A New Solution Path Algorithm in Support Vector Regression
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Abstract—In this paper, regularization path algorithms were proposed as a novel approach to the model selection problem by exploring the path of possibly all solutions with respect to some regularization hyperparameter in an efficient way. This approach was later extended to a support vector regression (SVR) model called ε-SVR. However, the method requires that the error parameter ε be set a priori. This is only possible if the desired accuracy of the approximation can be specified in advance. In this paper, we analyze the solution space for ε-SVR and propose a new solution path algorithm, called ε-path algorithm, which traces the solution path with respect to the hyperparameter ε rather than λ. Although both two solution path algorithms possess the desirable piecewise linearity property, our ε-path algorithm overcomes some limitations of the original λ-path algorithm and has more advantages. It is thus more appealing for practical use.

Index Terms—Model selection, solution path, support vector regression (SVR).

I. INTRODUCTION

In a typical regression problem, we are given a training set of independent and identically distributed (i.i.d.) examples in the form of \( n \) ordered pairs, \( \{(x_i, y_i)\}_{i=1}^{n} \subseteq \mathbb{R}^d \times \mathbb{R} \), where \( x_i \) and \( y_i \) denote the input and the output, respectively, of the \( i \)th training example. Linear regression is the simplest method to solve the regression problem where the regression function is a linear function of the input. As a nonlinear extension, support vector regression (SVR) is a kernel method that extends linear regression to nonlinear regression by exploiting the kernel trick [1], [2]. Essentially, each input \( x_i \) is mapped implicitly via a nonlinear feature map \( \phi(\cdot) \) to some kernel-induced feature space \( \mathcal{F} \) where linear regression is performed. Specifically, SVR learns the following regression function by estimating \( w \in \mathcal{F} \) and \( w_0 \in \mathbb{R} \) from the training data

\[
    f(x) = \langle w, \phi(x) \rangle + w_0 \tag{1}
\]

where \( \langle \cdot, \cdot \rangle \) denotes the inner product in \( \mathcal{F} \). The problem is solved by minimizing some empirical risk measure that is regularized a priori to control the model capacity.

One commonly used SVR model is called ε-SVR. In the ε-SVR model, the ε-insensitive loss function

\[
    |y - f(x)|_{\varepsilon} = \max \{0, |y - f(x)| - \varepsilon\}
\]

is used to define an empirical risk functional, which exhibits the same sparseness property as that for support vector classifiers (SVC) using the hinge loss function via the so-called support vectors (SV). If a data point \( x \) lies inside the insensitive zone called the ε-tube, i.e., \( |y - f(x)| \leq \varepsilon \), then it will not incur any loss. However, the error parameter \( \varepsilon \geq 0 \) has to be specified a priori by the user. The primal optimization problem for ε-SVR can be stated as follows:

\[
    \min_{w, \xi^+} \frac{1}{2} |w|^2 + \sum_{i=1}^{n} (\xi_i^+ + \xi_i^-) \tag{2}
\]

subject to

\[
    \begin{align*}
    y_i - (\langle w, \phi(x_i) \rangle + w_0) & \leq \varepsilon + \xi_i^+ \tag{3} \\
    (\langle w, \phi(x_i) \rangle + w_0) - y_i & \leq \varepsilon + \xi_i^- \tag{4} \\
    \xi_i^+ & \geq 0, \tag{5}
    \end{align*}
\]

Here and below, \( i = 1, \ldots, n \) and \( (\cdot)^+ \) denote both the variables with and without asterisks. The regularization hyperparameter \( \lambda > 0 \) plays a role in capacity control by maintaining a proper balance between empirical loss and model complexity. Like \( \varepsilon \), \( \lambda \) also has to be chosen in advance by the user. The two hyperparameters \( \lambda \) and \( \varepsilon \) play different roles in ε-SVR. Fig. 1 shows four different combinations of the hyperparameter values.

In practice, users often use some default values for \( \lambda \) and \( \varepsilon \) even though they are by no means optimal choices. A better approach is to specify some candidate hyperparameter values and then apply cross validation to select the best choices among the candidates. However, extensive exploration for the optimal hyperparameter values is seldom pursued because this requires training the model many times under different hyperparameter settings. To overcome the difficulty of selecting \( \varepsilon \), Scholkopf et al. [3] proposed the \( \nu \)-SVR model, which automatically adjusts the width of the tube so that at most a fraction \( \nu \) of the data points lie outside the tube. Although \( \nu \)-SVR can help alleviate much of the effort involved in choosing the \( \varepsilon \) value, it is still very time consuming to find an appropriate combination of values for \( \lambda \) and \( \varepsilon \) to fit the data.

More recently, a novel approach has emerged to address the model selection problem by exploring the entire regularization path1 for solutions without having to retrain the model multiple times [4]–[8]. Research efforts in this direction have resulted in several regularization path algorithms. For SVR, Gunter and Zhu [9] devised an algorithm that computes the entire solution path with respect to \( \lambda \) for some fixed \( \varepsilon \) value. The algorithm starts from \( \lambda = \infty \), with the initial solution obtained by solving a linear programming problem. As \( \lambda \) decreases, the algorithm

1For notation simplicity, we refer to the regularization path as \( \lambda \)-path.
computes the solution for every value of \( \lambda \). However, we observe that sometimes the regularization path cannot be traced successfully during the execution of the algorithm. When there exists only one or even no point at the boundaries, the tube has to move and rotate until two valid points enter the tube. There exist many possible combinations for two points to enter the elbows simultaneously. Hence, the algorithm needs to enumerate all the combinations and then pick a valid one from them. However, because the search strategy to enumerate all possible combinations is not systematic, this difficulty will pose a problem to the algorithm. As a consequence, it is not easy to realize the \( \lambda \)-path algorithm in practice.

In this paper, we also address the solution path problem for \( \epsilon \)-SVM. Our main contributions can be summarized by the following two key findings.

- We establish that the dual variables of the dual optimization problem corresponding to (2) are piecewise linear with respect to the 2-D hyperparameter vector \((\lambda, \epsilon)\).
- We propose an efficient solution path algorithm, called \( \epsilon \)-path algorithm, that traces the solution path with respect to \( \epsilon \) for the optimization problem in (2) when \( \lambda \) is fixed.

Our \( \epsilon \)-path algorithm possesses competitive advantages over the \( \lambda \)-path algorithm proposed by [9]. Not only can the \( \epsilon \)-path algorithm always proceed without difficulty, but it also has a very simple initialization step and is efficient in finding a good regression function that can generalize well. More specifically, the \( \epsilon \)-path algorithm initializes the tube width to infinity, implying that it starts with no SVs. The algorithm then reduces the tube width so that the number of SVs increases gradually as the algorithm proceeds. As a result, a good regression function with the desirable sparseness property can be obtained only after a small number of iterations in decreasing \( \epsilon \).

