Advances in Large Margin Classifiers

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edited by Alexander J. Smola Peter Bartlett Bernhard Schölkopf Dale Schuurmans

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Preface

 $Some\ good\ quote$

 $who\ knows$

some clever stuff ... and some more visionary comments

Alexander J. Smola, Peter Bartlett, Bernhard Schölkopf, Dale Schuurmans Berlin, Canberra, Waterloo, July 1999

1 Introduction to Large Margin Classifiers

The aim of this chapter is to provide a brief introduction to the basic concepts of large margin classifiers for readers unfamiliar with the topic. Moreover it is aimed at establishing a common basis in terms of notation and equations, upon which the subsequent chapters will build (and refer to) when dealing with more advanced issues.

1.1 A Simple Classification Problem

training data Assume that we are given a set of training data

$$X := \{\mathbf{x}_1, \dots, \mathbf{x}_m\} \subseteq \mathbb{R}^N \text{ where } m \in \mathbb{N}$$
 (1.1)

labels together with corresponding labels

$$Y := \{y_1, \dots, y_m\} \subseteq \{-1, 1\}. \tag{1.2}$$

The goal is to find some decision function $g: \mathbb{R}^N \to \{-1,1\}$ that accurately predicts the labels of unseen data points (\mathbf{x},y) . That is, we seek a function g that minimizes the classification error, which is given by the probability that $g(\mathbf{x}) \neq y$. A common approach to representing decision functions is to use a real valued prediction function $f: \mathbb{R}^N \to \mathbb{R}$ whose output is passed through a sign threshold to yield the final classification $g(\mathbf{x}) = \operatorname{sgn}(f(\mathbf{x}))$. Let us start with a simple example: linear decision functions. In this case the unthresholded prediction is given by a simple linear function of the input vector \mathbf{x}

linear decision function

$$g(\mathbf{x}) := \operatorname{sgn}(f(\mathbf{x})) \text{ where } f(\mathbf{x}) = (\mathbf{x} \cdot \mathbf{w}) + b \text{ for } \mathbf{w} \in \mathbb{R}^N \text{ and } b \in \mathbb{R}.$$
 (1.3)

This gives a classification rule whose decision boundary $\{\mathbf{x}|f(\mathbf{x})=0\}$ is an N-1 dimensional hyperplane separating the classes "+1" and "-1" from each other. Figure 1.1 depicts the situation. The problem of learning from data can be formulated as finding a set of parameters (\mathbf{w},b) such that $\mathrm{sgn}((\mathbf{w}\cdot\mathbf{x}_i)+b)=y_i$ for all $1\leq i\leq m$. However, such a solution may not always exist, in particular if we are dealing with noisy data. For instance, consider Figure 1.1 with the triangle replaced by an open circle. This raises the question what to do in such a situation.

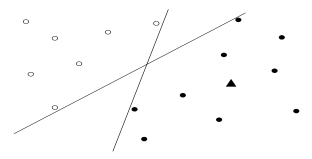


Figure 1.1 A linearly separable classification problem. Note that there may be several possible solutions as depicted by the two lines. The problem becomes non-separable if we replace the triangle by an open circle; in which case no solution (\mathbf{w}, b) exists.

1.1.1 Bayes Optimal Solution

Under the assumption that the data X, Y was generated from a probability distribution $p(\mathbf{x}, y)$ on $\mathbb{R}^N \times \{-1, 1\}$ and that p is known, it is straightforward to find a function that minimizes the probability of misclassification

$$R(g) := \int_{\mathbb{R}^{N} \times \{-1,1\}} 1_{\{g(\mathbf{x}) \neq y\}} p(\mathbf{x}, y) d\mathbf{x} dy.$$
 (1.4)

Bayes optimal decision function

This function satisfies

$$g(\mathbf{x}) = \operatorname{sgn} (p(\mathbf{x}, 1) - p(\mathbf{x}, -1)). \tag{1.5}$$

Consider a practical example.

Example 1.1 Two Gaussian Clusters

Assume that the two classes "+1" and "-1" are generated by two Gaussian clusters with the same covariance matrix Σ centered at μ_+ and μ_- respectively

$$p(\mathbf{x}, y) = \frac{1}{2(2\sigma)^{N/2} |\Sigma|^{1/2}} \begin{cases} e^{-\frac{1}{2}(\mathbf{x} - \boldsymbol{\mu}_{+})^{\mathsf{T}} \Sigma^{-1}(\mathbf{x} - \boldsymbol{\mu}_{+})} & \text{if } y = +1 \\ e^{-\frac{1}{2}(\mathbf{x} - \boldsymbol{\mu}_{-})^{\mathsf{T}} \Sigma^{-1}(\mathbf{x} - \boldsymbol{\mu}_{-})} & \text{if } y = -1. \end{cases}$$
(1.6)

Since the boundaries completely determine the decision function, we seek the set of points where $p(\mathbf{x}, +1) = p(\mathbf{x}, -1)$. In the case of (1.6) this is equivalent to seeking \mathbf{x} such that

$$(\mathbf{x} - \boldsymbol{\mu}_{+})^{\top} \Sigma^{-1} (\mathbf{x} - \boldsymbol{\mu}_{+}) = (\mathbf{x} - \boldsymbol{\mu}_{-})^{\top} \Sigma^{-1} (\mathbf{x} - \boldsymbol{\mu}_{-}). \tag{1.7}$$

By rearranging we find that this condition is equivalent to

$$\mathbf{x}^{\top} \Sigma^{-1} \mathbf{x} - 2\boldsymbol{\mu}_{+}^{\top} \Sigma^{-1} \mathbf{x} + \boldsymbol{\mu}_{+}^{\top} \Sigma^{-1} \boldsymbol{\mu}_{+} - \mathbf{x}^{\top} \Sigma^{-1} \mathbf{x} + 2\boldsymbol{\mu}_{-}^{\top} \Sigma^{-1} \mathbf{x} - \boldsymbol{\mu}_{-}^{\top} \Sigma^{-1} \boldsymbol{\mu}_{-} = 0$$

$$2(\boldsymbol{\mu}_{+}^{\top} \Sigma^{-1} - \boldsymbol{\mu}_{-}^{\top} \Sigma^{-1}) \mathbf{x} - (\boldsymbol{\mu}_{+}^{\top} \Sigma^{-1} \boldsymbol{\mu}_{+} - \boldsymbol{\mu}_{-}^{\top} \Sigma^{-1} \boldsymbol{\mu}_{-}) = 0 \quad (1.8)$$

The latter form is equivalent to having a linear decision function determined by

$$f(\mathbf{x}) = ((\mu_{+} - \mu_{-})^{\top} \Sigma^{-1}) \mathbf{x} - \frac{1}{2} (\mu_{+}^{\top} \Sigma^{-1} \mu_{+} - \mu_{-}^{\top} \Sigma^{-1} \mu_{-}).$$
 (1.9)

linear discriminant

Hence in this simple example the Bayes optimal classification rule is linear.

Problems arise, however, if $p(\mathbf{x}, y)$ is not known (as generally happens in practice). In this case one has to obtain a good *estimate* of $g(\mathbf{x}) = \operatorname{sgn}(f(\mathbf{x}))$ from the training data X, Y. A famous example of an algorithm for linear separation is the perceptron algorithm.

1.1.2 The Perceptron Algorithm

The perceptron algorithm is "incremental," in the sense that small changes are made to the weight vector in response to each labelled example in turn. For any learning rate $\eta > 0$, the algorithm acts sequentially as shown in Table 1.1. Notice

```
argument: Training sample, X = \{\mathbf{x}_1, \dots, \mathbf{x}_m\} \subset \mathcal{X}, Y = \{y_1, \dots, y_m\} \subset \{\pm 1\}
                    Learning rate, \eta
                    Weight vector \mathbf{w} and threshold b.
returns:
function Perceptron (X, Y, \eta)
     initialize \mathbf{w}, b = 0
     repeat
            for all i from i = 1, \ldots, m
                    Compute q(\mathbf{x}_i) = \operatorname{sgn}((\mathbf{w} \cdot \mathbf{x}_i) + b)
                    Update \mathbf{w}, b according to
                               \mathbf{w}' = \mathbf{w} + (\eta/2) (y_i - g(\mathbf{x}_i)) \mathbf{x}_i
                               b' = b + (\eta/2) (y_i - g(\mathbf{x}_i)).
            endfor
     until for all 1 \le i \le m we have g(\mathbf{x}_i) = y_i
     return f: \mathbf{x} \mapsto (\mathbf{w} \cdot \mathbf{x}) + b
end
```

 Table 1.1
 Basic Perceptron Algorithm.

perceptron algorithm

that (\mathbf{w}, b) is only updated on a labelled example if the perceptron in state (\mathbf{w}, b) misclassifies the example. It is convenient to think of the algorithm as maintaining the hypothesis $g: \mathbf{x} \mapsto \operatorname{sgn}((\mathbf{w} \cdot \mathbf{x}) + b)$, which is updated each time it misclassifies an example. The algorithm operates on a training sample by repeatedly cycling through the m examples, and when it has completed a cycle through the training data without updating its hypothesis, it returns that hypothesis.

The following result shows that if the training sample is consistent with some simple perceptron, then this algorithm converges after a finite number of iterations. In this theorem, \mathbf{w}^* and b^* define a decision boundary that correctly classifies all training points, and every training point is at least distance ρ from the decision boundary.

Theorem 1.1 Convergence of the Perceptron Algorithm

Suppose that there exists a $\rho > 0$, a weight vector \mathbf{w}^* satisfying $\|\mathbf{w}^*\| = 1$, and a threshold b^* such that

$$y_i\left((\mathbf{w}^* \cdot \mathbf{x}_i) + b^*\right) \ge \rho \text{ for all } 1 \le i \le m.$$
 (1.10)

Then for all $\eta > 0$, the hypothesis maintained by the perceptron algorithm converges after no more than $(b^{*2} + 1)(R^2 + 1)/\rho^2$ updates, where $R = \max_i ||x_i||^2$. Clearly, the limiting hypothesis is consistent with the training data (X, Y).

