Spectral Algorithms for Supervised Learning

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We discuss how a large class of regularization methods, collectively
known as spectral regularization and originally designed for solving
ill-posed inverse problems, gives rise to regularized learning algorithms.
All of these algorithms are consistent kernel methods that can be easily
implemented. The intuition behind their derivation is that the same
principle allowing for the numerical stabilization of a matrix inversion
problem is crucial to avoid overfitting. The various methods have a com-
mon derivation but different computational and theoretical properties.
We describe examples of such algorithms, analyze their classification
performance on several data sets and discuss their applicability to
real-world problems.

1 Introduction

A large amount of literature has pointed out the connection between algo-
rithms in learning theory and regularization methods in inverse problems
(see, e.g., Vapnik, 1982, 1998; Poggio & Girosi, 1992; Evgeniou, Pontil, &
Poggio, 2000; Hastie, Tibshirani, & Friedman, 2001; Schölkopf & Smola,
The main message is that regularization techniques provide stability with
respect to noise and sampling, therefore ensuring good generalization properties to the corresponding learning algorithms. Usually regularization in learning is based on the minimization of a functional in a suitable hypothesis space—for example, the penalized empirical error on a reproducing kernel Hilbert space. Hence, theoretical analysis focuses mainly on the choice of the loss function, the penalty term, and the kernel (Vapnik, 1998; Evgeniou et al., 2000; Schölkopf & Smola, 2002; Hastie et al., 2001).

From the seminal work of Tikhonov and others (Tikhonov & Arsenin, 1977), regularization has been rigorously defined in the theory of ill-posed inverse problems. In this context, the problem is to invert a linear operator (or a matrix) that might have an unbounded inverse (or a bad condition number). Regularization amounts to replacing the original operator with a bounded operator, namely, the regularization operator (Engl, Hanke, & Neubauer, 1996), whose condition number is controlled by a regularization parameter. The regularization parameter should be chosen according to the noise level in order to ensure stability. Many regularization algorithms are known; Tikhonov and truncated singular value decomposition (TSVD) are probably the most commonly used.

As Bertero and Boccacci (1998) noted, one can also regard regularization from a signal processing perspective introducing the notion of filter. This point of view gives a way of looking constructively at regularization; indeed, each regularization operator can be defined using spectral calculus as a suitable filter on the eigendecomposition of the operator defining the problem. The filter is designed to suppress the oscillatory behavior corresponding to small eigenvalues. In this view, it is known, for example, that Tikhonov regularization can be related to the Wiener filter (Bertero & Boccacci, 1998).

Regularization has a long history in learning, and our starting point is the theoretical analysis proposed in Bauer, Pereverzev, and Rosasco (2007), De Vito, Rosasco, and Verri (2005), Caponnetto and De Vito (2007), and Caponnetto (2006), showing that many regularization methods originally proposed in the context of inverse problems give rise to consistent kernel methods with optimal minimax learning rates. The analysis we propose in this letter focuses on three points.

First, in contrast to Bauer et al. (2007), we propose a more intuitive derivation of regularization based on the notion of spectral filter. We start by introducing the notion of filter functions and explain why, besides ensuring numerical stability, they can also provide a way to learn with generalization guarantees. This requires, in particular, discussing the interplay between filtering and random sampling. Our analysis is complementary to the theory developed in Bauer et al. (2007), De Vito, Rosasco, and Verri (2005), Yao, Rosasco, and Caponnetto (2007), and Caponnetto (2006). Note that the fact that algorithms ensuring numerical stability can also learn is not obvious but confirms the deep connection between stability and generalization (for
references, see Bousquet & Elisseeff, 2002; Poggio, Rifkin, Mukherjee, & Niyogi, 2004; Rakhlin, Mukherjee, & Poggio, 2005).

Second, we present and discuss several examples of filters inducing spectral algorithms for supervised learning. The filter function perspective provides a unifying framework for discussing similarities and differences among the various methods. Some of these algorithms, such as the \(\nu\)-method and iterated Tikhonov, are new to learning. Other algorithms are well known: spectral cut-off (TSVD) is related to principal component regression (PCR) and its kernel version; Landweber iteration is known as L2-boosting (Bühlmann & Yu, 2002), and Tikhonov regularization is also known as regularized least squares or ridge regression. Our analysis highlights the common regularization principle underlying algorithms originally motivated by seemingly unrelated ideas: penalized empirical risk minimization, like regularized least squares; early stopping of iterative procedures, like gradient descent; and (kernel) dimensionality reduction methods, like (kernel) principal component analysis.

Despite these similarities, spectral algorithms have differences from both computational and theoretical points of view. One of the main differences regards the so-called saturation effect affecting some regularization schemes. This phenomenon, which is well known in inverse problem theory, amounts to the impossibility for some algorithms to exploit the regularity of the target function beyond a certain critical value, referred to as the qualification of the method. We try to shed light on this, which is usually not discussed in the literature of learning rates, using some theoretical considerations and numerical simulations.

Another point that differentiates spectral algorithms concerns algorithmic complexity. An interesting aspect is the built-in property of iterative methods to recover solutions corresponding to the whole regularization path (Hastie, Rosset, Tibshirani, & Zhu, 2004).

Third, we evaluate the effectiveness of the proposed algorithms in learning tasks with an extensive experimental study. The performance of these spectral algorithms is assessed on various data sets, and it is compared against state-of-the-art techniques, such as support vector machines (SVMs). The above algorithms often improve the state-of-the-art results and have interesting computational properties, such as the fact that the implementation amounts to a few lines of code. Optimization issues that might be interesting starting points for future work are beyond the scope of this research (see the recent work of Li, Lee, & Leung, 2007, for references).

The plan of the letter is the following. Section 2 discusses previous works in the context of filtering and learning, section 3 reviews the regularized least-squares algorithm from a filter function perspective, and section 4 is devoted to extending the filter point of view to a large class of kernel methods. In section 5 we give several examples of such algorithms and discuss their properties and complexity in section 6. Section 7 reports...
results obtained with an experimental analysis on various data sets, and section 8 contains a final discussion.

2 Previous Work on Learning, Regularization, and Spectral Filtering

The idea of using regularization in statistics and machine learning has long been explored (see, e.g., Wahba, 1990; Poggio & Girosi, 1992), and the connection between large margin kernel methods such as support vector machines and regularization is well known (see Vapnik, 1998; Evgeniou et al., 2000; Schölkopf & Smola, 2002). Ideas coming from inverse problems mostly used Tikhonov regularization and were extended to several error measures other than the quadratic loss function. The gradient descent learning algorithm in Yao et al. (2007) can be seen as an instance of Landweber iteration (Engl et al., 1996) and is related to the L2 boosting algorithm (Bühlmann & Yu, 2002). For iterative methods, some partial results, which do not take into account the random sampling, are presented in Ong and Canu (2004) and Ong, Mary, Canu, and Smola (2004). The interplay of ill-posedness, stability, and generalization is not new to learning (Poggio & Girosi, 1992; Evgeniou et al., 2000; Bousquet & Elisseeff, 2002; Rakhlin et al., 2005; Poggio et al., 2004; De Vito, Rosasco, Caponnetto, et al., 2005).