The rest of this paper is organized as follows. Section II briefly reviews some existing solution path algorithms and Section III reviews the \( \epsilon \)-SVR formulation and introduces some basic terminology used later in this paper. In Section IV, we present a new approach for devising solution path algorithms and investigate the use of this approach for both the \( \lambda \)-path and the \( \epsilon \)-path. Some further discussions are given in Section V and experimental results are presented in Section VI. Finally, Section VII concludes this paper.

**II. SOLUTION PATH ALGORITHMS**

The basic idea underlying solution path algorithms comes from continuation methods [10], which compute the current solution based on an already obtained one. Specifically, we can interpret a solution path algorithm as follows: given a hyperparameter value \( \mu \) and its corresponding solution \( \hat{f}_\mu \), a solution path algorithm seeks to update the solution from \( \hat{f}_\mu \) to \( \hat{f}_{\mu+s} \) in an efficient way as \( \mu \) changes to \( \mu+s \) by a small value \( s \). The updating formula is often expressed as \( \hat{f}_{\mu+s} = \hat{f}_\mu + u(\mu, s) \). If the solution changes nonlinearly with \( s \), gradient descent or the Newton–Raphson method [11] can be used to estimate \( u(\mu, s) \). On the other hand, if the solution changes linearly with \( s \), \( u(\mu, s) \) can be expressed as \( s \cdot v(\mu) \) where \( v(\mu) \) does not depend on \( s \).

This approach makes it possible to trace the (entire) solution path as a function of the hyperparameter without having to train the model multiple times. Cross validation may then be used to estimate the hyperparameter value that gives the lowest generalization error. Because this approach has much lower computational demand without the need for training the model multiple times, we can afford to estimate the generalization errors for a much larger set of hyperparameter values in searching for the optimal choice. Previous works mainly focus on exploring the solution path with respect to the regularization hyperparameter, and hence, the resulting solution path is also called regularization path. Efron et al. [6] proposed an algorithm called the least angle regression (LARS) algorithm. It can be used to trace the regularization path for linear least square regression regularized with the \( L_1 \)-norm. An important finding is that the path of the solutions is piecewise linear, and hence, it is efficient to explore the entire path by monitoring the breakpoints between the linear segments only. Hastie et al. [7] proposed an algorithm to compute the regularization path for the standard \( L_2 \)-norm SVC and Zhu et al. [5] proposed one for the \( L_1 \)-norm SVC. They are again based on the property that the paths are piecewise linear with respect to the regularization hyperparameter. Rosset et al. [12] showed that boosting approximately follows the regularization path with an appropriate loss and an \( L_1 \) penalty. More generally, Rosset and Zhu [4] showed that any model with an \( L_1 \) regularization term and a quadratic, piecewise quadratic, piecewise linear, or linear loss function has a piecewise linear solution path, and hence, the entire path can be computed efficiently. For general loss functions and regularizers, the regularization paths are typically not (piecewise) linear. The predictor–corrector algorithm [10] is one of the fundamental strategies for implementing numerical continuation methods and can be used for

\[ \text{As the hyperparameter value } \mu \text{ changes, the solution estimator } \hat{f} \text{ will change accordingly. Thus, the estimator can be considered as a function of } \mu. \text{ We use } \hat{f}_\mu \text{ to indicate the dependence on } \mu. \]

In this paper, we investigate the solution space of \( \varepsilon \)-SVR based on a novel view. In particular, we consider two types of optimality conditions, namely, equality conditions and inequality conditions, which play different roles in exploring the solution space.

- The equalities establish conditions that must be satisfied throughout the entire solution path and thus determine the direction of movement between breakpoints.
- The inequalities determine which data points are involved in the equality constraints and thus determine the breakpoints.

Based on this approach, it is straightforward to investigate the path-following algorithm for either \( \lambda \) or \( \varepsilon \). An advantage of this approach is that it is very general and can be applied to explore the solution paths for other hyperparameters as well.

### III. SVR Problem Formulation

Applying the method of Lagrange multipliers to the primal optimization problem for \( \varepsilon \)-SVR in (2)–(5), we arrive at the following dual optimization problem:

\[
\max_{\alpha^{(s)}} - \frac{1}{2\lambda} \sum_{i,j=1}^{n}(\alpha_i - \alpha_i^*)^T (\alpha_j - \alpha_j^*) K(x_i, x_j) - \sum_{i=1}^{n} (\alpha_i + \alpha_i^*) y_i \tag{6}
\]

subject to

\[
0 \leq \alpha_i^{(s)} \leq 1, \quad \sum_{i=1}^{n} (\alpha_i - \alpha_i^*) = 0 \tag{7}
\]

where \( K(x_i, x_j) = \langle \phi(x_i), \phi(x_j) \rangle \) is the kernel function specified by the user and \( \alpha_i \) and \( \alpha_i^* \) are the Lagrange multipliers for constraints (3) and (4), respectively.

We also have

\[
w = \frac{1}{\lambda} \sum_{i=1}^{n} (\alpha_i - \alpha_i^*) \phi(x_i). \tag{9}
\]

By substituting (9) into (1), the regression function can be rewritten as

\[
f(x) = \langle w, \phi(x) \rangle + u_0 = \frac{1}{\lambda} \sum_{i=1}^{n} (\alpha_i - \alpha_i^*) K(x_i, x) + u_0. \tag{10}
\]

From the Karush–Kuhn–Tucker (KKT) optimality conditions, we can derive the following relationships:

\[
y_i - f(x_i) > \varepsilon \Rightarrow \alpha_i = 1, \quad \alpha_i^* = 0 \tag{11}
\]

\[
y_i - f(x_i) = \varepsilon \Rightarrow \alpha_i \in [0, 1], \quad \alpha_i^* = 0 \tag{12}
\]

\[
y_i - f(x_i) \in (-\varepsilon, \varepsilon) \Rightarrow \alpha_i = 0, \quad \alpha_i^* = 0 \tag{13}
\]

\[
y_i - f(x_i) = -\varepsilon \Rightarrow \alpha_i = 0, \quad \alpha_i^* \in [0, 1] \tag{14}
\]

\[
y_i - f(x_i) < -\varepsilon \Rightarrow \alpha_i = 0, \quad \alpha_i^* = 1 \tag{15}
\]

As a consequence, \( f(x) \) can be expressed as an expansion in terms of only a subset of data points for which either \( \alpha_i \) or \( \alpha_i^* \) is nonzero. These points are referred to as SVs that, like those for SVC, contribute to the sparseness property of \( f(x) \) with representational and computational advantages.