Proof Let (\mathbf{w}_j, b_j) be the state maintained immediately before the jth update occurring at, say, example (\mathbf{x}_i, y_i) . To measure the progress of the algorithm, we consider the evolution of the angle between (\mathbf{w}_j, b_j) and (\mathbf{w}^*, b^*) and note that the inner product $((\mathbf{w}_j, b_j) \cdot (\mathbf{w}^*, b^*))$ grows steadily with each update. To see this, note that (\mathbf{w}_j, b_j) is only updated when the corresponding hypothesis g_j misclassifies y_i , which implies that $y_i - g_j(\mathbf{x}_i) = 2y_i$. Therefore,

$$((\mathbf{w}_{j+1}, b_{j+1}) \cdot (\mathbf{w}^*, b^*)) = ([(\mathbf{w}_j, b_j) + (\eta/2)(y_i - g_j(\mathbf{x}_i))(\mathbf{x}_i, 1)] \cdot (\mathbf{w}^*, b^*))$$

$$= ((\mathbf{w}_j, b_j) \cdot (\mathbf{w}^*, b^*)) + \eta y_i((\mathbf{x}_i, 1) \cdot (\mathbf{w}^*, b^*))$$

$$\geq ((\mathbf{w}_j, b_j) \cdot (\mathbf{w}^*, b^*)) + \eta \rho$$

$$> j\eta\rho.$$

On the other hand, the norm of (\mathbf{w}_j, b_j) cannot grow too fast, because on an update we have $y_i((\mathbf{w}_i \cdot \mathbf{x}_i) + b_j) < 0$, and therefore

$$\begin{aligned} \|(\mathbf{w}_{j+1}, b_{j+1})\|^2 &= \|(\mathbf{w}_j, b_j) + \eta y_i(\mathbf{x}_i, 1)\|^2 \\ &= \|(\mathbf{w}_j, b_j)\|^2 + 2\eta y_i((\mathbf{x}_i, 1) \cdot (\mathbf{w}_j, b_j)) + \eta^2 \|(\mathbf{x}_i, 1)\|^2 \\ &\leq \|(\mathbf{w}_j, b_j)\|^2 + \eta^2 \|(\mathbf{x}_i, 1)\|^2 \\ &\leq j\eta^2 (R^2 + 1). \end{aligned}$$

Combining these two observations with the Cauchy-Schwarz inequality shows that

$$\sqrt{j\eta^{2}(R^{2}+1)} \ge \|(\mathbf{w}_{j+1}, b_{j+1})\|
\ge \frac{((\mathbf{w}_{j+1}, b_{j+1}) \cdot (\mathbf{w}^{*}, b^{*}))}{\sqrt{1+b^{*2}}}
\ge j\eta\rho,$$

and thus $j \leq (1 + b^{*2})(R^2 + 1)/\rho^2$ as desired.

Since the perceptron algorithm makes an update at least once in every cycle through the training data, and each iteration involves O(N) computation steps, this theorem implies that the perceptron algorithm has time complexity $O((R^2 + 1)mN/\rho^2)$.

1.1.3 Margins

The quantity ρ plays a crucial role in the previous theorem, since it determines how well the two classes can be separated and consequently how fast the perceptron

learning algorithm converges. This quantity ρ is what we shall henceforth call a margin.

Definition 1.1 Margin and Margin Errors

Denote by $f:\mathbb{R}^N\to\mathbb{R}$ a real valued hypothesis used for classification. Then

$$\rho_f(\mathbf{x}, y) := y f(\mathbf{x}), \tag{1.11}$$

margin

i.e. it is the margin by which the pattern \mathbf{x} is classified correctly (so that a negative value of $\rho_f(\mathbf{x}, y)$ corresponds to an incorrect classification). Moreover denote by

$$\rho_f := \min_{1 \le i \le m} \rho_f(\mathbf{x}_i, y_i) \tag{1.12}$$

minimum margin

the minimum margin over the whole sample. It is determined by the "worst" classification on the whole training set X, Y.

It appears to be desirable to have classifiers that achieve a large margin ρ_f since one might expect that an estimate that is "reliable" on the training set will also perform well on unseen examples. Moreover such an algorithm is more robust with respect to both patterns and parameters:

- Intuitively, for a pattern \mathbf{x} that is far from the decision boundary $\{\mathbf{x}|f(\mathbf{x})=0\}$ slight perturbations to \mathbf{x} will not change its classification $\operatorname{sgn}(f(\mathbf{x}))$. To see this, note that if $f(\mathbf{x})$ is a continuous function in \mathbf{x} then small variations in \mathbf{x} will translate into small variations in $f(\mathbf{x})$. Therefore, if $y_i f(\mathbf{x}_i)$ is much larger than zero, $y_i f(\mathbf{x}_i \pm \varepsilon)$ will also be positive for small ε . (See, for example, [21].)
- Similarly, a slight perturbation to the function f will not affect any of the resulting classifications on the training data $(\mathbf{x}_1, y_1), ..., (\mathbf{x}_m, y_m)$. Assume that $f_{\mathbf{w}}(\mathbf{x})$ is continuous in its parameters \mathbf{w} . Then, again, if $y_i f_{\mathbf{w}}(\mathbf{x}_i)$ is much larger than zero, $y_i f_{\mathbf{w} \pm \varepsilon}(\mathbf{x}_i)$ will also be positive for small ε .

robustness in patterns

robustness in parameters

1.1.4 Maximum Margin Hyperplanes

As pointed out in the previous section, it is desirable to have an estimator with a large margin. This raises the question whether there exists an estimator with maximum margin, i.e. whether there exists some f^* with

$$f^* := \underset{f}{\operatorname{argmax}} \ \rho_f = \underset{i}{\operatorname{argmax}} \ \underset{i}{\min} \ y_i f(\mathbf{x}_i). \tag{1.13}$$

Without some constraint on the size of \mathbf{w} , this maximum does not exist. In Theorem 1.1, we constrained \mathbf{w}^* to have unit length. If we define $f: \mathbb{R}^N \to \mathbb{R}$ by

$$f(\mathbf{x}) = \frac{(\mathbf{w} \cdot \mathbf{x}) + b}{\|\mathbf{w}\|},\tag{1.14}$$

then the maximum margin f is defined by the weight vector and threshold that satisfy

optimal hyperplane

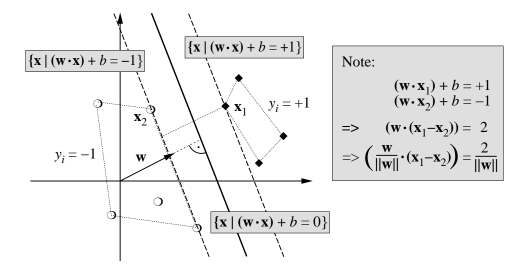


Figure 1.2 A binary classification toy problem: separate balls from diamonds. The optimal hyperplane is orthogonal to the shortest line connecting the convex hulls of the two classes (dotted), and intersects it half-way between the two classes. The problem being separable, there exists a weight vector \mathbf{w} and a threshold b such that $y_i((\mathbf{w} \cdot \mathbf{x}_i) + b) > 0$ $(i = 1, \dots, m)$. Rescaling \mathbf{w} and b such that the point(s) closest to the hyperplane satisfy $|(\mathbf{w} \cdot \mathbf{x}_i) + b| = 1$, we obtain a canonical form (\mathbf{w}, b) of the hyperplane, satisfying $y_i((\mathbf{w} \cdot \mathbf{x}_i) + b) > 1$. Note that in this case, the minimum Euclidean distance between the two classes (i.e. twice the margin), measured perpendicularly to the hyperplane, equals $2/\|\mathbf{w}\|$. This can be seen by considering two points \mathbf{x}_1 , \mathbf{x}_2 on opposite sides of the margin, i.e. $(\mathbf{w} \cdot \mathbf{x}_1) + b = 1$, $(\mathbf{w} \cdot \mathbf{x}_2) + b = -1$, and projecting them onto the hyperplane normal vector $\mathbf{w}/\|\mathbf{w}\|$.

$$\mathbf{w}^*, b^* = \underset{\mathbf{w}, b}{\operatorname{argmax}} \quad \min_{i=1}^{m} \quad \frac{y_i((\mathbf{w} \cdot \mathbf{x}_i) + b)}{\|\mathbf{w}\|}$$

$$= \underset{\mathbf{w}, b}{\operatorname{argmax}} \quad \min_{i=1}^{m} \quad y_i \operatorname{sgn}((\mathbf{w} \cdot \mathbf{x}_i) + b) \quad \left\| \frac{(\mathbf{w} \cdot \mathbf{x}_i)}{\|\mathbf{w}\|^2} \mathbf{w} + \frac{b}{\|\mathbf{w}\|^2} \mathbf{w} \right\|$$

$$(1.15)$$

$$= \underset{\mathbf{w}, b}{\operatorname{argmax}} \quad \min_{i=1}^{m} \quad y_{i} \operatorname{sgn}\left((\mathbf{w} \cdot \mathbf{x}_{i}) + b\right) \quad \left\| \frac{(\mathbf{w} \cdot \mathbf{x}_{i})}{\|\mathbf{w}\|^{2}} \mathbf{w} + \frac{b}{\|\mathbf{w}\|^{2}} \mathbf{w} \right\|$$
(1.16)

Euclidean Margin

The formulation (1.16) has a simple geometric interpretation: $-b\mathbf{w}/\|\mathbf{w}\|^2$ is the vector in direction w that ends right on the decision hyperplane (since $(\mathbf{w} \cdot (-b\mathbf{w}/\|\mathbf{w}\|^2)) = -b)$, and for a vector \mathbf{x}_i , $(\mathbf{w} \cdot \mathbf{x}_i)\mathbf{w}/\|\mathbf{w}\|^2$ is the projection of \mathbf{x}_i onto \mathbf{w} . Therefore, we are interested in maximizing the length of the vector differences $(\mathbf{w} \cdot \mathbf{x}_i) \mathbf{w} / ||\mathbf{w}||^2 - (-b\mathbf{w} / ||\mathbf{w}||^2)$ appropriatedly signed by $y_i g(\mathbf{x}_i)$.