The notion of filter function was previously studied in machine learning and gives a connection to the literature of function approximation in signal processing and approximation theory. The pioneering work of Poggio and Girosi (1992) established the relation between neural networks, radial basis functions, and regularization. Though the notion of reproducing kernel is not explicitly advocated, Green functions of second-order differential operator are used to define penalties for penalized empirical risk minimization. Filtering and reweighing of the Fourier transform of Green functions are used de facto to design new kernels (see also for the relation between kernel and penalty term in Tikhonov regularization). Poggio and Girosi (1992), as well as Girosi, Jones, and Poggio (1995), are suitable sources for reference and discussion. An important aspect that we would like to stress is that from a technical point of view, these works implicitly assume the data to be sampled according to a uniform distribution and make extensive use of Fourier theory. Indeed the extension to general probability distribution is not straightforward, and this is crucial since it is standard in learning theory to assume the point to be drawn according to a general, unknown distribution. A mathematical connection between sampling theory and learning theory has been recently proposed by Smale and Zhou (2004, 2005) whereas De Vito, Rosasco, Caponnetto, et al. (2005) and De Vito et al. (2006) gave an inverse problem perspective on learning. The analysis we present can be seen as a further step toward a deeper understanding of learning as a function approximation problem.

Recently filtering of the kernel matrix has been considered in the context of graph regularization (see Hastie et al., 2001; Chapelle, Weston, &
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Schölkopf, 2003; Zhu, Kandola, Ghahramani, & Lafferty, 2005; Smola & Kondor, 2003; Zhang & Ando, 2006). In this case, reweighing of a kernel matrix (filters) on a set of labeled and unlabeled input points is used to define new penalty terms, replacing the square of the norm in the adopted hypothesis space. It has been shown (see, e.g., Zhang & Ando, 2006) that this is equivalent to standard regularized least squares with data-dependent kernels. Note that in graph regularization, no sampling is considered, and the problem is truly a problem of transductive learning.

Our analysis relies on a different use of filter functions to define new algorithms rather than the new kernels. In fact, in our setting, the kernel is fixed, and each rescaling of the kernel matrix leads to a learning algorithm, which is not necessarily a penalized minimization. The dependence of the rescaling on the regularization parameter allows us to derive consistency results in a natural way.

3 Regularized Least Squares as a Spectral Filter

In this section we review how the generalization property of the regularized least-squares algorithm is a consequence of the algorithm being seen as a filter on the eigenvalues of the kernel matrix. This point of view naturally suggests a new class of learning algorithms defined in terms of filter functions, whose properties are discussed in the next section.

In the framework of supervised learning, the regularized least-squares algorithm is based on the choice of a Mercer kernel $K(x, t)$ on the input space $X$ and a regularization parameter $\lambda > 0$. Hence, for each training set $z = (x, y) = ((x_1, y_1), \ldots, (x_n, y_n))$ of $n$-examples $(x_i, y_i) \in X \times \mathbb{R}$, regularized least squares amounts to

$$f_\lambda^z(x) = \sum_{i=1}^{n} \alpha_i K(x, x_i) \quad \text{with} \quad \alpha = (K + n\lambda I)^{-1}y,$$  \hspace{1cm} (3.1)

where $K$ is the $n \times n$-matrix $(K)_{ij} = K(x_i, x_j)$.

Since $\lambda > 0$, it is clear that we are numerically stabilizing a matrix inversion problem that is possibly ill conditioned (i.e., numerically unstable). Before showing that regularized least squares can be seen as a suitable filtering of the kernel matrix, able to ensure good generalization properties of the estimator, we first need to recall some basic concepts of learning theory.

We assume that the examples $(x_i, y_i)$ are drawn identically and independently distributed according to an unknown probability measure $\rho(x, y) = \rho(y | x)\rho_X(x).$ Moreover, we assume that $X$ is a compact subset of $\mathbb{R}^d$, and the labels $y_i$ belong to a bounded subset $Y \subset \mathbb{R}$ (e.g., in a binary classification problem $Y = \{-1, 1\}$). Finally, we assume that the kernel $K$ is symmetric continuous function, which is positive definite (Aronszajn, 1950).
is bounded by 1 and is universal (see Micchelli, Xu, & Zhang, 2006, and references therein), that is, the set of functions

$$\mathcal{H} = \left\{ \sum_{i=1}^{N} \alpha_i K(x, x_i) \mid x_i \in X, \alpha_i \in \mathbb{R} \right\}$$

is dense in $L^2(X)$, the Hilbert space of functions that are square-integrable with respect to $\rho_X$.

With the choice of the square loss, the generalization property of the estimator means that the estimator $f^\lambda_z$ is a good approximation of the regression function,

$$f_\rho(x) = \int_Y y \, d\rho(y \mid x),$$

with respect to the norm of $L^2(X)$. In particular, the algorithm is (weakly) consistent (Vapnik, 1998) if, for a suitable choice of the parameter $\lambda = \lambda_n$ as a function of the examples,

$$\lim_{n \to \infty} \int_X \left( f^\lambda_z(x) - f_\rho(x) \right)^2 \, d\rho_X(x) = \lim_{n \to \infty} \| f^\lambda_z - f_\rho \|_\rho^2 = 0,$$

with high probability (see, e.g., Vapnik, 1998).

Notice that in classification, the goal is to approximate the Bayes rule sign

$$(f_\rho) = \text{sign}(\rho(1 \mid x) - 1/2)$$

with the plug-in estimator sign($f^\lambda_z$) with respect to the classification error $R(f^\lambda_z) = P(y f^\lambda_z(x) < 0)$. In any case the following bound holds,

$$R(f^\lambda_z) - R(f_\rho) \leq \| f^\lambda_z - f_\rho \|_\rho$$

(see, e.g., Bartlett, Jordan, & McAuliffe, 2006), so in the following discussion we consider only the square loss.

We start by rewriting equation 3.1 in a slightly different way:

$$(f^\lambda_z(x_1), \ldots, f^\lambda_z(x_n)) = \frac{K}{n} \left( \frac{K}{n} + \lambda \right)^{-1} y. \quad (3.3)$$

Observe that if $v$ is an eigenvector of $K/n$ with eigenvalue $\sigma$, then we have $K/n (K/n + \lambda)^{-1} v = \frac{\sigma}{\sigma + \lambda} v$, so that the regularized least-squares algorithm is in fact a filter on the eigenvalues of the kernel matrix.

The filter $\frac{\sigma}{\sigma + \lambda}$ ensures not only numerical stability but also the generalization properties of the estimator. To obtain insight into this point, consider the population case when we have knowledge of the probability
distribution $\rho$ generating the data. In this setting, the kernel matrix $K/n$ is replaced by the integral operator $L_K$ with kernel $K$,

$$L_K f(x) = \int_X K(x, s) f(s) d\rho_X(s) \quad f \in L^2(X),$$

(3.4)

and the data $y$ is replaced by the regression function $f_\rho$, so that equation 3.3 becomes

$$f^\lambda = L_K (L_K + \lambda I)^{-1} f_\rho.$$  

(3.5)

More explicitly, since $L_K$ is a positive compact operator bounded by 1 and $H$ is dense in $L^2(X)$, there is basis $(u_i)_{i \geq 1}$ in $L^2(X)$ such that $L_K u_i = \sigma_i u_i$ with $0 < \sigma_i \leq 1$ and $\lim_{i \to \infty} \sigma_i = 0$. Hence,

$$f_\rho = \sum_{i=1}^{\infty} \langle f_\rho, u_i \rangle_\rho u_i$$

and

$$f^\lambda = \sum_{i=1}^{\infty} \frac{\sigma_i}{\sigma_i + \lambda} \langle f_\rho, u_i \rangle_\rho u_i.$$

A comparison of the two equations shows that $f^\lambda$ is a good approximation of $f_\rho$ provided that $\lambda$ is small enough. For such $\lambda$, the filter $\frac{\sigma}{\sigma + \lambda}$ selects only the components of the $f_\rho$ corresponding to large eigenvalues, which are a finite number since the sequence of eigenvalues goes to zero. Hence, if we slightly perturb both $L_K$ and $f_\rho$, the corresponding solution of equation 3.4 is close to $f_\rho$ provided that the perturbation is small. The key idea is that now we can regard the sample case $K, y$ and the corresponding estimator $f^\lambda$, as a perturbation of $L_K, f_\rho$, and $f^\lambda$, respectively. A mathematical proof of the above intuition requires some work, and we refer to De Vito, Rosasco, and Verri (2005) and Bauer et al. (2007) for the technical details. The basic idea is that the law of large numbers ensures that the perturbation is small, provided that the number of examples is large enough and, as a consequence, $f^\lambda$ is close to $f_\rho$ and hence to $f_\rho$.