Following the convention in [9], we partition the set of all data points into the following five subsets as illustrated in Fig. 2:

- \( R = \{ i : y_i - f(x_i) > \varepsilon, \alpha_i = 1, \alpha_i^* = 0 \} \) (right of the elbows);
- \( \mathcal{E}_R = \{ i : y_i - f(x_i) = \varepsilon, \alpha_i \in [0, 1], \alpha_i^* = 0 \} \) (right elbow);
- \( C = \{ i : |y_i - f(x_i)| \leq \varepsilon, \alpha_i = 0, \alpha_i^* = 0 \} \) (center);
- \( \mathcal{E}_L = \{ i : y_i - f(x_i) = -\varepsilon, \alpha_i = 0, \alpha_i^* \in [0, 1] \} \) (left elbow);
- \( L = \{ i : y_i - f(x_i) \leq -\varepsilon, \alpha_i = 0, \alpha_i^* = 1 \} \) (left of the elbows).

As we change the value of \( \lambda \) or \( \varepsilon \), the tube will move, rotate, shrink, expand, or remain unchanged. Some events may occur during this process. An event is said to occur when a point enters or leaves an elbow, causing some point sets to change as a result. We categorize the different events as follows.

1) A point enters an elbow:
- From \( \mathcal{C} \) to \( \mathcal{E}_R \) with \( \alpha_i = 0 \);
- From \( \mathcal{C} \) to \( \mathcal{E}_L \) with \( \alpha_i^* = 0 \);
- From \( \mathcal{R} \) to \( \mathcal{E}_R \) with \( \alpha_i = 1 \);
- From \( \mathcal{L} \) to \( \mathcal{E}_L \) with \( \alpha_i^* = 1 \).

2) A point leaves an elbow:
- From \( \mathcal{E}_R \) to \( \mathcal{R} \) with \( \alpha_i = 1 \);
- From \( \mathcal{E}_R \) to \( \mathcal{C} \) with \( \alpha_i^* = 0 \);
- From \( \mathcal{E}_L \) to \( \mathcal{C} \) with \( \alpha_i^* = 0 \);
- From \( \mathcal{E}_L \) to \( \mathcal{L} \) with \( \alpha_i^* = 1 \).

For those points lying inside or outside the tube, i.e., in \( \mathcal{R} \cup \mathcal{L} \), their \( \alpha_i^{(s)} \) values remain fixed until an event occurs. Hence, it suffices to focus on the points at the elbows, i.e., in \( \mathcal{E}_R \cup \mathcal{E}_L \).

### IV. SVR Solution Paths

#### A. Optimality Conditions

The SVR dual optimization problem in (6)–(8) is a quadratic programming (QP) problem. When the kernel matrix \( K = (K(x_i, x_j)) \) is positive semidefinite, the problem has a unique optimal solution. For the convenience of subsequent

3The notion of sparsity here is somewhat different from that commonly used in statistics in the context of variable selection. As a kernel method, the sparseness property of \( f(x) \) refers to a regression function that is represented as a linear combination of a small number of kernel function terms.
derivation, we define \( \alpha_0 = \lambda w_0 \) and rewrite the regression function in (10) as

\[
f(x) = \frac{1}{\lambda} \left[ \sum_{i=1}^{n} (\alpha_i - \alpha_i^* \lambda) K(x_i, x) + \alpha_0 \right].
\]  

(16)

In the dual optimization problem, convex optimization is performed on a feasible set defined by the constraints (7) and (8). However, when the KKT conditions (11)–(15) play a role in defining the optimality conditions, \((\alpha^*, \alpha_0)\) is the optimal solution if and only if it satisfies all these conditions. Thus, (7) and (8) and (11)–(15) comprise the optimality conditions for SVR.

The optimality conditions can be distinguished into the equality conditions (8), (12), and (14) and the inequality conditions (7), (11), (13), and (15). The equality conditions (12) and (14), which hold only for the points at the elbows (i.e., in \(\mathcal{E}_R \cup \mathcal{E}_L\)), lead to the following linear system:

\[
\sum_{i \in \mathcal{E}_R} \alpha_i K(x_i, x_j) - \sum_{i \in \mathcal{E}_L} \alpha_i^* K(x_i, x_j) + \alpha_0 = \lambda (y_j - \epsilon) - \sum_{i \in \mathcal{R}} \alpha_i K(x_i, x_j) + \sum_{i \in \mathcal{L}} \alpha_i^* K(x_i, x_j) \quad \forall j \in \mathcal{E}_R
\]  

(17)

\[
\sum_{i \in \mathcal{E}_R} \alpha_i K(x_i, x_k) - \sum_{i \in \mathcal{E}_L} \alpha_i^* K(x_i, x_k) + \alpha_0 = \lambda (y_k + \epsilon) - \sum_{i \in \mathcal{R}} \alpha_i K(x_i, x_k) + \sum_{i \in \mathcal{L}} \alpha_i^* K(x_i, x_k) \quad \forall k \in \mathcal{E}_L.
\]  

(18)

From another equality condition (8), we have

\[
\sum_{i \in \mathcal{E}_R} \alpha_i - \sum_{i \in \mathcal{E}_L} \alpha_i^* = -\sum_{i \in \mathcal{R}} \alpha_i + \sum_{i \in \mathcal{L}} \alpha_i^*.
\]  

(19)

Suppose \(\mathcal{E}_R\) contains \(p_1\) indices and \(\mathcal{E}_L\) contains \(p_2\) indices. The value \(p = p_1 + p_2 + 1\) indicates the total number of points at the elbows. The size of the linear system defined by (17)–(19) is thus equal to \(p = p_1 + p_2 + 1\). The system of \(p\) linear equations can be represented in matrix form as

\[
\begin{bmatrix}
\alpha_0 \\
\alpha_{\mathcal{E}_R} \\
-\alpha_{\mathcal{E}_L}^*
\end{bmatrix}
= \lambda \mathbf{y}_\mathcal{E} + \lambda \epsilon \mathbf{1}_\mathcal{E} - \mathbf{K}^b
\begin{bmatrix}
0 \\
\alpha_{\mathcal{E}_R} \\
-\alpha_{\mathcal{E}_L}^*
\end{bmatrix}
\]  

(20)

where the undefined notations are introduced in the Appendix.

Suppose we have an SVR solution for certain hyperparameters \((\lambda, \epsilon)\). According to the regression function of this solution, data points are partitioned into five subsets as defined above. As the hyperparameter \(\lambda\) or \(\epsilon\) changes its value, the solution will change accordingly in order to satisfy (20). We consider the period between two consecutive events when \(\alpha_{\mathcal{E}_R}\) and \(\alpha_{\mathcal{E}_L}^*\) remain unchanged at 0 or 1. During this period, only those points at the elbows \((i \in \mathcal{E}_R \cup \mathcal{E}_L)\) can change their values. From the equality condition (8), we know that if one point at an elbow changes its value, then at least one other elbow point must also modify its value correspondingly. An immediate consequence of this is that there should be at least two points at the elbows, i.e., \(p_1 + p_2 \geq 2\).