The maxi-min problem (1.15) can be easily transformed into an equivalent constrained optimization task by conjecturing a lower bound on the margin, ρ , and maximizing ρ subject to the constraint that it really is a lower bound:

 $\mathbf{w}^*, b^*, \rho^*$

optimization problems

$$= \underset{\mathbf{w}, b, \rho}{\operatorname{argmax}} \quad \rho \quad \text{subject to } \frac{y_i((\mathbf{w} \cdot \mathbf{x}_i) + b)}{\|\mathbf{w}\|} \ge \rho \text{ for } 1 \le i \le m$$
 (1.17)

= argmax
$$\rho$$
 subject to $\|\mathbf{w}\| = 1$ and $y_i((\mathbf{w} \cdot \mathbf{x}_i) + b) \ge \rho$ for $1 \le i \le m$ (1.18)

=
$$\underset{\mathbf{w}}{\operatorname{argmin}} \|\mathbf{w}\|^2$$
 subject to $y_i((\mathbf{w} \cdot \mathbf{x}_i) + b) \ge 1$ for $1 \le i \le m$ (1.19)

quadratic program This last formulation is in the form of a quadratic programming problem, which can be easily handled using standard numerical routines [39, 10,].

Notice that (1.18) is in a particularly intuitive form. This formulation states that we are seeking a weight vector \mathbf{w} that obtains large dot products $y_i(\mathbf{w} \cdot \mathbf{x}_i)$, but constrain the weight vector to lie on the unit sphere to prevent obtaining such large dot products "for free" by scaling up \mathbf{w} . Interesting variants of problem (1.18) are obtained by choosing different norms to constrain the length of the weight vector. For example, constraining \mathbf{w} to lie on the unit ℓ_1 sphere instead of the unit ℓ_2 sphere gives the problem of determining

$$\mathbf{w}^*, b^*, \rho^*$$

$$= \underset{\mathbf{w}, b, \rho}{\operatorname{argmax}} \quad \rho \quad \text{subject to } \|\mathbf{w}\|_1 = 1 \text{ and } y_i((\mathbf{w} \cdot \mathbf{x}_i) + b) \ge \rho \text{ for } 1 \le i \le m \text{ (1.20)}$$

 ℓ_{∞} margin

which can easily be shown to be in the form of a linear programming problem. [41] shows that this is equivalent to finding the weight vector and threshold that maximize the minimum ℓ_{∞} distance between the training patterns and the decision hyperplane, in a direct analogue to the original Euclidean formulation (1.15).

Similarly, the constraint that **w** lie on the unit ℓ_{∞} sphere yields the problem

$$\mathbf{w}^*, b^*, \rho^*$$

$$= \underset{\mathbf{w}, b, \rho}{\operatorname{argmax}} \quad \rho \quad \text{subject to } \|\mathbf{w}\|_{\infty} = 1 \text{ and } y_i((\mathbf{w} \cdot \mathbf{x}_i) + b) \ge \rho \text{ for } 1 \le i \le m \text{ (1.21)}$$

 ℓ_1 margin

which is also a linear programming problem, but now equivalent to finding the weight vector and threshold that maximize the minimum ℓ_1 distance between the training patterns and the decision hyperplane. In general, constraining \mathbf{w} to lie on the unit ℓ_p sphere yields a convex programming problem

$$\mathbf{w}^*, b^*, \rho^*$$

$$= \underset{\mathbf{w}, b, \rho}{\operatorname{argmax}} \quad \rho \quad \text{subject to } \|\mathbf{w}\|_p = 1 \text{ and } y_i((\mathbf{w} \cdot \mathbf{x}_i) + b) \ge \rho \text{ for } 1 \le i \le m \text{ (1.22)}$$

 ℓ_q margin

which is equivalent to finding the weight vector and threshold that maximize the minimum ℓ_q distance between the training patterns and the decision hyperplane, where ℓ_p and ℓ_q are conjugate norms, i.e. such that $\frac{1}{p} + \frac{1}{q} = 1$ [41,].

In solving any of these constrained optimization problems, there is a notion of critical constraints; i.e. those inequality constraints that are satisfied as equalities by the optimal solution. In our setting, constraints correspond to training examples (\mathbf{x}_i, y_i) , $1 \leq i \leq m$, and the critical constraints are given by those training examples that lie right on the margin a distance ρ from the optimal hyperplane (cf. Figure 1.2). These critical training patterns are called Support Vectors.

Support Vectors

Notice that all the remaining examples of the training set are irrelevant: for non-critical examples the corresponding constraint $y_i((\mathbf{w} \cdot \mathbf{x}_i) + b) \ge 1$ in (1.19) does not play a role in the optimization, and therefore these points could be removed from the training set without affecting the results. This nicely captures our intuition of the problem: the hyperplane (cf. Figure 1.2) is completely determined by the patterns closest to it, the solution should not depend on the other examples.

soft margin hyperplane In practice, a separating hyperplane may not exist, e.g. if a high noise level causes a large overlap of the classes. The previous maximum margin algorithms perform poorly in this case because the maximum achievable minimum margin is negative, and this means the critical constraints are the mislabelled patterns that are furthest from the decision hyperplane. That is, the solution hyperplane is determined entirely by misclassified examples! To overcome the sensitivity to noisy training patterns, a standard approach is to allow for the possibility of examples violating the constraint in (1.19) by introducing slack variables [17, 74,]

slack variables

$$\xi_i \ge 0$$
, for all $i = 1, \dots, m$, (1.23)

along with relaxed constraints

$$y_i((\mathbf{w} \cdot \mathbf{x}_i) + b) \ge 1 - \xi_i$$
, for all $i = 1, \dots, m$. (1.24)

A classifier which generalizes well is then found by controlling both the size of \mathbf{w} and the number of training errors, minimizing the objective function

$$\tau(\mathbf{w}, \boldsymbol{\xi}) = \frac{1}{2} ||\mathbf{w}||^2 + C \sum_{i=1}^{m} \xi_i$$
 (1.25)

subject to the constraints (1.23) and (1.24), for some value of the constant C > 0. In the following section, we shall see why the size of \mathbf{w} is a good measure of the complexity of the classifier.

1.2 Theory

In order to provide a theoretical analysis of the learning problem we have to introduce a few definitions and assumptions about the process generating the data.

1.2.1 Basic Assumptions

independently identically distributed We assume that the training data X, Y is drawn independently and identically distributed (iid) according to some probability measure $p(\mathbf{x}, y)$. This means that all examples (\mathbf{x}_i, y_i) are drawn from $p(\mathbf{x}, y)$ regardless of the other examples or the index i.

This assumption is stronger than it may appear at first glance. For instance, time series data fails to satisfy the condition, since the observations are typically dependent, and their statistics might depend on the index i.

In (1.4), we defined the functional R(g) of a decision function g as the probability of misclassification. We can generalize this definition to apply to prediction functions f as well as thresholded decision functions g. This yields what we call the risk functional.

Definition 1.2 Risk Functional

Denote by $c(\mathbf{x}, y, f(\mathbf{x})) : \mathbb{R}^N \times \mathbb{R} \times \mathbb{R} \to [0, \infty)$ a cost function and by $p(\mathbf{x}, y)$ a probability measure as described above. Then the risk functional for a function $f : \mathbb{R}^N \to \mathbb{R}$ is defined as

Expected Risk

$$R(f) := \int_{\mathbb{R}^N \times \mathbb{R}} c(\mathbf{x}, y, f(\mathbf{x})) p(\mathbf{x}, y) d\mathbf{x} dy.$$
 (1.26)

Moreover the *empirical* risk functional for an m-sample X, Y is given by

$$R_{\text{emp}}(f) := \frac{1}{m} \sum_{i=1}^{m} c(\mathbf{x}_i, y_i, f(\mathbf{x}_i)). \tag{1.27}$$

Empirical Risk

For thresholded decision functions $g: \mathbb{R}^N \to \{-1,1\}$ we often use 0–1 classification error as the cost function $c(\mathbf{x},y,g(\mathbf{x})) = 1_{\{g(\mathbf{x})\neq y\}}$. In this case we obtain the risk functional defined in (1.4) (the probability of misclassification),

$$R(g) := \Pr\{g(\mathbf{x}) \neq y\}. \tag{1.28}$$

In this case, the empirical risk functional is

$$R_{\rm emp}(g) := \frac{1}{m} \sum_{i=1}^{m} 1_{\{g(\mathbf{x}_i) \neq y_i\}},\tag{1.29}$$

which is just the training error.

margin error

Finally we need a quantity called the *margin error* which is given by the proportion of training points that have margin less than ρ , i.e.

$$R_{\rho}(f) := \frac{1}{m} \sum_{i=1}^{m} 1_{\{y_i f(\mathbf{x}_i) < \rho\}}.$$
 (1.30)

This empirical estimate of risk counts a point as an error if it is either incorrectly classified or correctly classified by with margin less than ρ .

While one wants to minimize the risk R(g) this is hardly ever possible since $p(\mathbf{x}, y)$ is unknown. Hence one may only resort to minimizing $R_{\rm emp}(g)$ which is based on the training data. This, however, is not an effective method by itself—just consider an estimator that memorizes all the training data X,Y and generates random outputs for any other data. This clearly would have an empirical risk $R_{\rm emp}(g)=0$ but would obtain a true risk R(g)=0.5 (assuming the finite training sample has measure 0). The solution is to take the complexity of the estimate g into account as well, which will be discussed in the following sections.