The discussion suggests that one can replace $\frac{\sigma}{\sigma + \lambda}$ with other functions $\sigma g_\lambda(\sigma)$ that are filters on the eigenvalues of $L_K$ and obtain different regularization algorithms, as shown in the next section.

### 4 Kernel Methods from Spectral Filtering

In this section we discuss the properties of kernel methods based on spectral filtering. Our approach is inspired by inverse problems. A complete theoretical discussion of our approach can be found in De Vito, Rosasco, and Verri (2005), Bauer et al. (2007), and Caponnetto (2006).
Let $K$ be a Mercer kernel as in the above section. Looking at equation 3.1 suggests defining a new class of learning algorithm by letting

$$f^\lambda_Z(x) = \sum_{i=1}^{n} \alpha_i K(x, x_i) \quad \text{with} \quad \alpha = \frac{1}{n} g^\lambda \left( \frac{K}{n} \right) y,$$

where $g^\lambda: [0, 1] \to \mathbb{R}$ is a suitable function and $g^\lambda \left( \frac{K}{n} \right)$ is defined by spectral calculus; that is, if $v$ is an eigenvector of $K/n$ with eigenvalue $\sigma$ (since $K$ is a Mercer kernel bounded by 1, $0 \leq \sigma \leq 1$), then $g^\lambda \left( \frac{K}{n} \right) v = g^\lambda(\sigma) v$. In particular, on the given data, one has

$$(f^\lambda_Z(x_1), \ldots, f^\lambda_Z(x_n)) = \frac{K}{n} g^\lambda \left( \frac{K}{n} \right) y. \quad (4.2)$$

We note that unlike regularized least squares, such an estimator is not necessarily the solution of penalized empirical minimization. Clearly, to ensure both numerical stability and consistency, we need to make some assumptions on $g^\lambda$. Following De Vito, Rosasco, and Verri (2005) and Bauer et al. (2007), we say that a function $g^\lambda : [0, 1] \to \mathbb{R}$ parameterized by $0 < \lambda \leq 1$ is an admissible filter function if:

1. There exists a constant $B$ such that
   $$\sup_{0 < \sigma \leq 1} |g^\lambda(\sigma)| \leq \frac{B}{\lambda} \quad \forall \lambda \in [0, 1]. \quad (4.3)$$

2. There exists a constant $D$ such that
   $$\lim_{\lambda \to 0} \sigma g^\lambda(\sigma) = 1 \quad \forall \sigma \in [0, 1]$$
   $$\sup_{0 < \sigma \leq 1} |\sigma g^\lambda(\sigma)| \leq D \quad \forall \lambda \in [0, 1]. \quad (4.4)$$

3. There is a constant $\nu > 0$, namely, the qualification of the regularization $g^\lambda$, such that
   $$\sup_{0 < \sigma \leq 1} |1 - g^\lambda(\sigma)\sigma^{v} \sigma^{\nu} \leq \gamma_v \lambda^{\nu}, \quad \forall \ 0 < \nu \leq \nu,$$

where the constant $\gamma_v > 0$ does not depend on $\lambda$.

A simple computation shows that $g^\lambda(\sigma) = \frac{1}{\sigma + \lambda}$ is an admissible filter function; indeed equations 4.3 and 4.4 hold with $B = D = 1$, condition 4.5 is verified with $\gamma_v = 1$ for $0 < v \leq 1$, and hence the qualification equals 1. Other examples are discussed in the next section. Here we give a heuristic motivation of the above conditions, keeping in mind the discussion in the
previous section. First, observe that the population version of equation 4.1 becomes

$$f^\lambda = \sum_i \sigma_i g_{\lambda}(\sigma_i) \langle f_{\rho}, u_i \rangle_{\rho} u_i.$$ \hspace{1cm} (4.6)

We can make the following observations.

1) Equation 4.3 ensures that eigenvalues of $g_{\lambda}(K)$ are bounded by $\frac{B}{\lambda}$, so that equation 4.1 is numerically stable. Moreover, looking at equation 4.6, we see that it also implies that if $\sigma_i$ is much smaller than $\lambda$, the corresponding Fourier coefficient $\langle f^\lambda u_i \rangle_{\rho}$ is small. Hence, $f^\lambda$ has essentially only a finite number of nonzero Fourier coefficients on the basis $(u_i)_{i \geq 1}$, and we can argue that by the law of large numbers, $f^\lambda_z$ is a good approximation of $f^\lambda$ when $n$ is large enough.

2) Assumption 4.4 implies that $f^\lambda$ converges to $f_{\rho}$ if $\lambda$ goes to zero. In terms of the kernel matrix, this condition means that $g_{\lambda}(K)$ converges to $K^{-1}$ when $\lambda$ goes to zero, avoiding oversmoothing.

3) Condition 4.5 is related to the convergence rates of the algorithm. These rates depend on how fast the Fourier coefficients $\langle f_{\rho}, u_i \rangle_{\rho}$ converge to 0 with respect to the eigenvalues $\sigma_i$ (Bauer et al., 2007). This information is encoded by a priori assumptions on $f_{\rho}$ of the form

$$\sum_{i=1}^{\infty} \frac{\langle f_{\rho}, u_i \rangle_{\rho}^2}{\sigma_i^{2r}} < R,$$ \hspace{1cm} (4.7)

where the parameter $r$ encodes the regularity property of the regression function. If $r = 1/2$, this corresponds to assuming $f_{\rho} \in \mathcal{H}$ and, more generally, the larger is $r$, the smoother is the function. Condition 4.5 and the choice $\lambda_n = \frac{1}{n^{\nu+\tau}}$ ensure that if $r \leq \nu$,

$$\|f^\lambda_z - f_{\rho}\|_{\rho} \leq Cn^{-\frac{r}{\nu+\tau}} \text{ with high probability},$$ \hspace{1cm} (4.8)

whereas if $r \geq \nu$, the rate of convergence is always $n^{-\frac{r}{\nu+\tau}}$ (for proof and a complete discussion, see Bauer et al., 2007; Caponnetto, 2006). Hence, filter functions having a larger qualification $\nu$ give better rates, that is, the corresponding algorithms can better exploit the smoothness of $f_{\rho}$. This fact marks a big distinction among the various algorithms we consider, as we discuss in the following. Also, notice that in the classification setting we will consider in our experiments, if $r \leq \nu$ bounds 3.2 and equation 4.8 give that

$$\mathcal{R}(f^\lambda_z) - \inf_f \mathcal{R}(f) = O(n^{-\frac{r}{\nu+\tau}})$$

with high probability.
Considering the decomposition \( f^\lambda_z - f^\rho = (f^\lambda_z - f^\lambda) + (f^\lambda - f^\rho) \), from the above discussion, we have that the consistency of this class of learning algorithms depends on two opposite terms: the approximation error \( \| f^\lambda - f^\rho \| \) and the sample error \( \| f^\lambda_z - f^\lambda \| \). The approximation error depends on the examples only through \( \lambda = \lambda_n \), and it decreases if \( \lambda \) goes to zero, whereas the sample error is of a probabilistic nature and it increases if \( \lambda \) goes to zero. The optimal choice of the regularization parameter \( \lambda \) will be a trade-off between these two errors (see De Vito, Caponnetto, & Rosasco, 2005; Caponnetto & De Vito, 2006; Smale & Zhou, 2007; Wu, Ying, & Zhou, 2006), about the rates for regularized least-squares. See De Vito, Rosasco, and Verri (2005), Bauer et al. (2007), and Caponnetto (2006), for arbitrary filters.