When a new hyperparameter value is specified, the corresponding solution can be computed as

\[
\begin{bmatrix}
\alpha_0 \\
\alpha_{\mathcal{E}_R} \\
-\alpha_{\mathcal{E}_L}^*
\end{bmatrix}
= \lambda (\mathbf{K}^a)^{-1} (\mathbf{y}_\mathcal{E} + \epsilon \mathbf{1}_\mathcal{E}) - (\mathbf{K}^a)^{-1} \mathbf{K}^b
\begin{bmatrix}
0 \\
\alpha_{\mathcal{E}_R} \\
-\alpha_{\mathcal{E}_L}^*
\end{bmatrix}
\]  

(21)

It is easy to see that \(\begin{bmatrix}\alpha_0 \\
\alpha_{\mathcal{E}_R} \\
-\alpha_{\mathcal{E}_L}^*\end{bmatrix}\) is linear in either \(\lambda\) or \(\epsilon\). If the inverse of \(\mathbf{K}^a\) does not exist, the update of the solution will no longer be unique. For the sake of illustration, we consider an example where the points \(\{x_i\}\) are from a 1-D space. When a linear kernel is applied and there exist three points at the elbows, the kernel matrix \(\mathbf{K}^a\) is not of full rank. As a result, the solution path algorithm can have multiple updating possibilities with each one corresponding to choosing any two of the three points to compute a new solution. In this paper, we exclude this case by considering the positive–definite kernels only. When the kernels are positive semidefinite, the optimal solution may not be unique. We modify \(\mathbf{K}^a\) by adding a ridge term to ensure that \((\mathbf{K}^a)^{-1}\) always exists and an approximate path of solutions is obtained.

The \(p\) linear equations above have been used to derive the updating formula for a new solution. However, the derivation is based on the assumption that no event occurs during the period even though the hyperparameter value has changed. Hence, given a certain hyperparameter value and the solution obtained for that value, the solutions for new hyperparameter values within a local neighborhood in which no event occurs can be computed directly via (21). On the other hand, the inequality conditions serve to indicate when the assumption no longer holds. When a new solution along the path violates the inequality conditions, the point sets have to be updated to determine the next breakpoint. Here, we consider the algorithm to trace the \(\lambda\)-path from \(\lambda^1\) in the decreasing direction. Tracing the \(\lambda\)-path in the increasing direction and tracing the \(\epsilon\)-path are both very similar to this.

We simplify (21) to

\[
\begin{bmatrix}
\alpha_0 \\
\alpha_{\mathcal{E}_R} \\
-\alpha_{\mathcal{E}_L}^*
\end{bmatrix}
= \lambda \mathbf{v}^a - \mathbf{v}^b
\]  

(22)

where

\[
\mathbf{v}^a = (v_{0i}^a, v_i^a (\forall i \in \mathcal{E}_R), v_i^a (\forall i \in \mathcal{E}_L))^T
\]  

(23)

\[
\mathbf{v}^b = (v_{0i}^b, v_i^b (\forall i \in \mathcal{E}_R), v_i^b (\forall i \in \mathcal{E}_L))^T
\]  

(24)

are independent of \(\lambda\). The algorithm monitors the occurrence of any of the following possible events.

\footnote{If \(p_1 + p_2 < 2\), we face a problem in the initialization setup. The algorithms for the \(\lambda\)-path and the \(\epsilon\)-path use different methods to handle this case. We will discuss them in Sections IV-B and IV-C, respectively.}
The $\alpha_i$ or $\alpha^*_i$ value of a point $i \in E_R \cup E'_R$ reaches 0 or 1. The $\lambda_i$ value of the point can be calculated by $\lambda_i = (\alpha_i + \nu^*_i)/\nu_i \ (\forall i \in \mathcal{E}_R)$ or $\lambda_i = (-\alpha^*_i + \nu^*_i)/\nu^*_i \ (\forall i \in \mathcal{E}_L)$.

A point $i \in \mathcal{R} \cup \mathcal{C} \cup \mathcal{L}$ hits an elbow, i.e., $y_i - f(x_i) = \epsilon$. The $\lambda_i$ value of the point can be calculated directly by plugging (22) into (16).

Because the solution is linear in either $\lambda$ or $\epsilon$, the $\lambda_i$ values for the possible events above can be computed exactly. Through monitoring these possible breakpoint values, we can find the next breakpoint $X^{t+1} = \arg\max_{\lambda_i < X^t} \{ \lambda_i \}$, at which the next event occurs. The algorithm then updates the point sets before decreasing the hyperparameter value further. This updating step is essential, or else the solutions computed for the hyperparameter values beyond the breakpoint without updating the point sets will lead to violation of the inequality conditions. We thus have an algorithm for tracing the SVR solutions along the $\lambda$-path or the $\epsilon$-path. However, because the $\lambda$-path and $\epsilon$-path have different properties, we will discuss them separately in Section IV-B and IV-C.

B. $\lambda$-Path

The $\lambda$-path algorithm discussed above is essentially the same as the regularization path algorithm proposed by [9]. However, the algorithm derived based on our approach for computing the next breakpoint via (22) is simpler and easier to understand. The $\lambda$-path algorithm explores the correspondence between every $\lambda$ value and the corresponding solution $(\alpha^{(s)}(\lambda), u_{0}(\lambda))$ while $\epsilon$ is fixed. The path may start from the solution of an $\epsilon$-SVR model for any initial value of $\lambda$, because the values of $\alpha^{(s)}$ fully determine the sets $\mathcal{R}_c, \mathcal{E}_c, \mathcal{C}, \mathcal{L}_c,$ and $\mathcal{L}$. However, this requires solving a QP problem, which is computationally demanding if it is solved directly. The problem becomes simpler if we set $\lambda = \infty$ initially. Doing so will make the first term of the objective function (6) vanish, leaving only the last two terms. Thus, the QP problem degenerates to a linear programming problem, which is easier to solve. Similarly, the first term of (10) vanishes so that the regression function becomes $f(x) = u_0$, which corresponds to the case shown in Fig. 1(b). The initial values of $\alpha^{(s)}_i$, denoted as $\alpha^{(s)}_i$, are either 0 or 1 if all the values of $y_i$ are distinct. The $\epsilon$-tube is bounded by the sets $\mathcal{R}_c, \mathcal{E}_c, \mathcal{C}, \mathcal{L}_c,$ and $\mathcal{L}$. The tube can move around by changing $\lambda$ and $u_0$, while no point is allowed to pass through any elbow. Hence, the following constraints are satisfied:

$$y_j - \frac{1}{\lambda} \sum_{i} (\alpha^0_i - \alpha^{(s)}_i) K(x_i, x_j) - u_0 > \epsilon, \quad \text{for } j \in \mathcal{R}$$

$$\left| y_j - \frac{1}{\lambda} \sum_{i} (\alpha^0_i - \alpha^{(s)}_i) K(x_i, x_j) - u_0 \right| < \epsilon, \quad \text{for } j \in \mathcal{C}$$

$$y_j - \frac{1}{\lambda} \sum_{i} (\alpha^0_i - \alpha^{(s)}_i) K(x_i, x_j) - u_0 < -\epsilon, \quad \text{for } j \in \mathcal{L}.$$