1.2.2 Error Bounds for Thresholded Decision Functions

The central result of this analysis is to relate the number of training examples, the training set error, and the complexity of the hypothesis space to the generalization error. For thresholded decision functions, an appropriate measure for the complexity of the hypothesis space is the Vapnik-Chervonenkis (VC) dimension.

VC dimension

Definition 1.3 VC dimension

The VC dimension h of a space of $\{-1,1\}$ -valued functions, G, is the size of the largest subset of domain points that can be labelled arbitrarily by choosing functions only from G [79,].

The VC dimension can be used to prove high probability bounds on the error of a hypothesis chosen from a class of decision functions G—this is the famous result of [79]. The bounds have since been improved slightly by [70]—see also [3,].

Theorem 1.2 VC Upper Bound

Let G be a class of decision functions mapping \mathbb{R}^N to $\{-1,1\}$ that has VC dimension h. For any probability distribution $p(\mathbf{x},y)$ on $\mathbb{R}^N \times \{-1,1\}$, with probability at least $1-\delta$ over m random examples \mathbf{x} , for any hypothesis g in G the risk functional with 0–1 loss is bounded by

$$R(g) \le R_{\rm emp}(g) + \sqrt{\frac{c}{m} \left(h + \ln\left(\frac{1}{\delta}\right)\right)}$$
 (1.31)

where c is a universal constant. Furthermore, if $g^* \in G$ minimizes $R_{\text{emp}}(\cdot)$, then with probability $1 - \delta$

$$R(g^*) \le \inf_{g \in G} R(g) + \sqrt{\frac{c}{m} \left(h + \ln\left(\frac{1}{\delta}\right)\right)}$$
 (1.32)

(A short proof of this result is given by [38], but with worse constants than Talagrand's.)

These upper bounds are asymptotically close to the best possible, since there is also a lower bound with the same form:

Theorem 1.3 VC Lower Bound

Let G be a hypothesis space with finite VC dimension $h \ge 1$. Then for any learning algorithm there exist distributions such that with probability at least δ over m random examples, the error of its hypothesis g satisfies

$$R(g) \ge \inf_{g' \in G} R(g') + \sqrt{\frac{c}{m}} \left(h + \ln\left(\frac{1}{\delta}\right) \right)$$
 (1.33)

where c is a universal constant.

(Results of this form have been given by [20, 65, 5], using ideas from [23].)

Theorems 1.2 and 1.3 give a fairly complete characterization of the generalization error that can be achieved by choosing decision functions from a class G. However,

1.2 Theory 11

this characterization suffers from two drawbacks.

■ The first drawback is that the VC dimension must actually be determined (or at least bounded) for the class of interest—and this is often not easy to do. (However, bounds on the VC dimension h have been computed for many natural decision function classes, including parametric classes involving standard arithmetic and boolean operations. See [5] for a review of these results.)

■ The second (more serious) drawback is that the analysis ignores the *structure* of the mapping from training samples to hypotheses, and concentrates solely on the range of the learner's possible outputs. Ignoring the details of the learning map can omit many of the factors that are *crucial* for determining the success of the learning algorithm in real situations.

For example, consider learning algorithms that operate by first computing a real valued prediction function f from some class F and then thresholding this hypothesis to obtain the final decision function $g(\mathbf{x}) = \operatorname{sgn}(f(\mathbf{x}))$. Here, the VC dimension is a particularly weak method for measuring the representational capacity of the resulting function class $G = \operatorname{sgn}(F)$.

One reason is that the VC dimension of G is not sensitive to the *scale* of F at the accuracy level of interest. That is, it does not pay attention to whether the complexity of the hypothesis class is at a scale that is relevant for the outcome of the predictions.

The first step towards a more refined analysis that takes scale into account is given by [72]. Consider a set $X_0 \subset \mathbb{R}^N$ of input points with norm bounded by R > 0 (that is, $||\mathbf{x}_i|| \leq R$ for $\mathbf{x} \in X_0$), and the set F of bounded linear functions defined on X_0 ,

$$F = \{ \mathbf{x} \mapsto (\mathbf{w} \cdot \mathbf{x}) \mid ||\mathbf{w}|| \le 1, \ \mathbf{x} \in X_0 \}$$
 (1.34)

satisfying $|f(\mathbf{x})| \ge \rho$ for all patterns \mathbf{x} in X_0 . Then if we consider the set G of linear decision functions obtained by thresholding functions in F, [72] shows

$$VCdim(G) \le min\{R^2/\rho^2, N\} + 1.$$
 (1.35)

Note that this can be much smaller than the VC dimension of $\operatorname{sgn}(F)$ obtained without taking ρ into account, which is N+1 in this case. Therefore, one could hope to obtain significant benefits by using scale sensitive bounds which give much tighter results for large margin classifiers. Unfortunately, the bound (1.35) does not yet suffice for our purposes, because note that it requires that *all* points (including the test points) satisfy the margin condition, and therefore theorem 1.2 does not apply in this case. Rigorously obtaining these scale sensitive improvements is the topic we now address. In the following section, we consider scale-sensitive versions of the VC dimension, and obtain upper and lower bounds on risk in terms of these dimensions.

Margin Dependent Error Bounds for Real Valued Predictors

Definition 1.4 Fat Shattering Dimension

Let F be a set of real valued functions. We say that a set of points $S \subset \mathcal{X}$, which we will index as a vector $\mathbf{x} \in \mathcal{X}^{|S|}$, is ρ -shattered by F if there is a vector of real numbers $\mathbf{b} \in \mathbb{R}^{|S|}$ such that for any choice of signs $\mathbf{y} \in \{-1,1\}^{|S|}$ there is a function f in F that satisfies

$$y_i(f(x_i) - b_i) > \rho \text{ for } 1 < i < |S|.$$
 (1.36)

(That is, $f(x_i) \geq b_i + \rho$ if $y_i = 1$, and $f(x_i) \leq b_i - \rho$ if $y_i = -1$, for all x_i in S. Notice how similar this is to the notion of a minimum margin defined by (1.12).) The fat shattering dimension $\operatorname{fat}_F(\rho)$ of the set F is a function from the positive real numbers to the integers which maps a value ρ to the size of the largest ρ -shattered

set, if this is finite, or infinity otherwise.

the VC dimension obtained by thresholding but requiring that outputs are ρ above the threshold for positive classification and ρ below for negative.

We may think of the fat-shattering dimension of a set of real-valued functions as

The fat-shattering dimension is closely related to a more basic quantity, the covering number of a class of functions.

Definition 1.5 Covering Numbers of a Set

Denote by (S, d) a pseudometric space, $B_r(\mathbf{x})$ the closed ball in S centred at \mathbf{x} with radius r, T a subset of S, and ε some positive constant. Then the covering number $\mathcal{N}(\varepsilon,T)$ is defined as the minimum cardinality (that is, number of elements) of a set of points $T' \subset S$ such that

$$T \subseteq \bigcup_{\mathbf{x}_i \in T'} B_{\varepsilon}(\mathbf{x}_i), \tag{1.37}$$

i.e. such that the maximum difference of any element in T and the closest element in T' is less than or equal to ε .

Covering a class of functions F with an ε -cover means that one is able to approximately represent F (which may be of infinite cardinality) by a finite set. For learning, it turns out that it suffices to approximate the restrictions of functions in a class F to finite samples. For a subset X of some domain \mathcal{X} , define the pseudometric $\ell_{\infty,X}$ by

$$\ell_{\infty,X}(f,f') = \max_{\mathbf{x} \in X} |f(\mathbf{x}) - f'(\mathbf{x})| \tag{1.38}$$

where f and f' are real-valued functions defined on \mathcal{X} . Let $\mathcal{N}(\varepsilon, F, m)$ denote the maximum, over all $X \subset \mathcal{X}$ of size |X| = m, of the covering number $\mathcal{N}(\varepsilon, F)$ with respect to $\ell_{\infty,X}$. The following theorem shows that the fat-shattering dimension is intimately related to these covering numbers. (The upper bound is due to [4], and the lower bound to [9].)

fat shattering

covering number

Theorem 1.4 Bounds on N in terms of fat_F

Let F be a set of real functions from a domain \mathcal{X} to the bounded interval [0, B]. Let $\varepsilon > 0$ and let $m \ge \operatorname{fat}_F(\varepsilon/4)$. Then

$$\frac{\log_2 e}{8} \operatorname{fat}_F(16\varepsilon) \leq \log_2 \mathcal{N}(\varepsilon, F, m) \leq 3 \operatorname{fat}_F(\frac{\varepsilon}{4}) \log_2^2 \left(\frac{4eBm}{\varepsilon}\right). \tag{1.39}$$

Unfortunately, directly bounding \mathcal{N} can be quite difficult in general. Useful tools from functional analysis (which deal with the functional inverse of \mathcal{N} wrt. ϵ , the so called entropy number) for obtaining these bounds have been developed for classes of functions F defined by linear mappings from Hilbert spaces [15,], and linear functions over kernel expansions [89,].

The following result shows that we can use covering numbers to obtain upper bounds on risk in terms of margin error [64, 8,].

Theorem 1.5 Bounds on R(f) in terms of N and ρ

Suppose that F is a set of real-valued functions defined on \mathcal{X} , $\varepsilon \in (0,1)$ and $\rho > 0$. Fix a probability distribution on $\mathcal{X} \times \{-1,1\}$ and a sample size m. Then the probability that some f in F has $R_{\rho}(f) = 0$ but $R(f) \geq \varepsilon$ is no more than

$$2 \mathcal{N}\left(\frac{\rho}{2}, F, 2m\right) 2^{-\varepsilon m/2}. \tag{1.40}$$

Furthermore,

Pr ("some
$$f$$
 in F has $R(f) \ge R_{\rho}(f) + \varepsilon$ ") $\le 2 \mathcal{N}\left(\frac{\rho}{2}, F, 2m\right) e^{-\varepsilon^2 m/8}$. (1.41)

In fact, it is possible to obtain a similar result that depends only on the behaviour of functions in F near the threshold (see [5,] for details).