Before giving several examples of algorithms that fit into the above general framework, we observe that the considered algorithms can be regarded as filters on the expansion of the target function on a suitable basis. In principle, this basis can be obtained from the spectral decomposition of the integral operator \( L_K \) and in practice is approximated by considering the spectral decomposition of the kernel matrix \( K \). Interestingly, the basis thus obtained has a natural interpretation: if the data are centered (in the feature space), then the elements of the basis are the principal components of the expected (and empirical) covariance matrix in the feature space. In this respect, the spectral methods we discussed rely on the assumption that most of the information is actually encoded in the first principal components.

5 The Proposed Algorithms

In this section we give specific examples of kernel methods based on spectral regularization. All of these algorithms are known in the context of regularization for linear inverse problems, but only some of them have been used for statistical inference problems. These methods have many interesting features: from the algorithmic point of view, they are simple to implement; usually they amount to a few lines of code. They are appealing for applications: their model selection is simple since they depend on few parameters, and overfitting may be dealt with in a very transparent way. Some of them represent a very good alternative to regularized least squares (RLS) because they are faster without compromising classification performance (see section 7). Note that for regularized least squares, the algorithm has the following variational formulation,

\[
\min_{f \in \mathcal{H}} \frac{1}{n} \sum_{i=1}^{n} (y_i - f(x_i))^2 + \lambda \| f \|_{\mathcal{H}}^2 ,
\]

which can be interpreted as an extension of empirical risk minimization. In general, the class of regularization might not be described by a
variational problem so that the filter point of view provides us with a suitable description.

More details on the derivation of these algorithms can be found in Engl et al. (1996).

5.1 Iterative Landweber. Landweber iteration is characterized by the filter function

$$g_t(\sigma) = \tau \sum_{i=0}^{t-1} (1 - \tau \sigma)^i,$$

where we identify $\lambda = t^{-1}, t \in \mathbb{N}$ and take $\tau = 1$ (since the kernel is bounded by 1). In this case, we have $B = D = 1$, and the qualification is infinite since equation 4.5 holds with $\gamma_\nu = 1$ if $0 < \nu \leq 1$ and $\gamma_\nu = \nu^n$ otherwise. The above filter can be derived from a variational point of view. In fact, as shown in Yao et al. (2007), this method corresponds to empirical risk minimization via gradient descent. If we denote with $\|\cdot\|_n$ the norm in $\mathbb{R}^n$, we can impose

$$\nabla \|K\alpha - y\|_n^2 = 0,$$

and by a simple calculation, we see that the solution can be rewritten as the following iterative map,

$$\alpha_i = \alpha_{i-1} + \frac{\tau}{n} (y - K\alpha_{i-1}), \quad i = 1, \ldots, t,$$

where $\tau$ determines the step size. We may start from a very simple solution: $\alpha_0 = 0$. Clearly, if we let the number of iterations grow, we are simply minimizing the empirical risk and are bound to overfit. Early stopping of the iterative procedure allows us to avoid overfitting; thus, the iteration number plays the role of the regularization parameter. In Yao et al. (2007), the fixed step size $\tau = 1$ was shown to be the best choice among the variable step size $\tau = \frac{1}{(t+1)^{\theta}}$, with $\theta \in [0, 1)$. This suggests that $\tau$ does not play any role in regularization. Landweber regularization was introduced under the name of L2-boosting for splines in a fixed design statistical model (Bühlmann & Yu, 2002) and eventually generalized to general RKH spaces and random design in Yao et al. (2007).

5.2 Semi-Iterative Regularization. An interesting class of algorithms consists of the so-called semi-iterative regularization or accelerated Landweber iteration. These methods can be seen as a generalization of Landweber iteration where the regularization is now

$$g_t(\sigma) = p_t(\sigma),$$
with $p_t$, a polynomial of degree $t - 1$. In this case, we can identify $\lambda = t^{-2}$, $t \in \mathbb{N}$. One can show that $D = 1$, $B = 2$, and the qualification of this class of methods is usually finite (Engl et al., 1996).

An example that turns out to be particularly interesting is the so-called $\nu$ method. The derivation of this method is fairly complicated and relies on the use of orthogonal polynomials to obtain acceleration of the standard gradient descent algorithm (see Chapter 10 in Golub & Van Loan, 1996). Such a derivation is beyond the scope of this presentation and we refer interested readers to Engl et al. (1996). In the $\nu$ method, the qualification is $\nu$ (fixed) with $\gamma_\nu = c$ for some positive constant $c$. The algorithm amounts to solving (with $\alpha_0 = 0$) the following map,

$$\alpha_i = \alpha_{i-1} + u_i(\alpha_{i-1} - \alpha_{i-2}) + \frac{\omega_i}{n}(y - K\alpha_{i-1}), \quad i = 1, \ldots, t,$$

where

$$u_i = \frac{(i - 1)(2i - 3)(2i + 2\nu - 1)}{(i + 2\nu - 1)(2i + 4\nu - 1)(2i + 2\nu - 3)} \quad \omega_i = 4 \frac{(2i + 2\nu - 1)(i + \nu - 1)}{(i + 2\nu - 1)(2i + 4\nu - 1)} \quad t > 1.$$

The interest of this method lies in the fact that since the regularization parameter here is $\lambda = t^{-2}$, we just need the square root of the number of iterations needed by Landweber iteration. In inverse problems, this method is known to be extremely fast and is often used as a valid alternative to conjugate gradient (see Engl et al., 1996, Chap. 6, for details). To our knowledge, semi-iterative regularization has not been previously used in learning.

5.3 Spectral Cut-Off. This method, also known as truncated singular values decomposition (TSVD), is equivalent to the so-called (kernel) principal component regression. The filter function is simply

$$g_\lambda(\sigma) = \begin{cases} 1 & \sigma \geq \lambda, \\ 0 & \sigma < \lambda. \end{cases}$$

In this case, $B = D = 1$. The qualification of the method is arbitrary, and $\gamma_\nu = 1$ for any $\nu > 0$. The corresponding algorithm is based on the following simple idea. Perform SVD of the kernel matrix $K = USU^T$, where $U$ is an orthogonal matrix and $S = \text{diag}(\sigma_1, \ldots, \sigma_n)$ is diagonal with $\sigma_i \geq \sigma_{i+1}$. Then
discard the singular values smaller than the threshold $\lambda$, and replace them with 0. The algorithm is then given by

$$a = K^{-1}_\lambda y,$$

(5.1)

where $K^{-1}_\lambda = U^T S^{-1}_\lambda U$ and $S^{-1}_\lambda = \text{diag}(1/\sigma_1, \ldots, 1/\sigma_m, 0, \ldots)$ where $\sigma_m \geq \lambda$ and $\sigma_{m+1} < \lambda$. The regularization parameter is the threshold $\lambda$ or, equivalently, the number $m$ of components that we keep.