(25)

To explore the solution path with respect to $\lambda$, there should be at least two points at the elbows, i.e., $p_1 + p_2 \geq 2$. Gunter and Zhu [9] proposed a strategy for finding a feasible $(\lambda, u_0)$ combination so that two points enter the elbows simultaneously. The first moves the tube until a point enters one elbow through changing $u_0$. Then, it decreases $\lambda$ until another point also enters an elbow. However, we find that this strategy is very difficult to implement in practice. The reason is that, given the sets $\mathcal{R}, \mathcal{C}$, and $\mathcal{L}$, there exist many possible combinations for two points to enter the elbows simultaneously. However, most of these combinations are invalid because the $\lambda$-path algorithm cannot proceed further with the corresponding point sets. Fig. 3 depicts some possible valid and invalid cases. For the upper invalid case, we assume that a cross point $i \in \mathcal{R}$ enters one elbow with $\alpha_i = 1$ and a circle point $j \in \mathcal{C}$ enters another elbow simultaneously with $\alpha^*_j = 0$. To continue the $\lambda$-path, these two points should pass through the elbows during which $\alpha_j$ should decrease to 0 but $\alpha^*_j$ should increase to 1. However, doing so will lead to violation of the constraints in (7), showing that the two points cannot stay at the elbows to continue the $\lambda$-path. Thus, the algorithm has to make the tube move and rotate without changing the point sets $\mathcal{R}, \mathcal{C}$, and $\mathcal{L}$ until another two points enter the elbows. Because the movement of the tube to search for two valid points is not systematic, it is difficult to implement a program to explore all possible combinations for two points to enter the elbows simultaneously. It is also difficult to estimate the number of iterations required to find two such valid points. In fact, whenever the elbows contain fewer than two points while the $\lambda$-path is being traced, the algorithm has to perform such a random search, making this approach unappealing in practice.

C. $\epsilon$-Path

In this section, we illustrate some interesting properties of our proposed $\epsilon$-path algorithm. If we set $\epsilon = \infty$, it is trivial to solve the optimization problem in (6)–(8). The solution is simply $\alpha^{(s)} = 0$ for all $i$, meaning that all the points are inside the tube (i.e., in $\mathcal{C}$) and, from (10), $f(x) = u_0$. This corresponds to the case shown in Fig. 1(c). Because $y_i - f(x_i) = \epsilon$ for all $i$, we set $\epsilon > (y_{\max} - y_{\min})/2$ and $u_0 = (y_{\max} + y_{\min})/2$ initially. Compared with the $\lambda$-path algorithm that has to solve a linear programming problem, the initialization problem for the $\epsilon$-path algorithm is much easier to solve.

We assume that $\lambda$ is prespecified by the user and it remains fixed during the execution of the $\epsilon$-path algorithm. Along the $\epsilon$-path, we always have $|E_R| > 0$ and $|E_L| > 0$ and we apply (21) to update the solution. If there is an elbow containing no points, we reduce $\epsilon$ until each elbow contains at least one point. Let $i_+ = \arg\max_{i \in \mathcal{C}} (y_i - f(x_i))$ and $i_- = \arg\min_{i \in \mathcal{C}} (y_i - f(x_i))$. Then, $E_R = \{i_+\}$, $E_L = \{i_-\}$, and $\epsilon = (y_{i_+} - y_{i_-})/2$.
This procedure is very efficient; it only involves shrinking the tube to reduce its width without rotating it. It is deterministic to find two points at the elbows simultaneously and is thus much easier to implement than the random search in the $\lambda$-path algorithm when $p_1 + p_2 < 2$.

The process along the $c$-path can be understood geometrically in the linear space. If $d = 1$ and each elbow contains one point, then decreasing $\epsilon$ will rotate the tube with the two points at the two elbows as the centers of rotation. Fig. 1(d) shows one such example. The resulting rotation causes the width of the tube to decrease while the two elbow points remain at the elbows. Considering further the above example, the $c$-path algorithm cannot proceed if a new point enters an elbow, i.e., $|e^L_R| + |e^L_L| > 2$, $|e^L_R| \geq 1$, and $|e^L_L| \geq 1$. In the $d = 1$ linear space, the width of the tube will be fixed by these elbow points, and hence, the tube cannot rotate nor shrink. As a result, $\epsilon$ cannot decrease. This problem always occurs in the linear space. If the dimensionality of the linear space is $d$, the rank of $K_c$ is at most $d$, no matter how many points are involved. Thus, the inverse of $K^d$ does not exist when the elbows contain more than $d$ points. This problem can also be overcome by using a positive–definite kernel or adding a ridge term. For example, if the Gaussian kernel is used, we can always execute the $c$-path algorithm no matter how many points are at the elbows.

D. Complexity Analysis

The optimization problems for both $c$-SVR and $\nu$-SVR can be formulated as QP problems, with $O(n^2)$ time complexity and $O(n^3)$ space complexity. Many attempts have been made to reduce the computational complexity of support vector machine (SVM) algorithms. For example, one popular approach is to obtain low-rank approximation of the kernel matrix by using the Nyström method [13], greedy approximation [14], or other methods. Another approach is to use chunking and decomposition methods [1], [15]–[18], which perform optimization on only a small subset of all the training data at a time. The sequential minimal optimization (SMO) algorithm [17], [19] has been considered as the state-of-the-art SVM implementation, with training time complexity empirically observed to be between $O(n)$ and $O(n^2/3)$. In practice, users often prespecify a small set of candidate hyperparameter values and then perform cross validation to estimate the generalization error of the model trained for each choice of the hyperparameter value. The choice that gives the lowest generalization error is considered the best choice. A disadvantage of this approach is that the search for the “optimal” choice is limited to be within a relatively small set of choices that have to be prespecified in advance by the user. This approach is computationally prohibitive if we want to perform an extensive search for a good hyperparameter value. To analyze the computational complexity of the solution path algorithm, we rewrite the regression function $f(\mathbf{x})$ as

$$b_{RL}(\mathbf{x}) = \sum_{i \in R} \alpha^c_i K(\mathbf{x}_i, \mathbf{x}) - \sum_{i \in L} \alpha^c_i K(\mathbf{x}_i, \mathbf{x}).$$