Let us have a close look at the bound (1.41) on the probability of excessive error. The factor $e^{-\varepsilon^2 m/8}$ in (1.41) stems from a bound of [30] on the probability of a large deviation of a sum of random variables from its mean. The factor $\mathcal{N}\left(\frac{\rho}{2}, F, 2m\right)$ stems from the fact that the continuous class of functions F was approximated (to accuracy $\rho/2$) by a finite number of functions. The 2m is due to the use of a symmetrization argument which is needed to make the overall argument work. Theorem 1.4 shows that this term is bounded by an exponential function of the fat-shattering dimension at scale $\rho/8$.

Interestingly, a similar result holds in regression. (For a review of these uniform convergence results, see [5,].

Theorem 1.6 Bounds on R(f) for Regression

Suppose that F is a set of functions defined on a domain \mathcal{X} and mapping into the real interval [0,1]. Let p be any probability distribution on $\mathcal{X} \times [0,1]$, ε any real number between 0 and 1, and $m \in \mathbb{N}$. Then for the quadratic cost function $c(\mathbf{x}, y, f(\mathbf{x})) = (y - f(\mathbf{x}))^2$ we have

$$\Pr\left(\sup_{f\in F}|R(f)-R_{\rm emp}(f)|\geq\varepsilon\right) \leq 4\mathcal{N}\left(\frac{\varepsilon}{16},F,2m\right)e^{-\varepsilon^2m/32}.$$
 (1.42)

anatomy of a uniform convergence bound Comparing with (1.41), notice that the scale of the covering number depends on the desired accuracy ε , whereas in (1.41) it depends on the scale ρ at which the margins are examined.

1.2.4 Error Bounds for Linear Decision Functions

The following result, due to [7], gives a bound on the fat-shattering dimension of large margin linear classifiers. It has a similar form to the bound (1.35) on the VC dimension of linear functions restricted to certain sets. It improves on a straightforward corollary of that result, and on a result of [28].

Theorem 1.7 Fat Shattering Dimension for Linear Classifiers

Suppose that B_R is the ℓ_2 ball of radius R in \mathbb{R}^n , centered at the origin, and consider the set

$$F := \{ f_{\mathbf{w}} \mid f_{\mathbf{w}}(\mathbf{x}) = (\mathbf{w} \cdot \mathbf{x}) \text{ with } ||\mathbf{w}|| \le 1, \mathbf{x} \in B_R \}.$$
 (1.43)

Ther

$$fat_F(\rho) \le \left(\frac{R}{\rho}\right)^2$$
. (1.44)

Using this result together with Theorems 1.4 and 1.5 gives the following theorem.

Theorem 1.8 Error Bounds for Linear Classifiers

Define the class F of real-valued functions on the ball of radius R as in (1.43). There is a constant c such that, for all probability distributions, with probability at least $1-\delta$ over m independently generated training examples, every $\rho > 0$ and every function $f \in F$ with margin at least ρ on all training examples (i.e. $R_{\rho}(f) = 0$) satisfies

$$R(f) \leq \frac{c}{m} \left(\frac{R^2}{\rho^2} \log^2 \left(\frac{m}{\rho} \right) + \log \left(\frac{1}{\delta} \right) \right). \tag{1.45}$$

Furthermore, with probability at least $1 - \delta$, for all $\rho > 0$, every function f in F has error

$$R(f) \leq R_{\rho}(f) + \sqrt{\frac{c}{m} \left(\frac{R^2}{\rho^2} \log^2 \left(\frac{m}{\rho}\right) + \log\left(\frac{1}{\delta}\right)\right)}. \tag{1.46}$$

For estimators using a linear programming approach as in [40,] one may state the following result, which then, via Theorem 1.4 can be transformed into a generalization bound as well.

Theorem 1.9 Capacity Bounds for Linear Classifiers

There is a constant c such that for the class

$$F_R = \left\{ \mathbf{x} \mapsto \mathbf{w}^T \mathbf{x} \mid \|\mathbf{x}\|_{\infty} \le 1, \|\mathbf{w}\|_1 \le R \right\}$$
(1.47)

we have

$$fat_{F_R}(\varepsilon) \leq c \left(\frac{R}{\varepsilon}\right)^2 \ln(2N+2).$$
(1.48)

Finally, we can obtain bounds for convex combinations of arbitrary hypotheses from a class G of $\{-1,1\}$ -valued functions,

$$co(G) = \left\{ \sum_{i} \alpha_{i} g_{i} \mid \alpha_{i} > 0, \sum_{i} \alpha_{i} = 1, g_{i} \in G \right\}.$$

$$(1.49)$$

See [53,]. These bounds are useful in analysing boosting algorithms; see Section 1.4.

Theorem 1.10 Bounds for Convex Combinations of Hypotheses

Let $p(\mathbf{x}, y)$ be a distribution over $\mathcal{X} \times \{-1, 1\}$, and let X be a sample of m examples chosen iid according to p. Suppose the base-hypothesis space G has VC dimension h, and let $\delta > 0$. Then with probability at least $1 - \delta$ over the random choice of the training set X, Y, every convex combination of functions $f \in \text{co}(G)$ satisfies the following bound for all $\rho > 0$.

$$R(f) \le R_{\rho}(f) + \sqrt{\frac{c}{m} \left(\frac{h \log^{2}(m/h)}{\rho^{2}} + \log\left(\frac{1}{\delta}\right)\right)}$$
(1.50)

1.3 Support Vector Machines

1.3.1 Optimization Problem

To construct the *Optimal Hyperplane* (cf. Figure 1.2), one solves the following optimization problem:

minimize
$$\tau(\mathbf{w}) = \frac{1}{2} ||\mathbf{w}||^2$$
 (1.51)

subject to
$$y_i((\mathbf{w} \cdot \mathbf{x}_i) + b) > 1$$
, for all $i = 1, \dots, m$. (1.52)

Lagrangian

This constrained optimization problem is dealt with by introducing Lagrange multipliers $\alpha_i \geq 0$ and a Lagrangian

$$L(\mathbf{w}, b, \boldsymbol{\alpha}) = \frac{1}{2} ||\mathbf{w}||^2 - \sum_{i=1}^{m} \alpha_i \left(y_i ((\mathbf{x}_i \cdot \mathbf{w}) + b) - 1 \right).$$
 (1.53)

The Lagrangian L has to be minimized with respect to the *primal variables* \mathbf{w} and b and maximized with respect to the *dual variables* α_i (i.e. a saddle point has to be found). Let us try to get some intuition for this. If a constraint (1.52) is violated, then $y_i((\mathbf{w} \cdot \mathbf{x}_i) + b) - 1 < 0$, in which case L can be increased by increasing the corresponding α_i . At the same time, \mathbf{w} and b will have to change such that L decreases. To prevent $-\alpha_i (y_i((\mathbf{w} \cdot \mathbf{x}_i) + b) - 1)$ from becoming arbitrarily large, the change in \mathbf{w} and b will ensure that, provided the problem is separable, the constraint will eventually be satisfied.

KKT conditions

Similarly, one can understand that for all constraints which are not precisely met as equalities, i.e. for which $y_i((\mathbf{w} \cdot \mathbf{x}_i) + b) - 1 > 0$, the corresponding α_i must be 0: this is the value of α_i that maximizes L. The latter is the statement of the Karush-Kuhn-Tucker complementarity conditions of optimization theory [34, 37, 10,].

The condition that at the saddle point, the derivatives of L with respect to the primal variables must vanish,

$$\frac{\partial}{\partial b}L(\mathbf{w}, b, \boldsymbol{\alpha}) = 0 \text{ and } \frac{\partial}{\partial \mathbf{w}}L(\mathbf{w}, b, \boldsymbol{\alpha}) = 0,$$
 (1.54)

leads to

$$\sum_{i=1}^{m} \alpha_i y_i = 0 \tag{1.55}$$

and

$$\mathbf{w} = \sum_{i=1}^{m} \alpha_i y_i \mathbf{x}_i. \tag{1.56}$$

support vector expansion

The solution vector thus has an expansion in terms of a subset of the training patterns, namely those patterns whose Lagrange multiplier α_i is non-zero. By the Karush-Kuhn-Tucker complementarity conditions these training patterns are the ones for which

$$\alpha_i(y_i((\mathbf{x}_i \cdot \mathbf{w}) + b) - 1) = 0, \quad i = 1, \dots, m, \tag{1.57}$$

and therefore they correspond precisely to the *Support Vectors* (i.e. critical constraints) discussed in Section 1.1.4. Thus we have the satisfying result that the Support Vectors are the only training patterns that determine the optimal decision hyperplane; all other training patterns are irrelevant and do not appear in the expansion (1.56).

By substituting (1.55) and (1.56) into L, one eliminates the primal variables and arrives at the Wolfe dual of the optimization problem [?, e.g.]]Bertsekas95: find multipliers α_i which

dual optimization problem

maximize
$$W(\alpha) = \sum_{i=1}^{m} \alpha_i - \frac{1}{2} \sum_{i,j=1}^{m} \alpha_i \alpha_j y_i y_j \left(\mathbf{x}_i \cdot \mathbf{x}_j \right)$$
 (1.58)

subject to
$$\alpha_i \ge 0$$
 for all $i = 1, \dots, m$, and $\sum_{i=1}^m \alpha_i y_i = 0$. (1.59)

The hyperplane decision function can thus be written as

$$f(\mathbf{x}) = \operatorname{sgn}\left(\sum_{i=1}^{m} y_i \alpha_i \left(\mathbf{x} \cdot \mathbf{x}_i\right) + b\right)$$
(1.60)

where b is computed using (1.57).

The structure of the optimization problem closely resembles those that typically arise in Lagrange's formulation of mechanics [?, e.g.]]Goldstein86. In that case also, it is often only a subset of the constraints that are active. For instance, if we keep

a ball in a box, then it will typically roll into one of the corners. The constraints corresponding to the walls which are not touched by the ball are irrelevant, the walls could just as well be removed.