Finally, notice that if the data are centered in the feature space, then the columns of the matrix $U$ are the principal components of the covariance matrix in the feature space, and the spectral cut-off is a filter that discards the projection on the last principal components. The procedure is well known in literature as kernel principal component analysis (see, e.g., Schölkopf & Smola, 2002).

5.4 Iterated Tikhonov. We conclude this section by mentioning a method that is a mixture between Landweber iteration and Tikhonov regularization. Unlike Tikhonov regularization, which has finite qualification and cannot exploit the regularity of the solution beyond a certain regularity level, iterated Tikhonov overcomes this problem by means of the following regularization:

$$g_\lambda(\sigma) = \frac{(\sigma + \lambda)^v - \lambda^v}{\sigma(\sigma + \lambda)^v}, \quad v \in \mathbb{N}.$$

In this case, we have $D = 1$ and $B = t$, and the qualification of the method is now $v$ with $\gamma_v = 1$ for all $0 < v \leq t$. The algorithm is described by the following iterative map,

$$(K + n\lambda I)a_i = y + n\lambda a_{i-1} \quad i = 1, \ldots, v,$$

choosing $a_0 = 0$. It is easy to see that for $v = 1$, we simply recover the standard Tikhonov regularization, but as we let $v > 1$, we improve the qualification of the method with respect to standard Tikhonov. Moreover, we note that by fixing $\lambda$, we can think of the above algorithms as an iterative regularization with $v$ as the regularization parameter.

6 Different Properties of Spectral Algorithms

In this section, we discuss the differences of the proposed algorithms from theoretical and computational viewpoints.

6.1 Qualification and Saturation Effects in Learning. As we mentioned in section 4, one of the main differences among the various spectral methods
is their qualification. Each spectral regularization algorithm has a critical value (the qualification) beyond which learning rates no longer improve despite the regularity of the target function $f_\rho$. If this is the case, we say that methods saturate. In this section, we recall the origin of this problem and illustrate it with some numerical simulations.

Saturation effects have their origin in analytical and geometrical properties rather than in statistical properties of the methods. To see this, recall the error decomposition $f_\lambda^\rho - f_\rho = (f_\lambda^\rho - f_\lambda) + (f_\lambda - f_\rho)$, where the latter term is the approximation error that, recalling equation 4.6, is related to the behavior of

$$f_\rho - f_\lambda = \sum_i (f_\rho, u_i)_\rho u_i - \sum_i \sigma_i g_\lambda(\sigma_i) (f_\rho, u_i)_\rho u_i$$

$$= \sum_i (1 - \sigma_i g_\lambda(\sigma_i)) \sigma_i^r \frac{(f_\rho, u_i)_\rho}{\sigma_i^r} u_i.$$  \hfill (6.1)

If the regression function satisfies equation 4.7, we have

$$\|f_\rho - f_\lambda\|_\rho \leq R \sup_{0 < \sigma \leq 1} ((1 - g_\lambda'(\sigma)) \sigma^r).$$

The above formula clearly motivates condition 4.5 and the definition of qualification. In fact, it follows that if $r \leq \bar{v}$, then $\|f_\rho - f_\lambda\|_\rho = O(\lambda^\bar{v})$, whereas if $r > \bar{v}$, we have $\|f_\rho - f_\lambda\|_\rho = O(\lambda^\bar{v})$. To avoid confusion, note that the index $r$ in the equation 6.1 encodes a regularity property of the target function, whereas $\bar{v}$ in equation 4.5 encodes a property of the given algorithm.

In Figure 1, we show the behaviors of the residual $(1 - \sigma g_\lambda(\sigma)) \sigma^r$ as a function of $\sigma$ for different values of $r$ and fixed $\lambda$. For Tikhonov regularization (see Figure 1, left), in the two top plots, where $r < 1$, the
Figure 2: The behaviors of the approximation errors for Tikhonov regularization (left) and TSVD (right) as a function of $\lambda$ for different values of $r$.

maximum of the residual changes and is achieved within the interval $0 < \sigma < 1$, whereas in the two bottom plots, where $r \geq 1$, the maximum of the residual remains the same and is achieved for $\sigma = 1$. For TSVD (see Figure 1, right), the maximum of the residual changes for all the values of the index $r$ and is always achieved at $\sigma = \lambda$. An easy calculation shows that the behavior of iterated Tikhonov is the same as Tikhonov, but the critical value is now $\nu$ rather than 1. Similarly one can recover the behavior of the $\nu$ method and Landweber iteration.

In Figure 2 we show the corresponding behavior of the approximation error as a function of $\lambda$ for different values of $r$. Again the difference between finite (Tikhonov) and infinite (TSVD) qualification is apparent. For Tikhonov regularization (see Figure 2, right), the approximation error is $O(\lambda^r)$ for $r < 1$ (see the two top plots) and $O(\lambda)$ for $r \geq 1$ (the plots for $r = 1$ and $r = 2$ overlap) since the qualification of the method is 1. For TSVD (Figure 2, left), the approximation error is always $O(\lambda^r)$ since the qualification is infinite. Again similar considerations can be done with iterated Tikhonov as well as for the other methods.

To further investigate the saturation effect, we consider a regression toy problem and evaluate the effect of finite qualification on the expected error. Clearly this is more difficult since the effect of noise and sampling contributes to the error behavior through the sampling error as well. In our toy example, $X$ is simply the interval $[0, 1]$ endowed with the uniform probability measure $d\rho_X(x) = dx$. The hypotheses space $\mathcal{H}$ is the Sobolev space of absolutely continuous, with square integrable first derivative and boundary condition $f(0) = f(1) = 0$. This is a Hilbert space of function endowed with the norm

$$\|f\|^2_{\mathcal{H}} = \int_0^1 |f''(x)|^2 \, dx$$
and it can be shown to be a reproducing Kernel Hilbert (RKH) space with kernel

$$K(x, s) = \Theta(x \geq s)(1 - x)s + \Theta(x \leq s)(1 - s)x,$$

where $\Theta$ is the Heaviside step function. In this setting we compare the performance of spectral regularization methods in two different learning tasks. In both cases, the output is corrupted by gaussian noise. The first task is to recover the regression function given by $f_\rho(x) = K(x_0, x)$ for a fixed point $x_0$ given a priori, and the second task is to recover the regression function $f_\rho(x) = \sin(x)$. The two cases should correspond roughly to $r = 1/2$ and $r \gg 1$. In Figure 3, we show the behavior, for various training set sizes, of

$$\Delta(n) = \min_\lambda \| f_\rho - f_\lambda \|^2_\rho,$$

where we took a sample of cardinality $N \gg n$ to approximate $\| f \|^2_\rho$ with $\frac{1}{N} \sum_{i=1}^{N} | f(x_i) |^2$. We considered 70 repeated trials and show the average learning rates plus and minus one standard deviation. The results in Figure 3 confirm the presence of a saturation effect. For the first learning task (top) the learning rates of Tikhonov and TSVD are essentially the same, but TSVD has better learning rates than Tikhonov in the second learning task (bottom), where the regularity is higher. We performed similar simulations, not reported here, comparing the learning rates for Tikhonov and iterated Tikhonov regularization recalling that the latter has a higher qualification. As expected, iterated Tikhonov has better learning rates in the second learning task and essentially the same learning rates in the first task. Interestingly, we found the real behavior of the error to be better than the one expected from the probabilistic bound, and we conjecture that this behavior is due to the fact that our bound on the sample error is too crude.