While the solution path is being explored, all the $f(\mathbf{x}_i)$ values are repeatedly used for estimating the next breakpoint. Because $\alpha_R$ and $\alpha_L$ remain unchanged between two consecutive events, it will lead to significant computational saving if we cache the values of $b_{RL}(\mathbf{x}_i)$ $\forall \mathbf{x}_i$. Let $|R| + |L| = q$. Computing $b_{RL}(\mathbf{x}_i)$ $\forall \mathbf{x}_i$ during the preprocessing step has $O(nq)$ time complexity and $O(n)$ space complexity. When the next event occurs, the point sets are changed and the algorithm updates $b_{RL}(\mathbf{x}_i) \forall \mathbf{x}_i$ with $O(n)$ complexity. To calculate $\mathbf{v}^c$ and $\mathbf{v}^L$ in the updating formula (22), some matrix operations are performed with $O(p^3)$ time complexity. Because there is only one point difference in the elbows between two consecutive events, the Sherman–Morrison–Woodbury formula for block matrix inversion can be applied, and hence, the computational cost can be reduced to $O(p^2)$. To find the first inequality condition violated by the path, it requires scanning through all inequality conditions to estimate the next breakpoint. Because $b_{RL}(\mathbf{x})$ has been computed in advance, it requires $O(np^2)$ time complexity to find the first point that will hit or leave the elbows. Therefore, the overall computational cost between two breakpoints is $O(p^2 + np + n)$. The total number of breakpoints along the solution path depends on the range of the hyperparameter values we want to explore. Starting from $\epsilon = \infty$, most points move from inside the tube to outside as the width of the tube decreases. When nonlinear kernel functions are used, some points may reenter the tube after leaving it and pass through the elbows multiple times as $\epsilon$ decreases. Empirically, we found that the number of breakpoints is a small multiple of $n$ even when exploring a wide range of hyperparameter values such as $\lambda \in (0, \infty)$ or $\epsilon \in (0, \infty)$.

However, it is usually not necessary to explore the entire $c$-path in SVR. When $\epsilon$ is initialized to infinity in the beginning, all points are inside the tube, and hence, there is no SV. As $\epsilon$ decreases, the points pass through the elbows from inside the tube to outside and the number of SVs increases. This has a similar effect as increasing $n$ from 0 to 1 in $\nu$-SVR, but the number of SVs can be controlled exactly in our method. To obtain a desirable regression function with the sparseness property, we only need to run the algorithm for a small number of steps. Therefore, a regression function with the desired modeling ability can be obtained very efficiently. Note that the $\lambda$-path for SVC is explored in a reverse direction as compared with the $c$-path for SVR. In the SVC formulation, the SVs are those points inside the margin. To simplify the initialization step for the $\lambda$-path algorithm, $\lambda$ is initialized to be very large so that most points are inside the margin. At this moment, most of the points are SVs. As $\lambda$ decreases, both the width of the margin and the number of SVs decreases. Because a classifier that generalizes well typically has a sparse representation involving a small number of SVs, almost the entire solution path has to be explored until $\lambda$ becomes very small so that most points are excluded from the margin. Thus, the total number of moves is $O(n)$. Based on empirical findings, Hastie et al. [7] suggested that this number is some small multiple of $n$.

Because the solution path proceeds in a piecewise linear manner, any solution between two breakpoints can be computed efficiently via interpolation based on the two solutions obtained.
at the breakpoints. As a result, it suffices to store the solutions for the breakpoints only.

V. DISCUSSIONS

From (21), we have seen that the update is linear in either \( \lambda \) or \( \epsilon \). When the values of \( \lambda \) and \( \epsilon \) change simultaneously, the corresponding solution can still be obtained directly from (21). Even if we change the kernel parameter value, which will lead to changes in \( K_n \) and \( K^0 \), this simple update can still be used for computing the new solution. The updating formula remains valid between two consecutive events when the point sets do not change. If further tracing the path of solutions after an event occurred, the solution path will lead to a breakpoint at which an event occurs. The point sets have to be updated before the solution path is traced beyond this breakpoint, and a new formula is used to update the solution. However, the update with changes in \( (\lambda, \epsilon) \) simultaneously or in the kernel parameter is nonlinear, thus the next breakpoint cannot be calculated in advance like that in the \( \epsilon \)-path or the \( \lambda \)-path. One straightforward approach [20] to estimate the new breakpoint is to change the hyperparameter value by a small increment and update the new solution. It then checks whether the next event has occurred. If the next event has not occurred, we keep on changing the hyperparameter value further. Otherwise, the increment of the change is decreased to a smaller value. Following this procedure, the algorithm can iteratively approximate the next breakpoint and continue to trace a nonlinear solution path. As we see, it is more difficult to calculate the next breakpoint during the nonlinear path exploration. If we are given a certain direction \( (\lambda, \epsilon) \) in which the path is traced, it will be more efficient to use \( \epsilon \)-path and \( \lambda \)-path interchangeably.

After exploring the path of solutions, it is necessary to estimate the generalization errors of these solutions and then pick an optimal one from the solutions. Efron [21] gave a comprehensive discussion on the estimation of generalization errors and Zou et al. [22] studied the degrees of freedom of LASSO in the framework of Stein’s unbiased risk estimation (SURE). Based on these works, Gunter and Zhu [9] proposed an unbiased estimator for the degrees of freedom of the SVR model, i.e.,

\[
\tilde{d}_f = |E_R| + |E_C|, \tag{31}
\]

The number of points at the elbows is a simple unbiased estimator for the degrees of freedom of \( f(x) \). This estimator can be applied to derive AIC [23] and BIC [24] for model selection. However, because there is an assumption that the response \( y \) is generated according to a homoskedastic model, an additional variable, i.e., the variance, has to be estimated. The generalized cross-validation (GCV) criterion [25], which is an approximation of the leave-one-out cross-validation criterion, is another criterion for estimating the generalization error without requiring a known variance

\[
\text{GCV} = \frac{\sum_{i=1}^{n}(y_i - f(x_i))^2}{(1 - \tilde{d}_f/n)^2}. \tag{32}
\]

Gunter and Zhu [9] used this criterion to estimate the generalization errors of the SVR solutions. The degree of freedom provides an intuitive and informative measure of the complexity of a fitted model. When the elbows contain many points, the complexity of the regression function will become too high to generalize well. The number of SVs indicates the sparsity of the function, but it is by no means an indicator of the generalization performance. Our experimental results in Section VI also illustrate this phenomenon.

VI. EXPERIMENTAL RESULTS

The behaviors of the above algorithms can best be illustrated using video. We have prepared some videos to show several illustrative examples.\(^5\)

A. Toy Example: Noisy Sinc-Function

We generate a set of 100 data points \( \{(x_i, y_i)\} \) with \( x_i \) drawn uniformly from \([-3,3]\) and \( y_i = \sin(\pi x_i)/(\pi x_i) + e_i \), where \( e_i \) is a Gaussian noise term with zero mean and a variance of

\(^5\)The videos and the code for the \( \lambda \)-path and \( \epsilon \)-path algorithms are available at http://www.cse.ust.hk/wanggang/svrpath.htm
function, it is not a good indicator of the generalization performance. Because the elbow size remains almost unchanged as $\epsilon$ is less than a certain value when $\gamma = 0.5$, the generalization error performs well along the corresponding $\epsilon$-path.

