by a negative or zero real number p. For two positive vectors \boldsymbol{x} and \boldsymbol{y} with $x_i, y_i > 0$ $(1 \le i \le d)$, the power mean kernel M_p is defined as

$$M_p(\mathbf{x}, \mathbf{y}) = \sum_{i=1}^d M_p(x_i, y_i) = \sum_{i=1}^d \left(\frac{x^p + y^p}{2}\right)^{1/p}.$$

Power mean kernels were proposed by J. Wu in a paper titled "Power Mean SVM for Large Scale Visual Classification", CVPR 2012. Show that the three additive kernels discussed in the previous problem (histogram intersection, χ^2 and Hellinger's) are all special cases of the power mean kernels.

(c) (Conditionally positive definite kernel) A function κ is called a *conditionally positive definite kernel* if for an arbitrary positive integer n and any $c_1, c_2, \ldots, c_n \in \mathbb{R}$ which satisfies $\sum_{i=1}^n c_i = 0$, the inequality $\sum_{i=1}^n \sum_{j=1}^n c_i c_j \kappa(x_i, x_j) \geq 0$ holds for arbitrary $x_1, x_2, \ldots, x_n \in \mathbb{R}$. Prove

that $-\frac{x^p+y^p}{2}$ is conditionally positive definite for positive data.

(d) Use the following theorem to prove that power mean kernels are positive definite kernels $(-\infty \le p \le 0)$.

Theorem. If a kernel κ is conditionally positive definite and negative valued, then $\frac{1}{(-\kappa)^{\delta}}$ is positive definite for all $\delta \geq 0$.

- (e) When we use the power mean kernel $M_{-\infty}$ to replace the histogram intersection kernel in a programming environment, we have to use a p value that is far away from the zero value to replace $-\infty$. For example, we can use p=-32, i.e., using M_{-32} to approximate $M_{-\infty}$ / $\kappa_{\rm HI}$. Write a simple Matlab or Octave program to evaluate the largest absolute error and relative error between $M_{-32}(x,y)$ and $M_{-\infty}(x,y)$, where x and y are generated by the command x=0.01:0.01:1 and y=0.01:0.01:1, respectively. Is M_{-32} a good approximation for $\kappa_{\rm HI}$? That is, to compare $\kappa_{\rm HI}(x,y)$ and $M_{-32}(x,y)$ for the 10000 pairs of x and y values.
- (f) To illustrate the effectiveness of power mean kernels, visit https://sites.google.com/site/wujx2001/home/power-mean-svm and read the instructions there for installation and usage instructions. Use the resources in this page to generate training and testing data for the Caltech 101 dataset. When you generate the data, use K = 100 rather than K = 2000. Try different p values and apply the PmSVM software to this dataset. Use the LIBSVM software and the RBF kernel on the same dataset. Which one has higher accuracy on this dataset (whose feature vectors are histograms?") Note that if you use the tool provided in LIBSVM to choose optimal C and gamma hyper-parameter values, it will take very long to finish. Which software has higher learning and testing speed?
- 4. (SVM without bias) In the linear SVM formulation, we assume a classification boundary of the form $\boldsymbol{w}^T\boldsymbol{x} + b$, which includes a bias term $b \in \mathbb{R}$. However, it is also possible to learn a linear SVM classifier without the bias term, i.e., using $\boldsymbol{w}^T\boldsymbol{x}$ as the classification boundary.
 - (a) Without the bias term, what is the optimization problem in the primal space? Use the notations in this chapter.
 - (b) Without the bias term, show that the dual form is as follows.

$$\min_{\alpha} f(\alpha) = \frac{1}{2} \sum_{i=1}^{n} \sum_{j=1}^{n} \alpha_i \alpha_j y_i y_j x_i^T x_j - \sum_{i=1}^{n} \alpha_i$$
 (70)

s.t.
$$0 \le \alpha_i \le C$$
, $i = 1, 2, \dots, n$. (71)

If the solution for α is α^* , how do you find the optimal decision boundary w^* ?

(c) When a bias is preferred, this formulation (without bias) can still be useful. Given a training dataset (x_i, y_i) $(1 \le i \le n)$, one can convert

any $x \in \mathbb{R}^d$ into \hat{x} in \mathbb{R}^{d+1} by adding an extra dimension to x. The added dimension always has constant value 1. Suppose α^* is the optimal solution for the dual formulation, and suppose the classification boundary is $w^T x + b$, what is the optimal b value?

- 5. (Dual coordinate descent) In this problem, we introduce the *dual coordinate descent* algorithm for solving a linear SVM without the bias term in the dual space.
 - (a) Using the notations in the previous problem, find $\frac{\partial f}{\partial \alpha_i}$. Find a way to calculate $\frac{\partial f}{\partial \alpha_i}$ in $\mathcal{O}(d)$ steps. We use $f'(\alpha)$ to denote the partial derivative of f with respect to α , i.e., $f'(\alpha)_i = \frac{\partial f}{\partial \alpha_i}$.
 - (b) The dual coordinate descent (DCD) algorithm was proposed by Hsieh et al. in a paper titled "A Dual Coordinate Descent Method for Large-scale Linear SVM", which was published in ICML 2008. Among the n Lagrange multipliers α_i ($1 \le i \le n$), DCD updates one multiplier at a time. One training epoch updates $\alpha_1, \alpha_2, \ldots, \alpha_n$ sequentially one by one. The DCD algorithm was proved to converge and in practice it often converges in a few epochs. Now, suppose we fix $\alpha_1, \ldots, \alpha_{i-1}$ and $\alpha_{i+1}, \ldots, \alpha_n$, and want to find a better value for α_i . Denote the set of Lagrange multipliers after this update as α' , in which $\alpha'_i = \alpha_i + d$ and $\alpha'_i = \alpha_j$ for $j \ne i$. Show that

$$f(\boldsymbol{\alpha}') - f(\boldsymbol{\alpha}) = \frac{1}{2} \|\boldsymbol{x}_i\|^2 d^2 + f'(\boldsymbol{\alpha}) d.$$

- (c) Remember that α_i and α_i' should both be between 0 and C. Then, what is the optimal value for α_i' ?
- (d) A practical implementation of the DCD algorithm needs to pay attention to a lot of implementation details, e.g., randomize the order of updating for different α_i values in different epochs. Read the DCD paper and understand what details are important. This paper provides proof for DCD's convergence, too. A C++ implementation is provided at https://www.csie.ntu.edu.tw/~cjlin/liblinear/. Download this software and learn how to use it. This software has different solvers for linear classification and regression problems and several parameters. How do you specify parameters to perform classification in equations 70–71?
- (e) Download the rcv1.binary training set from the LIBSVM data page (https://www.csie.ntu.edu.tw/~cjlin/libsvmtools/datasets/).

Use 5-fold cross validation to obtain the training time and the cross-validation accuracy. Compare LIBSVM and LIBLINEAR on this task: use the linear kernel in LIBSVM and use the problem specified by equations 70-71 in LIBLINEAR. Set C=4 in these experiments.