### 6.2 Algorithmic Complexity and Regularization Path

In this section we comment on the properties of spectral regularization algorithms in terms of algorithmic complexity.

Having in mind that each of the algorithms we discuss depends on at least one parameter, we are going to distinguish between (1) the computational cost of each algorithm for one fixed parameter value and (2) the computational cost of each algorithm to find the solution corresponding to many parameter values. The first situation corresponds to the case when a

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2In general, besides the regularization parameter, there might be some kernel parameter. In our discussion, we assume the kernel (and its parameters) to be fixed.
Figure 3: Comparisons of the learning rates for Tikhonov regularization and TSVD on two learning tasks with very different regularity indexes. In the first learning task (top), the regression function is less regular than in the second learning task (bottom). The continuous plots represent the average learning rates over 70 trials, while the dashed plots represent the average learning rates plus and minus one standard deviation.

Correct value of the regularization parameter is given a priori or has already been computed. The complexity analysis in this case is fairly standard, and we compute it in a worst-case scenario, though for nicely structured kernel matrices (e.g., sparse or block structured), the complexity can be drastically reduced.
The second situation is more interesting in practice since one usually has to find a good parameter value; therefore, the real computational cost includes the parameter selection procedure. Typically one computes solutions corresponding to different parameter values and then chooses the one minimizing some estimate of the generalization error, for example, hold-out or leave-one-out estimates (Hastie et al., 2001). This procedure is related to the concept of the regularization path (Hastie et al., 2004). Roughly speaking, the regularization path is the sequence of solutions, corresponding to different parameters, that we need to compute to select the best parameter estimate. Ideally one would like the cost of calculating the regularization path to be as close as possible to that of calculating the solution for a fixed parameter value. In general, this is a strong requirement, but, for example, SVM algorithm has a step-wise linear dependence on the regularization parameter (Pontil & Verri, 1998), and this can be exploited to efficiently find the regularization path (Hastie et al., 2004).

Given the above premises, in analyzing spectral regularization algorithms, we notice a substantial difference between iterative methods (Landweber and \( \nu \) method) and the others. At each iteration, iterative methods calculate a solution corresponding to \( t \), which is both the iteration number and the regularization parameter (as mentioned above, equal to \( 1/\lambda \)). In this view, iterative methods have the built-in property of computing the entire regularization path. Landweber iteration at each step \( i \) performs a matrix-vector product between \( K \) and \( \alpha_{i-1} \), so that at each iteration, the complexity is \( O(n^2) \). If we run \( t \) iterations, the complexity is then \( O(t \cdot n^2) \). Similar to Landweber iteration, the \( \nu \) method involves a matrix-vector product so that each iteration costs \( O(n^2) \). However, as discussed in section 5, the number of iterations required to obtain the same solution of Landweber iteration is the square root of the number of iterations needed by Landweber (see also Table 2). Such a rate of convergence can be shown to be optimal among iterative schemes (see Engl et al., 1996). In the case of RLS, in general, one needs to perform a matrix inversion for each parameter value that costs in the worst case \( O(n^3) \). Similarly for spectral cut-off, the cost is that of finding the singular value decomposition of the kernel matrix, which is again \( O(n^3) \). Finally we note that computing solutions for different parameter values is in general very costly for a standard implementation of RLS, while for spectral cut-off, one can perform only one SVD. This suggests the use of SVD decomposition also for solving RLS in case a parameter tuning is needed.

7 Experimental Analysis

This section reports experimental evidence on the effectiveness of the algorithms discussed in section 5. We apply them to a number of classification problems, first considering a set of well-known benchmark data and comparing the results we obtain with the ones reported in the literature; then we
Table 1: The Benchmark Data Sets Used: Size (Training and Test), Space Dimension and Number of Splits in Training and Testing.

<table>
<thead>
<tr>
<th>Number Trained</th>
<th>Number Tested</th>
<th>Dimension</th>
<th>Number Resampled</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Banana</td>
<td>400</td>
<td>4900</td>
<td>2</td>
</tr>
<tr>
<td>2. Breast Cancer</td>
<td>200</td>
<td>77</td>
<td>9</td>
</tr>
<tr>
<td>3. Diabetes</td>
<td>468</td>
<td>300</td>
<td>8</td>
</tr>
<tr>
<td>4. Flare Solar</td>
<td>666</td>
<td>400</td>
<td>9</td>
</tr>
<tr>
<td>5. German</td>
<td>700</td>
<td>300</td>
<td>20</td>
</tr>
<tr>
<td>6. Heart</td>
<td>170</td>
<td>100</td>
<td>13</td>
</tr>
<tr>
<td>7. Image</td>
<td>1300</td>
<td>1010</td>
<td>18</td>
</tr>
<tr>
<td>8. Ringnorm</td>
<td>400</td>
<td>7000</td>
<td>20</td>
</tr>
<tr>
<td>9. Splice</td>
<td>1000</td>
<td>2175</td>
<td>60</td>
</tr>
<tr>
<td>10. Thyroid</td>
<td>140</td>
<td>75</td>
<td>5</td>
</tr>
<tr>
<td>11. Titanic</td>
<td>150</td>
<td>2051</td>
<td>3</td>
</tr>
<tr>
<td>12. Twonorm</td>
<td>400</td>
<td>7000</td>
<td>20</td>
</tr>
<tr>
<td>13. Waveform</td>
<td>400</td>
<td>4600</td>
<td>21</td>
</tr>
</tbody>
</table>

consider a more specific application, face detection, analyzing the results obtained with a spectral regularization algorithm and comparing them with SVM, which many authors in the post have applied with success. For these experiments, we consider both a benchmark data set available on the Web and a set of data acquired by a video-monitoring system designed in our lab.

7.1 Experiments on Benchmark Data Sets. In this section we analyze the classification performance of the regularization algorithms on various benchmark data sets. In particular we consider the IDA benchmark, containing one toy data set (banana; see Table 1), and several real data sets (available at http://ida.first.fraunhofer.de/projects/bench/). These data sets have been previously used to assess many learning algorithms, including AdaBoost, RBF networks, SVMs, and kernel projection machines. The benchmarks Web page reports the results obtained with these methods, and we compare them against the results obtained with our algorithms.

For each data set, 100 resamplings into training and test sets are available from the Web site. The structure of our experiments follows the one reported on the benchmarks Web page: we perform parameter estimation with five-fold cross-validation on the first five partitions of the data set; then we compute the median of the five estimated parameters and use it as an optimal parameter for all the resamplings. As for the choice of parameter $\sigma$ (i.e., the standard deviation of the RBF kernel), first we set the value to the average of square distances of training set points of two different resamplings: let it be $\sigma_c$. Then we compute the error on two randomly chosen partitions on the range $[\sigma_c - \delta, \sigma_c + \delta]$ for a small $\delta$ on several values of $\lambda$ and choose the most appropriate $\sigma$. After we select $\sigma$, the parameter $t$ (corresponding to $1/\lambda$) is tuned with 5-CV on the range $[1, \infty]$. Regarding
the choice of the parameter $\nu$ for the $\nu$ method and iterated Tikhonov (where $\nu$ is the number of iteration), we tried different values and obtained very similar results. The saturation effect on real data seemed much harder to spot, and all the errors were very close. In the end, we chose $\nu = 5$ for both methods.