Fig. 7 shows the effect of different values of $\lambda$ on the $\epsilon$-path algorithm. It shows that the regression function is not very sensitive to $\lambda$ if it is given a moderate value. When $\lambda = 0.01$ and 1, the solution paths show similar generalization error curves as $\epsilon$ decreases. Their generalization errors decrease dramatically in the beginning and good regression functions can be obtained. As the $\epsilon$-path further proceeds, the performance of the regression functions remains stable and the generalization errors change slightly. However, when $\lambda$ is quite large ($\lambda = 10$), it always tends to give a flat function leading to large error. It is interesting to notice that when $\lambda$ is very small ($\lambda = 0.001$), its beginning part of the generalization error curve is overlapped with those when $\lambda = 0.01$ or 1. The regression function can fit the data well after a few iterations of the algorithm. Nevertheless, the generalization error increases as the algorithm proceeds, thus overfitting occurs. Both $\epsilon$ and $\lambda$ control the model complexity in SVR. This property is different from that of many statistical models, because only setting the regularization hyper-parameter to be small does not necessarily lead to “poor” generalization ability. The generalization error curve for the $\epsilon$-path will first decrease and then increase for very small values of $\lambda$. To avoid the overfitting cases, the value of $\lambda$ should not be set very small. In this data set, we observe that $\lambda$ taking values from $[0.01, 1]$ is a good choice. Fig. 7(b) shows that $\epsilon$ decreases rapidly during the first few steps of the $\epsilon$-path algorithm. Afterwards, the rate of decrease in $\epsilon$ slows down significantly. As $\epsilon$ decreases, more and more points move towards the elbows. There is an inflexion point after which both $\epsilon$ and the regression function only change slightly. Executing more steps of the algorithm beyond this point leads to imperceptible changes to the regression function. We next examine the relationships between the generalization error and the number of steps in Fig. 7(c). Similar to Fig. 7(b), the generalization error decreases rapidly during the first few steps. The generalization error is minimized at around the position where the inflexion point occurs. At this time, only a small number of steps have been executed. Further executing the $\epsilon$-path algorithm cannot lead to continued improvement in the generalization performance. Instead, the resulting regression function becomes more redundant and may lead to overfitting. Moreover, it incurs unnecessarily high computational cost. Consequently, it is not necessary to explore all solutions along the $\epsilon$-path. The optimal solution preserving the sparseness property can be obtained very efficiently.

We also give some preliminary comparisons on the computational expense between our solution path algorithm and

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0.1. We randomly partition the data set into a training set of 50 points and a validation set of 50 points. The Gaussian radial basis function (RBF) kernel $K(x_i, x_j) = \exp(||x_i - x_j||^2/\gamma)$ is used. Because the input points are from a 1-D space, the regression function can be shown as a 2-D plot. Fig. 4 plots the SVR results at three breakpoints along the $\epsilon$-path. We run the $\epsilon$-path algorithm with three different $\gamma$ values, 0.05, 0.5, and 5, while setting $\lambda = 0.1$. The algorithm terminates when $\epsilon$ decreases to 0.03, at which most of the points become SVs. For each $\epsilon$-path, we compute the mean squared error (MSE) on the validation set to estimate the generalization error. Fig. 5 plots the training data, the sinc function, and the regression functions that minimize the MSE along the whole $\epsilon$-paths for different kernel values. We can see that the optimal regression function overfits the data when $\gamma = 0.05$ but underfits the data when $\gamma = 5$. On the other hand, it fits the data well when $\gamma = 0.5$.

Fig. 5 plots the number of SVs, the elbow size, and the estimated generalization error as a function of $\epsilon$ for different values of $\gamma$. As the $\epsilon$-path proceeds, the tube width always decreases and more points become SVs. The number of SVs increases accordingly regardless of what the $\gamma$ value is. When $\gamma = 0.05$, the tube and the regression function are very elastic. The elbow size generally increases as $\epsilon$ decreases. During this process, more and more points move into the elbows and then settle down there. The regression function is thus sensitive to having many points at the elbows. It leads to a large degree of freedom, which overfits the data. In other words, the model has low bias but high variance. When $\gamma = 5$, on the other hand, the elbow size remains small along the $\epsilon$-path. Because the function is not flexible enough, many points cannot stay at the elbows simultaneously. It induces a model with high bias but low variance, thus underfitting the data. Setting $\gamma = 0.5$ can fit the data satisfactorily as the bias and variance are balanced well. Therefore, the elbow size measures the complexity of the regression function. When its value is either too high or too low, it cannot generalize well. From Fig. 6(a), we notice that the number of SVs always increases. Although it determines the sparsity of the regression function, it is not a good indicator of the generalization performance. Because the elbow size remains almost unchanged as $\epsilon$ is less than a certain value when $\gamma = 0.5$, the generalization error performs well along the corresponding $\epsilon$-path.
Fig. 6. Results on (a) the number of SVs, (b) the elbow size, and (c) the generalization error along the SVR $\epsilon$-path for different $\gamma$ values. $\lambda$ is set to 0.1.

LIBSVM [18] to give readers some practical sense. Our algorithm is implemented in MATLAB. LIBSVM is written in C++ and we use its interface for MATLAB. The experiments are performed on a ThinkPad T61 notebook with Intel T7300 Core 2 Duo (2.0-GHz, 800-MHz FSB, 4-MB cache) processor and 2-GB memory. We generate the sinc-function data sets with the size of 100, 200, 400, 800. There are five data sets generated for each size in order to alleviate random sampling. For each data set, the $\epsilon$-path algorithm is executed first until 50% of the points become SVs. Thus, a sequence of breakpoints is computed. LIBSVM is then executed for all $\epsilon$ values at breakpoints and we record the total elapsed time (in seconds). From the results shown in Table I, we can see that the solution path algorithm has much computational advantage over the original model selection approach. The number of breakpoints increases almost linearly with the size of the training set. Comparing the number of breakpoints and the ratio in the table, we notice that the computational cost of exploring the entire solution path is similar to that of training LIBSVM once when the data size is 100. As the size of training data increases, the expense of the path exploration becomes higher than computing LIBSVM once.