Seen in this light, it is not too surprising that it is possible to give a mechanical interpretation of optimal margin hyperplanes [14,]: If we assume that each support vector \mathbf{x}_i exerts a perpendicular force of size α_i and sign y_i on a solid plane sheet lying along the hyperplane, then the solution satisfies the requirements of mechanical stability. The constraint (1.55) states that the forces on the sheet sum to zero; and (1.56) implies that the torques also sum to zero, via $\sum_i \mathbf{x}_i \times y_i \alpha_i \mathbf{w} / ||\mathbf{w}|| = \mathbf{w} \times \mathbf{w} / ||\mathbf{w}|| = 0$.

1.3.2 Feature Spaces and Kernels

To construct Support Vector Machines, the optimal hyperplane algorithm is augmented by a method for computing dot products in feature spaces that are nonlinearly related to input space [2, 12,]. The basic idea is to map the data into some other dot product space (called the feature space) \mathcal{F} via a nonlinear map

$$\Phi: \mathbb{R}^N \to \mathcal{F},\tag{1.61}$$

and then in the space \mathcal{F} perform the linear algorithm described above.

For instance, suppose we are given patterns $\mathbf{x} \in \mathbb{R}^N$ where most information is contained in the d-th order products (monomials) of entries x_j of \mathbf{x} , i.e. $x_{j_1}x_{j_2}\cdots x_{j_d}$, where $j_1,\ldots,j_d\in\{1,\ldots,N\}$. In that case, we might prefer to extract these monomial features first, and work in the feature space \mathcal{F} of all products of d entries.

This approach, however, fails for realistically sized problems: for N-dimensional input patterns, there exist (N+d-1)!/(d!(N-1)!) different monomials. Already 16×16 pixel input images (e.g. in character recognition) and a monomial degree d=5 yield a dimensionality of 10^{10} .

This problem can be overcome by noticing that both the construction of the optimal hyperplane in \mathcal{F} (cf. (1.58)) and the evaluation of the corresponding decision function (1.60) only require the evaluation of dot products $(\Phi(\mathbf{x}) \cdot \Phi(\mathbf{x}'))$, and never require the mapped patterns $\Phi(\mathbf{x})$ in explicit form. This is crucial, since in some cases, the dot products can be evaluated by a simple kernel [2, 12,].

 $k(\mathbf{x}, \mathbf{x}') = (\Phi(\mathbf{x}) \cdot \Phi(\mathbf{x}')). \tag{1.62}$

For instance, the polynomial kernel

$$k(\mathbf{x}, \mathbf{x}') = (\mathbf{x} \cdot \mathbf{x}')^d \tag{1.63}$$

can be shown to correspond to a map Φ into the space spanned by all products of exactly d dimensions of \mathbb{R}^N ([48, 12]). For a proof, see [55]. For d=2 and $\mathbf{x}, \mathbf{x}' \in \mathbb{R}^2$, for example, we have [74,]

$$(\mathbf{x} \cdot \mathbf{x}')^2 = (x_1^2, x_2^2, \sqrt{2} \ x_1 x_2) (y_1^2, y_2^2, \sqrt{2} \ y_1 y_2)^\top = (\Phi(\mathbf{x}) \cdot \Phi(\mathbf{x}')), \tag{1.64}$$

feature space

Mercer kernel

polynomial kernel defining $\Phi(\mathbf{x}) = (x_1^2, x_2^2, \sqrt{2} x_1 x_2).$

By using $k(\mathbf{x}, \mathbf{x}') = ((\mathbf{x} \cdot \mathbf{x}') + c)^d$ with c > 0, we can take into account all product of order up to d (i.e. including those of order smaller than d).

More generally, the following theorem of functional analysis shows that kernels k of positive integral operators give rise to maps Φ such that (1.62) holds [42, 2, 12, 22,]:

Theorem 1.11 Mercer

positive integral operator If k is a continuous symmetric kernel of a positive integral operator T, i.e.

$$(Tf)(\mathbf{x}') = \int_{\mathcal{X}} k(\mathbf{x}, \mathbf{x}') f(\mathbf{x}) d\mathbf{x}$$
(1.65)

with

$$\int_{\mathcal{X}\times\mathcal{X}} k(\mathbf{x}, \mathbf{x}') f(\mathbf{x}) f(\mathbf{x}') d\mathbf{x} d\mathbf{x}' \ge 0$$
(1.66)

for all $f \in L_2(\mathcal{X})$ (\mathcal{X} being a compact subset of \mathbb{R}^N), it can be expanded in a uniformly convergent series (on $\mathcal{X} \times \mathcal{X}$) in terms of T's eigenfunctions ψ_j and positive eigenvalues λ_j ,

$$k(\mathbf{x}, \mathbf{x}') = \sum_{j=1}^{N_F} \lambda_j \psi_j(\mathbf{x}) \psi_j(\mathbf{x}'), \tag{1.67}$$

where $N_{\mathcal{F}} \leq \infty$ is the number of positive eigenvalues.

An equivalent way to characterize Mercer kernels is that they give rise to positive matrices $K_{ij} := k(\mathbf{x}_i, \mathbf{x}_j)$ for all $\{\mathbf{x}_1, \dots, \mathbf{x}_m\}$ [51,].

From (1.67), it is straightforward to construct a map Φ into a potentially infinite-dimensional l_2 space which satisfies (1.62). For instance, we may use

$$\Phi(\mathbf{x}) = (\sqrt{\lambda_1}\psi_1(\mathbf{x}), \sqrt{\lambda_2}\psi_2(\mathbf{x}), \dots). \tag{1.68}$$

Rather than thinking of the feature space as an l_2 space, we can alternatively represent it as the Hilbert space \mathcal{H}_k containing all linear combinations of the functions $f(.) = k(\mathbf{x}_i, .)$ ($\mathbf{x}_i \in \mathcal{X}$). To ensure that the map $\Phi : \mathcal{X} \to \mathcal{H}_k$, which in this case is defined as

$$\Phi(\mathbf{x}) = k(\mathbf{x}, .), \tag{1.69}$$

satisfies (1.62), we need to endow \mathcal{H}_k with a suitable dot product $\langle .,. \rangle$. In view of the definition of Φ , this dot product needs to satisfy

$$\langle k(\mathbf{x},.), k(\mathbf{x}',.) \rangle = k(\mathbf{x}, \mathbf{x}'), \tag{1.70}$$

reproducing kernel which amounts to saying that k is a reproducing kernel for \mathcal{H}_k . For a Mercer kernel (1.67), such a dot product does exist. Since k is symmetric, the ψ_i $(i = 1, ..., N_{\mathcal{F}})$ can be chosen to be orthogonal with respect to the dot product in $L_2(C)$, i.e. $(\psi_j, \psi_n)_{L_2(C)} = \delta_{jn}$, using the Kronecker δ_{jn} . From this, we can construct $\langle .,. \rangle$

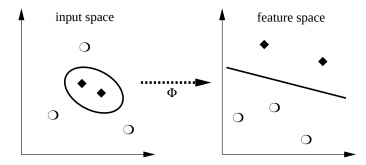


Figure 1.3 The idea of SV machines: map the training data nonlinearly into a higher-dimensional feature space via Φ , and construct a separating hyperplane with maximum margin there. This yields a nonlinear decision boundary in input space. By the use of a kernel function (1.62), it is possible to compute the separating hyperplane without explicitly carrying out the map into the feature space.

such that

$$\langle \sqrt{\lambda_j} \psi_j, \sqrt{\lambda_n} \psi_n \rangle = \delta_{jn}. \tag{1.71}$$

Substituting (1.67) into (1.70) then proves the desired equality (for further details, see [6, 86, 26, 55]).

sigmoid kernel Besides (1.63), SV practictioners use sigmoid kernels

$$k(\mathbf{x}, \mathbf{x}') = \tanh(\kappa(\mathbf{x} \cdot \mathbf{x}') + \Theta) \tag{1.72}$$

for suitable values of gain κ and threshold Θ , and radial basis function kernels, as for instance [2, 12, 62,]

$$k(\mathbf{x}, \mathbf{x}') = \exp\left(-\|\mathbf{x} - \mathbf{x}'\|^2 / (2\sigma^2)\right),\tag{1.73}$$

with $\sigma > 0$. Note that when using Gaussian kernels, for instance, the feature space \mathcal{H}_k thus contains all superpositions of Gaussians on \mathcal{X} (plus limit points), whereas by definition of Φ (1.69), only single bumps $k(\mathbf{x}, ...)$ do have pre-images under Φ .