Table 2 shows the average generalization performance (with standard deviation) over the partitions of the data sets. It also reports the parameters $\sigma$ and $l (= 1/\lambda)$ chosen to find the best model. The results obtained with the five methods are very similar, with the exception of Landweber, whose performances are less stable. The $\nu$ method performs very well and converges to a solution in fewer iterations.

From this analysis, we conclude that the $\nu$ method shows the best combination of generalization performance and computational efficiency among the four regularization methods analyzed. We choose it as a representative

Table 2: Comparison of the Five Methods Discussed.

<table>
<thead>
<tr>
<th></th>
<th>Landweber</th>
<th>$\nu$-Method</th>
<th>RLS</th>
<th>TSVD</th>
<th>IT ($\nu = 5$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>11.70 ± 0.68</td>
<td><strong>10.67 ± 0.53</strong></td>
<td>11.22 ± 0.61</td>
<td>11.74 ± 0.63</td>
<td>10.96 ± 0.56</td>
</tr>
<tr>
<td></td>
<td>(116/1)</td>
<td>(70/1)</td>
<td>(350/1)</td>
<td>(301/1)</td>
<td>(141/1)</td>
</tr>
<tr>
<td></td>
<td>(5/2)</td>
<td>(5/2)</td>
<td>(41/2)</td>
<td>(120/2)</td>
<td>(4/2)</td>
</tr>
<tr>
<td>3</td>
<td>23.70 ± 1.80</td>
<td><strong>23.60 ± 1.82</strong></td>
<td>24.40 ± 1.79</td>
<td>24.29 ± 0.2</td>
<td>23.63 ± 1.88</td>
</tr>
<tr>
<td></td>
<td>(18/2)</td>
<td>(11/2)</td>
<td>(400/2)</td>
<td>(300/2)</td>
<td>(10/2)</td>
</tr>
<tr>
<td>4</td>
<td>34.27 ± 1.57</td>
<td><strong>34.25 ± 1.59</strong></td>
<td>34.31 ± 1.607</td>
<td>32.43 ± 0.90</td>
<td>30.92 ± 10.47</td>
</tr>
<tr>
<td></td>
<td>(25/1)</td>
<td>(8/1)</td>
<td>(51/1)</td>
<td>(140/1)</td>
<td>(6/1)</td>
</tr>
<tr>
<td>5</td>
<td>23.20 ± 2.28</td>
<td><strong>23.14 ± 2.34</strong></td>
<td>23.37 ± 2.11</td>
<td>24.67 ± 2.60</td>
<td>23.31 ± 2.24</td>
</tr>
<tr>
<td></td>
<td>(119/3)</td>
<td>(16/3)</td>
<td>(600/3)</td>
<td>(1150/3)</td>
<td>(51/3)</td>
</tr>
<tr>
<td>6</td>
<td>15.94 ± 3.37</td>
<td><strong>15.48 ± 3.25</strong></td>
<td>15.71 ± 3.20</td>
<td>15.58 ± 3.41</td>
<td>15.60 ± 3.41</td>
</tr>
<tr>
<td></td>
<td>(63/12)</td>
<td>(16/12)</td>
<td>(500/12)</td>
<td>(170/12)</td>
<td>(21/12)</td>
</tr>
<tr>
<td>7</td>
<td>6.42 ± 0.82</td>
<td>2.78 ± 0.56</td>
<td><strong>2.68 ± 0.54</strong></td>
<td>2.99 ± 0.48</td>
<td>2.72 ± 0.53</td>
</tr>
<tr>
<td></td>
<td>(7109/1)</td>
<td>(447/2.6)</td>
<td>(17900/2.6)</td>
<td>(28000/2.6)</td>
<td>(20001/2.6)</td>
</tr>
<tr>
<td>8</td>
<td>9.09 ± 0.89</td>
<td>3.09 ± 0.42</td>
<td>4.68 ± 0.7</td>
<td><strong>2.85 ± 0.33</strong></td>
<td>3.83 ± 0.52</td>
</tr>
<tr>
<td></td>
<td>(514/3)</td>
<td>(37/3)</td>
<td>(820/3)</td>
<td>(510/3)</td>
<td>(151/3)</td>
</tr>
<tr>
<td>9</td>
<td>14.71 ± 0.75</td>
<td><strong>10.79 ± 0.67</strong></td>
<td>11.43 ± 0.72</td>
<td>11.67 ± 0.68</td>
<td>10.92 ± 0.72</td>
</tr>
<tr>
<td></td>
<td>(816/6)</td>
<td>(72/6)</td>
<td>(1250/6)</td>
<td>(1400/6)</td>
<td>(501/6)</td>
</tr>
<tr>
<td>10</td>
<td>4.53 ± 2.34</td>
<td>4.55 ± 2.35</td>
<td><strong>4.48 ± 2.33</strong></td>
<td>4.49 ± 2.21</td>
<td>4.59 ± 2.34</td>
</tr>
<tr>
<td></td>
<td>(65/1)</td>
<td>(28/1)</td>
<td>(100/1)</td>
<td>(200/1)</td>
<td>(21/1)</td>
</tr>
<tr>
<td>11</td>
<td>23.53 ± 1.82</td>
<td>22.96 ± 1.21</td>
<td>22.82 ± 1.81</td>
<td><strong>21.28 ± 0.67</strong></td>
<td>20.20 ± 7.17</td>
</tr>
<tr>
<td></td>
<td>(5/1)</td>
<td>(1/1)</td>
<td>(1.19/1)</td>
<td>(12/1)</td>
<td>(1/1)</td>
</tr>
<tr>
<td>12</td>
<td>2.39 ± 0.13</td>
<td><strong>2.36 ± 0.13</strong></td>
<td>2.42 ± 0.14</td>
<td>2.39 ± 0.13</td>
<td>2.56 ± 0.30</td>
</tr>
<tr>
<td></td>
<td>(20/3)</td>
<td>(7/3)</td>
<td>(100/3)</td>
<td>(61/3)</td>
<td>(1/3)</td>
</tr>
<tr>
<td>13</td>
<td>9.53 ± 0.45</td>
<td>9.63 ± 0.49</td>
<td>9.53 ± 0.44</td>
<td><strong>9.77 ± 0.35</strong></td>
<td>9.52 ± 0.44</td>
</tr>
<tr>
<td></td>
<td>(8/3.1)</td>
<td>(12/3.1)</td>
<td>(150/3.1)</td>
<td>(171/3.1)</td>
<td>(21/3.1)</td>
</tr>
</tbody>
</table>

Notes: The average and standard deviation of the generalization error on the 13 data sets (numbered as in Table 1) is reported on top and the value of the regularization parameter and the gaussian width, ($l/\sigma$), in parentheses. The best result for each data set is in boldface type.
Table 3: Comparison of the $v$ Method Against the Best of the Seven Methods Taken from the Benchmark Web Page on the Benchmark Data Sets.