The main lesson from the study of kernel functions, is that the use of kernels can turn any algorithm that only depends on dot products into a nonlinear algorithm which is linear in feature space. In the time since this was explicitly pointed out [59,] a number of such algorithms have been proposed: until then the applications of the kernel trick were a proof of the convergence of rbf network training by [2,] and the nonlinear variant of the SV algorithm by [12] (see Figure 1.3). To construct SV machines, one computes an optimal hyperplane in feature space. To this end, we substitute $\Phi(\mathbf{x}_i)$ for each training example \mathbf{x}_i . The weight vector (cf. (1.56)) then becomes an expansion in feature space. Note that \mathbf{w} will typically no more correspond to the image of just a single vector from input space (cf. [56] for a formula to compute the pre-image if it exists), in other words, \mathbf{w} may not be directly accessible any more. However, since all patterns only occur in dot products, one can substitute Mercer kernels k for the dot products [12, 29,], leading to decision

kernel

kernel

Gaussian RBF

decision function

functions of the more general form (cf. (1.60))

$$g(\mathbf{x}) = \operatorname{sgn}\left(\sum_{i=1}^{m} y_i \alpha_i \left(\Phi(\mathbf{x}) \cdot \Phi(\mathbf{x}_i)\right) + b\right) = \operatorname{sgn}\left(\sum_{i=1}^{m} y_i \alpha_i k(\mathbf{x}, \mathbf{x}_i) + b\right)$$
(1.74)

and the following quadratic program (cf. (1.58)):

maximize
$$W(\alpha) = \sum_{i=1}^{m} \alpha_i - \frac{1}{2} \sum_{i,j=1}^{m} \alpha_i \alpha_j y_i y_j k(\mathbf{x}_i, \mathbf{x}_j)$$
 (1.75)

subject to
$$\alpha_i \ge 0, \ i = 1, ..., m, \ \text{and} \ \sum_{i=1}^m \alpha_i y_i = 0.$$
 (1.76)

soft margin and kernels

Recall that, as discussed in Section 1.1.4 a separating hyperplane may not always exist, even in the expanded feature space \mathcal{F} . To cope with this difficulty, slack variables were introduced to yield the *soft margin* optimal hyperplane problem (1.25). Incorporating kernels, and rewriting (1.25) in terms of Lagrange multipliers, this again leads to the problem of maximizing (1.75), but now subject to the constraints

$$0 \le \alpha_i \le C, \quad i = 1, \dots, m, \text{ and } \sum_{i=1}^m \alpha_i y_i = 0.$$
 (1.77)

The only difference from the separable case (1.76) is the upper bound C on the Lagrange multipliers α_i . This way, the influence of the individual patterns (which could always be outliers) gets limited. As above, the solution takes the form (1.74). The threshold b can be computed by exploiting the fact that for all SVs \mathbf{x}_i with $\alpha_i < C$, the slack variable ξ_i is zero (this again follows from the Karush-Kuhn-Tucker complementarity conditions), and hence

$$\sum_{j=1}^{m} y_j \alpha_j k(\mathbf{x}_i, \mathbf{x}_j) + b = y_i.$$
(1.78)

If one uses an optimizer that works with the double dual [?, e.g.]] Vanderbei 97, one can also recover the value of the primal variable b directly from the corresponding double dual variable.

Finally, the algorithm can be modified such that it does not require the regularization constant C. Instead, one specifies an upper bound $0 \le \nu \le 1$ on the fraction of points allowed to lie in the margin (asymptotically, the number of SVs) [60,]. This leaves us with a homogeneous target function made up by the quadratic part of (1.75), and an additional lower bound constraint on the sum over all Lagrange multipliers.

1.3.3 Smoothness and Regularization

For kernel-based function expansions, one can show [67,] that given a regularization operator P mapping the functions of the learning machine into some dot product

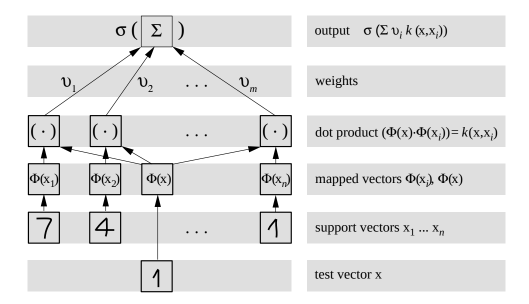


Figure 1.4 Architecture of SV machines. The input \mathbf{x} and the Support Vectors \mathbf{x}_i are nonlinearly mapped (by Φ) into a feature space \mathcal{F} , where dot products are computed. By the use of the kernel k, these two layers are in practice computed in one single step. The results are linearly combined by weights v_i , found by solving a quadratic program (in pattern recognition, $v_i = y_i \alpha_i$; in regression estimation, $v_i = \alpha_i^* - \alpha_i$). The linear combination is fed into the function σ (in pattern recognition, $\sigma(x) = \operatorname{sgn}(x + b)$; in regression estimation, $\sigma(x) = x + b$).

space, the problem of minimizing the regularized risk

$$R_{\text{reg}}(f) := R_{\text{emp}}(f) + \frac{\lambda}{2} ||Pf||^2$$
 (1.79)

regularized risk

(with a regularization parameter $\lambda \geq 0$) can be written as a constrained optimization problem. For particular choices of the loss function, it further reduces to a SV type quadratic programming problem. The latter thus is not specific to SV machines, but is common to a much wider class of approaches. What gets lost in the general case, however, is the fact that the solution can usually be expressed in terms of a small number of SVs (cf. also [26], who establishes a connection between SV machines and basis pursuit denoising [16,]). This specific feature of SV machines is due to the fact that the type of regularization and the class of functions that the estimate is chosen from are intimately related [27, 66, 68,]: the SV algorithm is equivalent to minimizing the regularized risk $R_{\rm reg}(f)$ on the set of functions

$$f(\mathbf{x}) = \sum_{i} \alpha_{i} k(\mathbf{x}_{i}, \mathbf{x}) + b, \tag{1.80}$$

provided that k and P are interrelated by

$$k(\mathbf{x}_i, \mathbf{x}_j) = ((Pk)(\mathbf{x}_i, .) \cdot (Pk)(\mathbf{x}_j, .)). \tag{1.81}$$

To this end, k is chosen as a Green's function of P^*P , for in that case, the right hand side of (1.81) equals $(k(\mathbf{x}_i,.)\cdot(P^*Pk)(\mathbf{x}_j,.))=(k(\mathbf{x}_i,.)\cdot\delta_{\mathbf{x}_j}(.))=k(\mathbf{x}_i,\mathbf{x}_j)$. For instance, an RBF kernel corresponds to regularization with a functional containing a specific differential operator.

In SV machines, the kernel thus plays a dual role: firstly, it determines the class of functions (1.80) that the solution is taken from; secondly, via (1.81), the kernel determines the type of regularization that is used. The next question, naturally, is what type of regularization (i.e. kernel) we should use in order to get the best generalization performance. Using bounds on covering numbers of Hilbert spaces [15,], one can show [89, 88, 57,] that the eigenspectrum of the matrix $k(x_i, x_j)$ is closely connected to the latter and also to the eigenspectrum of the kernel k.

For arbitrary expansions of f into basis functions, say f_i , the considerations about smoothness of the estimate still hold, provided ||Pf|| is a norm in the space spanned by the basis functions f_i (otherwise one could find functions $f \in \text{span}\{f_i\}$ with ||Pf|| = 0, however $f \neq 0$). In this case the existing bounds for kernel expansions can be readily applied to regularization networks as well (cf. e.g. [89, 69,] for details). However, one can show [36, 18,], that such an expansion may not fully minimize the regularized risk functional (1.79). This is one of the reasons why often only kernel expansions are considered.

Finally it is worth while pointing out the connection between Gaussian Processes and Support Vector machines. The similarity is most obvious in regression, where the Support Vector solution is the maximum a posteriori estimate of the corresponding Bayesian inference scheme [87,]. In particular, the kernel k of Support Vector machines plays the role of a covariance function such that the prior probability of a function $f = \sum_i \alpha_i k(\mathbf{x}_i, \mathbf{x})$ is given by

$$P(f) \propto \exp\left(-\frac{1}{2}\|Pf\|^2\right) = \exp\left(-\frac{1}{2}\sum_{i,j}\alpha_i\alpha_j k(\mathbf{x}_i, \mathbf{x}_j)\right).$$
 (1.82)

Bayesian methods, however, require averaging over the posterior distribution P(f|X,Y) in order to obtain the final estimate and to derive error bounds. In classification the situation is even more complicated, since we have Bernoulli distributed random variables for the labels of the classifier. See [87,] for more details on this subject.

1.3.4 A Bound on the Leave-One-Out Estimate

Besides the bounds directly involving large margins, which are useful for stating uniform convergence results, one may also try to estimate R(f) by using leave-one-out estimates. Denote by f_i the estimate obtained from $X \setminus \{\mathbf{x}_i\}, Y \setminus \{y_i\}$. Then

regularization networks

Gaussian processes

1.4 Boosting 23

$$R_{\text{out}}(f) := \frac{1}{m} \sum_{i=1}^{m} c(\mathbf{x}_i, y_i, f_i(\mathbf{x}_i))$$
(1.83)

One can show (cf. e.g. [72,]) that the latter is an unbiased estimator of R(f). Unfortunately, $R_{\rm out}(f)$ is hard to compute and thus rarely used. In the case of Support Vector classification, however, an upper bound on $R_{\rm out}(f)$ is not too difficult to obtain. [74] showed that the fraction of Support Vectors is an upper bound on $R_{\rm out}(f)$. [32] have generalized this result as follows

$$R_{\text{out}}(f) \leq \frac{1}{m} \sum_{i=1}^{m} 1_{\left\{ y_{i} \sum_{j \neq i} \alpha_{j} y_{j} k(\mathbf{x}_{j}, \mathbf{x}_{i}) + y_{i} b > 0 \right\}}$$

$$= \frac{1}{m} \sum_{i=1}^{m} 1_{\left\{ y_{i} f(\mathbf{x}_{i}) - \alpha_{i} k(\mathbf{x}_{i}, \mathbf{x}_{i})) > 0 \right\}}.$$
(1.84)

The latter can be obtained easily without explicitly solving the optimization problem again for the reduced samples. In particular, for kernels with $k(\mathbf{x}, \mathbf{x}) = 1$ like many RBF kernels the condition reduces to testing whether $y_i f(\mathbf{x}_i) - \alpha_i > 0$. The remaining problem is that $R_{\text{out}}(f)$ itself is a random variable and thus it does not immediately give a *bound* on R(f). See also chapters 6 and 4 for futher details on how to exploit these bounds in practical cases.

1.4 Boosting

[24] proposed the AdaBoost algorithm for combining classifiers produced by other learning algorithms. AdaBoost has been very successful in practical applications (see Section 1.5). It turns out that it is also a large margin technique.

Table 1.2 gives the pseudocode for the algorithm. It returns a convex combination of classifiers from a class G, by using a learning algorithm L that takes as input a training sample X, Y and a distribution D on X (not to be confused with the true distribution p), and returns a classifier from G. The algorithm L aims to minimize training error on X, Y, weighted according to D. That is, it aims to minimize

$$\sum_{i=1}^{m} D_i 1_{\{h(\mathbf{x}_i) \neq y_i\}}. \tag{1.85}$$

AdaBoost iteratively combines the classifiers returned by L. The idea behind AdaBoost is to start with a uniform weighting over the training sample, and progressively adjust the weights to emphasize the examples that have been frequently misclassified by the classifiers returned by L. These classifiers are combined with convex coefficients that depend on their respective weighted errors. The following theorem shows that Adaboost produces a large margin classifier, provided L is successful at finding classifiers with small weighted training error. See [53,]. Recall (1.30) that the margin error of a function f with respect to ρ on a sample X,Y is

argument: Training sample,
$$X = \{\mathbf{x}_1, \dots, \mathbf{x}_m\} \subset \mathcal{X}, \ Y = \{y_1, \dots, y_m\} \subset \{\pm 1\}$$
 Number of iterations, T returns: Convex combination of functions from G , $f = \sum_{t=1}^T \alpha_t g_t$. function AdaBoost (X, Y, T) for all i from $i = 1, \dots, m$
$$D_1(i) := 1/m$$
 endfor for all t from $\{1, \dots, T\}$
$$g_t := L(X, Y, D_t)$$

$$\varepsilon_t := \sum_{i=1}^m D_t(i) 1_{g_t(x_i) \neq y_i}$$

$$\alpha_t := \frac{1}{2} \ln \left(\frac{1-\varepsilon_t}{\varepsilon_t}\right)$$

$$Z_t := 2\sqrt{\varepsilon_t(1-\varepsilon_t)}$$
 for all i from $i = 1, \dots, m$
$$D_{t+1}(i) := \left\{\begin{array}{c} D_t(i) e^{-\alpha_t}/Z_t & \text{if } y_i = g_t(x_i) \\ D_t(i) e^{\alpha_t}/Z_t & \text{otherwise,} \end{array}\right.$$
 endfor endfor return $f = \frac{\sum_{t=1}^T \alpha_t g_t}{\sum_{i=1}^T \alpha_t}$.

Table 1.2 Pseudocode for the Adaboost algorithm. (L is a learning algorithm that chooses a classifier from G to minimize weighted training error.)

$$R_{\rho}(f) = \frac{1}{m} \sum_{i=1}^{m} 1_{\{y_i f(\mathbf{x}_i) < \rho\}}$$

Theorem 1.12 Margin Error of AdaBoost

If, at iteration t, L returns a function with weighted training error $\varepsilon_t < 1/2$, then AdaBoost returns a function f that satisfies

$$R_{\rho}(f) \le 2^{T} \prod_{t=1}^{T} \sqrt{\varepsilon_{t}^{1-\rho} (1-\varepsilon_{t})^{1+\rho}}.$$
(1.86)

In particular, if $\varepsilon_t \leq 1/2 - 2\rho$, then

$$R_{\rho}(f) < (1 - \rho^2)^{T/2},$$
 (1.87)

and this is less than ε for $T > (2/\rho^2) \ln(1/\varepsilon)$.

1.5 Empirical Results, Implementations, and Further Developments

Large margin classifiers are not only promising from the theoretical point of view. They also have proven to be competitive or superior to other learning algorithms in practical applications. In the following we will give references to such situations.

1.5.1 Boosting

Experimental results show that boosting is able to improve the performance of classifiers significantly. Extensive studies on the UC Irvine dataset, carried out by [25] and [49] with tree classifiers show the performance of such methods. However, also other learning algorithms can benefit from boosting. [63] achieve record performance on an OCR task on the UC Irvine database, using neural networks as the base classifiers. See [50] and chapter 7 for further results on the performance of improved versions of boosted classifiers.

1.5.2 Support Vector Machines

SV Machines perform particularly well in feature rich highdimensional problems. [54, 61, 58, 14, 55] achieve state of the art, or even record performance in several Optical Character Recognition (OCR) tasks such as the digit databases of the United Postal Service (USPS) and the National Institute of Standards and Technology (NIST). The latter can be obtained at

http://www.research.att.com/~yann/ocr/mnist/

Similar results have been obtained for face recognition by [44, 46] and object recognition [11, 55,]. Finally, also on large noisy problems SV Machines are very competitive as shown in [69,].

1.5.3 Implementation and Available Code

Whilst Boosting can be easily implemented by combining a base learner and following the pseudocode of table 1.2. Hence one only has to provide a base learning algorithm satisfying the properties of a weak learner, which defers all problems to the underlying algorithm.

http://www.research.att.com/~yoav/adaboost/

provides a Java applet demonstrating the basic properties of AdaBoost.

The central problem in Support Vector Machines is a quadratic programming problem. Unfortunately, off-the-shelf packages developed in the context of mathematical programming like MINOS [43,], LOQO [71,], OSL [31,], or CPLEX [19,] are often prohibitively expensive or unsuitable for optimization problems in more than several thousand variables (whilst the number of variables may be in the tens of thousands in practical applications). Furthermore these programs are often optimized to deal with sparse matrix entries, causing unneeded overhead when solving generic SV optimization problems (which are sparse in the solution, not in the

matrix entries).

This situation led to the development of several quadratic optimization algorithms specifically designed to suit the needs of SV machines. Starting from simple subset selection algorithms as initially described by [72] and subsequently implemented in e.g. [54,], more advanced chunking methods were proposed [45,] (see also [33,] for a detailed description of the algorithm) for splitting up the optimization problem into smaller subproblems that could be easily solved by standard optimization code. Other methods exploit constrained gradient descent techniques [35,], or minimize very small subproblems, such as the Sequential Minimal Optimization algorithm (SMO) by [47]. See also chapter 5 for further methods for training a SV classifier. Implementations include SvmLight by [33],

http://www-ai.cs.uni-dortmund.de/thorsten/svm light.html

the Royal Holloway / ATT / GMD Support Vector Machine by [52], available at

http://svm.dcs.rhbnc.ac.uk/

and the implementation by Steve Gunn which can be downloaded from

http://www.isis.ecs.soton.ac.uk/resources/svminfo/.

The first two of these optimizers use the GMD (Smola) implementation of an interior point code along the lines of [71] as the core optimization engine. It is available as a standalone package at

http://www.svm.first.gmd.de/software.html.

This site will also contain pointers to further toolboxes as they become available. Java applets for demonstration purposes can be found at

http://http://svm.dcs.rhbnc.ac.uk/pagesnew/GPat.shtml http://http://svm.research.bell-labs.com/SVT/SVMsvt.html. 1.6 Notation 27

1.6 Notation

We conclude the introduction with a list of symbols which are used throughout the book, unless stated otherwise.

```
\mathbb{N}
              the set of natural numbers
\mathbb{R}
              the set of reals
X
              a sample of input patterns
Y
              a sample of output labels
\mathcal{X}
              an abstract domain
              logarithm to base e
ln
              logarithm to base 2
\log_2
(\mathbf{x} \cdot \mathbf{x}')
              inner product between vectors \mathbf{x} and \mathbf{x}'
              2-norm (Euclidean distance), \|\mathbf{x}\| := \sqrt{(\mathbf{x} \cdot \mathbf{x})}
||.||
              p\text{-norm} , \|\mathbf{x}\|_p := \left(\sum_{i=1}^N |x_i|^p\right)^{1/p}
||.||_{p}
              \infty-norm, \|\mathbf{x}\|_{\infty} := \max_{i=1}^{N} |x_i|
||.||_{\infty}
\ell_p
              \ell_p metric
L_2(X)
              space of functions on X square integrable wrt. Borel-Lebesgue measure
\mathbf{E}(\xi)
              expectation of random variable \xi
Pr(\cdot)
              probability of an event
N
              dimensionality of input space
              number of training examples
m
\mathbf{x}_{i}
              input patterns
              target values, or (in pattern recognition) classes
              weight vector
W
              constant offset (or threshold)
b
              VC dimension
h
              a real valued function f: \mathbb{R}^N \to \mathbb{R} (unthresholded)
              a family of real valued functions f
F
              a decision function g: \mathbb{R}^N \to \{-1, 1\}
g
F
              a family of decision functions g
              margin of function f on the example (\mathbf{x}, y), i.e. y f(\mathbf{x})
\rho_f(\mathbf{x}, y)
              minimum margin, i.e. \min_{1 \leq i \leq m} \rho_f(\mathbf{x}_i, y_i)
\rho_f
```

$c(\mathbf{x}, y, f(\mathbf{x}))$	cost function
R(g)	risk of g , i.e. expected fraction of errors
$R_{\mathrm{emp}}(g)$	empirical risk of g , i.e. fraction of training errors
R(f)	risk of f
$R_{ m emp}(f)$	empirical risk of f
k	Mercer kernel
$\mathcal F$	Feature space induced by a kernel
Φ	map into feature space (induced by k)
α_i	Lagrange multiplier
α	vector of all Lagrange multipliers
ξ_i	slack variables
ξ	vector of all slack variables
C	regularization constant for SV Machines
λ	regularization constant $(C = \frac{1}{\lambda})$

I The Key Insight

2 Successful Paper Writing with Support Vector Machines and Other Classifiers

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We believe that Large Margin Classifiers offer a great opportunity to generate a serious amount of publications that would have ended up in the trashcan otherwise. Therefore we think that it is of utmost importance to keep the field growing and foster further research in this area.

2.1 Introduction

general blurb

Besides the correct way of quoting previous results [1, 13, 17, 77, 78, 79, 80, 81, 82, 83, 84, 85, 72, 74, 73, 76, 75] one has to keep in mind not to forget the important issue of finding the right trend in machine learning. Hence one will have to fact the fact that sometimes trends do not last too long, hence one has to write even faster than expected.

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3 Generalized Support Vector Machines

4 Gaussian Process Classification and SVM: Mean Field Results

5 ... sv training algorithm ...

6 Leave-One-Out Machines

7 DOOM2

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