<table>
<thead>
<tr>
<th>Method</th>
<th>Best of Seven</th>
<th>SVM</th>
<th>$v$-Method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Banana</td>
<td>LP_REG-Ada</td>
<td>10.73 ± 0.43</td>
<td>11.53 ± 0.66</td>
</tr>
<tr>
<td></td>
<td>KFD</td>
<td>24.77 ± 4.63</td>
<td>26.04 ± 4.74</td>
</tr>
<tr>
<td>Breast Cancer</td>
<td>KFD</td>
<td>23.21 ± 1.63</td>
<td>23.53 ± 1.73</td>
</tr>
<tr>
<td>Diabetes</td>
<td>SVM-RBF</td>
<td>32.43 ± 1.82</td>
<td>32.43 ± 1.82</td>
</tr>
<tr>
<td>Flare Solar</td>
<td>KFD</td>
<td>23.61 ± 2.07</td>
<td>23.61 ± 2.07</td>
</tr>
<tr>
<td></td>
<td>SVM-RBF</td>
<td>15.95 ± 3.26</td>
<td>15.95 ± 3.26</td>
</tr>
<tr>
<td>German</td>
<td>ADA_REG</td>
<td>2.67 ± 0.61</td>
<td>2.96 ± 0.6</td>
</tr>
<tr>
<td>Heart</td>
<td>ADA_REG</td>
<td>1.58 ± 0.12</td>
<td>1.66 ± 0.2</td>
</tr>
<tr>
<td>Image</td>
<td>ADA_REG</td>
<td>9.50 ± 0.65</td>
<td>10.88 ± 0.66</td>
</tr>
<tr>
<td></td>
<td>KFD</td>
<td>4.20 ± 2.07</td>
<td>4.80 ± 2.19</td>
</tr>
<tr>
<td>Titanic</td>
<td>SVM-RBF</td>
<td>22.42 ± 1.02</td>
<td>22.42 ± 1.02</td>
</tr>
<tr>
<td></td>
<td>KFD</td>
<td>2.61 ± 0.15</td>
<td>2.96 ± 0.23</td>
</tr>
<tr>
<td>Twonorm</td>
<td>KFD</td>
<td>9.86 ± 0.44</td>
<td>9.88 ± 0.44</td>
</tr>
<tr>
<td>Waveform</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note: The middle column shows the results for SVM from the same Web page.

for comparisons with other approaches. Table 3 compares the results obtained with the $v$ method with an SVM with RBF kernel and, for each data set, with the classifier performing best among the seven methods considered on the benchmark page (including RBF networks, AdaBoost and Regularized AdaBoost, Kernel Fisher Discriminant, and SVMs with RBF kernels). The results obtained with the $v$ method compare favorably with the ones achieved by the other methods.

7.2 Experiments on Face Detection. This section reports the analysis we carried out on the problem of face detection in order to evaluate the effectiveness of the $v$ method in comparison to SVMs. The structure of the experiments, including model selection and error estimation, follows the one reported above. The data we consider are image patches. We represent them in the simplest way, unfolding the patch matrix in a one-dimensional vector of integer values—the gray levels. All the images of the two data sets are $19 \times 19$; thus, the size of our data is 361.
Table 4: Average and Standard Deviation of the Classification Error of SVM and \( \nu \)-Method Trained on Training Sets of Increasing Size.

<table>
<thead>
<tr>
<th>Number Trained Plus Number Tested</th>
<th>600 + 1400</th>
<th>700 + 1300</th>
<th>800 + 1200</th>
</tr>
</thead>
<tbody>
<tr>
<td>Classifier</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RBF-SVM</td>
<td>2.41 ± 1.39</td>
<td>1.99 ± 0.82</td>
<td>1.60 ± 0.71</td>
</tr>
<tr>
<td>( \sigma = 800 ) ( C = 1 )</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>( \nu )-method</td>
<td>1.63 ± 0.32</td>
<td>1.53 ± 0.33</td>
<td>1.48 ± 0.34</td>
</tr>
<tr>
<td>( \sigma = 341 ) ( t = 85 )</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note: The data are the CBCL-MIT benchmark data set of frontal faces (see text).

The first data set we used for training and testing is the well-known CBCL data set for frontal faces composed of thousands of small images of positive and negative examples of size (available online at http://cbcl.mit.edu/software-datasets/FaceData2.html). The face images obtained from this benchmark are clean and nicely registered.

The second data set we consider is made of low-quality images acquired by a monitoring system installed in our department (the data set is available on request). The data are very different from the previous set since they have been obtained from video frames (therefore, they are noisy and often blurred by motion), faces have not been registered, and gray values have not been normalized. The RBF kernel may take into account slight data misalignment due to the intraclass variability, but in this case, model selection is more crucial, and the choice of an appropriate parameter for the kernel is advisable.

The experiments performed on these two sets follow the structure discussed in the previous section. Starting from the original set of data, in both cases we randomly extracted 2000 data that we use for most of our experiments: for a fixed training set size, we generate 50 resamplings of training and test data. Then we vary the training set size from 600 (300 + 300) to 800 (400 + 400) training examples. The results obtained are reported in Tables 4 and 5. The tables show a comparison between the \( \nu \) method and SVM as the size of the training set grows. The results obtained are slightly different: while on the CBCL data set the \( \nu \) method performance is clearly above the SVM classifier, in the second set of data, the performance of the \( \nu \) method increases as the training set size grows.

At the end of this evaluation process, we retrained the \( \nu \)-method on the full set of 2000 data and again tuned the parameters with KCV, obtaining \( \sigma = 200 \) and \( t = 58 \). Then we used this classifier to test a batch of newly acquired data (the size of this new test set is of 6000 images), obtaining a classification error of 3.67%. These results confirm the generalization ability of the algorithm. For completeness, we report that the SVM classifier trained and tuned on the entire data set (\( \sigma = 600 \) and \( C = 1 \)) leads to an error rate of 3.92%.
Table 5: Average and Standard Deviation of the Classification Error of SVM and \( \nu \)-Method Trained on Training Sets of Increasing Size.

<table>
<thead>
<tr>
<th>Classifier</th>
<th>Number Trained Plus Number Tested</th>
<th>600 + 1400</th>
<th>700 + 1300</th>
<th>800 + 1200</th>
</tr>
</thead>
<tbody>
<tr>
<td>RBF-SVM</td>
<td>( \sigma = 570 ) C = 2 ( \sigma = 550 ) C = 1 ( \sigma = 550 ) C = 1</td>
<td>3.99 ± 1.21</td>
<td>3.90 ± 0.92</td>
<td>3.8 ± 0.58</td>
</tr>
<tr>
<td>( \nu )-method</td>
<td>( \sigma = 250 ) t = 67 ( \sigma = 180 ) t = 39 ( \sigma = 200 ) t = 57</td>
<td>4.36 ± 0.53</td>
<td>4.19 ± 0.50</td>
<td>3.69 ± 0.54</td>
</tr>
</tbody>
</table>

Note: The data have been acquired by a monitoring system developed in our laboratory (see text).

8 Conclusion

In this letter, we present and discuss several spectral algorithms for supervised learning. Starting from the standard regularized least squares, we show that a number of methods from inverse problems theory lead to consistent learning algorithms. We provide a unifying theoretical analysis based on the concept of filter function showing that these algorithms, which differ from the computational viewpoint, are all consistent kernel methods. The iterative methods, like the \( \nu \) method and the iterative Landweber, and projections methods, like spectral cut-off or PCA, give rise to regularized learning algorithms in which the regularization parameter is the number of iterations or the number of dimensions in the projection, respectively.

We report an extensive experimental analysis on a number of data sets showing that all the proposed spectral algorithms are a good alternative, in terms of generalization performances and computational efficiency, to state-of-the-art algorithms for classification like SVM and AdaBoost. One of the main advantages of the methods we propose is their simplicity: each spectral algorithm is an easy-to-use linear method whose implementation is straightforward. Indeed, our experience suggests that this helps dealing with overfitting in a transparent way and makes the model selection step easier. In particular, the search for the best choice of the regularization parameter in iterative schemes is naturally embedded in the iteration procedure.

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References


Spectral Algorithms for Supervised Learning


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