B. Friedman’s Benchmark Functions

Friedman’s benchmark functions were introduced in [26] and have become a widely used benchmark for regression models. There are three nonlinear prediction problems in this benchmark. Friedman’s $F1$ function has ten independent variables $x = (x_1, \ldots, x_{10})^T$ that are uniformly distributed in $[0,1]$

$$F1: \ y = 10\sin(\pi x_1 x_2) + 20(x_3 - 0.5)^2 + 10x_4 + 5x_5 + e$$ (33)

where $e \sim N(0,1)$. This function depends on only five of the ten variables. The $F2$ and $F3$ functions both have four indepen-
dent variables \( \mathbf{x} = (x_1, \ldots, x_d)^T \) that are uniformly distributed in the following ranges: \( 0 \leq x_1 \leq 100, \quad 40\pi \leq x_2 \leq 500\pi, \quad 0 \leq x_3 \leq 1, \) and \( 1 \leq x_4 \leq 11. \) The functions are defined as

\[
F2: \quad y = \sqrt{x_1^2 + \left[ x_2 x_3 - (x_2 x_4)^{-1} \right]^2} + \epsilon, \quad e \sim \mathcal{N}(0, 125) \tag{34}
\]

\[
F3: \quad y = \tan^{-1}(x_1^{-1} (x_2 x_3) - (x_2 x_4)^{-1}) + \epsilon, \quad e \sim \mathcal{N}(0, 0.1) \tag{35}
\]

The standard deviations of the noise terms are set in such a way that the signal-to-noise ratio is 3:1. Thus, the variance of the signal part of the function accounts for 90% of the total variance. For each function, we generate a set of 400 points with 50% randomly chosen for training and the rest for validation. For consistent evaluation of the different data sets, we scale each of the input variables \( \mathbf{x} \) and the output \( y \) linearly to the range \([-1, 1]\). The RBF kernel is thus defined as \( K(x_i, x_j) = \exp(-\|x_i - x_j\|^2/(2\gamma)) \), where \( \gamma \) is the dimensionality of \( \mathbf{x} \) and \( \gamma \) is set to 10, 1, 0.1, and 0.01 separately.

Fig. 8 shows the results on the three benchmarks for different kernel parameter values. The curves for the elbow size can help us to choose an optimal value for \( \gamma \). If the elbow set contains most of the data points, it indicates that the regression function is probably too complex with a high chance of overfitting the data. On the other hand, if it contains too few points, the regression function is unlikely to fit the data well. The curves for the estimated generalization error in Fig. 8(d)–(f) support this speculation. Therefore, we choose \( \gamma = 1 \) for \( F1 \) and \( \gamma = 0.5 \) for \( F2 \) and \( F3 \).
Based on the kernel parameter values selected, we apply the $\varepsilon$-path algorithm with different $\lambda$ values. The results are shown in Fig. 9. The generalization error curves exhibit a similar trend as that of the sinc-function data. When $\lambda$ is very large, the regression functions underfit the data. On the contrary, when $\lambda$ is too small such as $0.0001$, overfitting always occurs when $\varepsilon$ also decreases so as to be small in the $\varepsilon$-path algorithm. The generalization error first decreases and then increases, giving a U-shape curve. $\lambda$ takes desirable value from $[0.1, 0.001]$ for $F_1$, $[1, 0.01]$ for $F_2$, and $[0.1, 0.001]$ for $F_3$. For each data set, the generalization error curves for these $\lambda$ values are almost overlapped most of the time; it generally decreases rapidly in the beginning and then only changes slightly as the algorithm continues. Although the generalization error shows no significant further improvement after the initial stage, the number of SVs continues to increase. Because the elbow size only changes slightly after a few iterations, the generalization of the regression function becomes stable.

VII. TWO REAL-WORLD DATA SETS

We further apply the $\varepsilon$-path algorithm to two real-world regression data sets, abalone and housing, from the UCI Machine Learning Repository. The abalone data set has 4177 examples and each example has eight attributes, while the housing data set has 506 examples and each example has 13 attributes. In each data set, the value of the output attribute to predict is provided for each example. We first scale each attribute linearly to the range $[0, 1]$. Each data set is randomly partitioned into two subsets with $60\%$ of the data for training and $40\%$ for validation. As we can see from Fig. 10(a), (b), (d), and (e), the elbow size and the generalization error again show similar behaviors as before for different kernel parameter values along the $\varepsilon$-path when $\lambda$ is set to $0.1$. For the abalone data set, the elbows include very few points when $\gamma \leq 10$ or when $\gamma \equiv 1$. We choose $\gamma \equiv 0.1$ as a better choice, because at this value the elbows contain $1\%$ of the points and become stable in the algorithm. For the housing data set, we choose $\gamma = 0.5$. The generalization errors for different $\lambda$ values are plotted in Fig. 10(c) and (f). We observe that $\varepsilon$-paths of solutions are very similar to each other for moderate values of $\lambda$ from $[0.1, 0.001]$. Hence, we choose such a value for $\lambda$ in practice and explore the $\varepsilon$-path until the point after which the generalization error does not change significantly. This ensures that a good regression function with the sparseness property can be obtained efficiently.
VIII. CONCLUSION AND FUTURE WORK

In this paper, we analyze the SVR solution paths by considering two types of optimality conditions, i.e., equality and inequality conditions. From the equality conditions, we obtain the solution updating formula (21) for both $\lambda$ and $\epsilon$. If we do not take the inequality conditions into account, the updating formula (21) is very general and can be used to compute the solution for any hyperparameter value. Note that it bears resemblance to ridge regression. In ridge regression, the optimality condition is simply the equation obtained by setting the first derivative to zero, which gives the optimal solution in a closed form. Because there is only one (equality) optimality condition for ridge regression, the solutions for different hyperparameter values can be computed directly without having to trace the solution path. The complexity of updating the solution from the current one is equivalent to that of computing the next solution directly. This is not the case for SVR though. Due to the sparseness property of the function, the optimality conditions contain both equality and inequality conditions. The solution path consists of multiple (linear) segments where each segment is governed by the equality conditions. Any solution within a segment can be explored easily by focusing on the equality conditions only. On the other hand, the inequality conditions correspond to the boundaries between consecutive segments. They serve to localize the valid ranges of the equality conditions. As the solution path proceeds, it needs to monitor the inequality conditions to identify the first one that no longer holds. The inequalities determine which data points are involved in the equality constraints and thus determine the breakpoints. These relationships are illustrated in Fig. 11. The solution path algorithm then sequentially extends from one segment of the path to the next. This analysis is general and can also be extended to other sparse modeling methods such as SVC [7] and LASSO [27].
Fig. 10. The $\epsilon$-path results on the [(a)–(c)] abalone data set and [(d)–(f)] housing data set. $\lambda = 0 : 1$ in (a), (b), (d), and (e). $\lambda = 0 : 1$ in (c) and $\lambda = 0 : 5$ in (f).

Fig. 11. Each curve corresponds to an inequality condition. The region corresponding to the candidate solutions specified by all the inequality conditions can be explored freely to update the solution. However, when the solution update goes beyond this region, some procedures are needed to continue the path exploration.

Our proposed $\epsilon$-path algorithm has some advantages over the $\lambda$-path algorithm.

- The $\epsilon$-path algorithm has a very simple initialization, while the $\lambda$-path algorithm requires solving a linear programming problem.
- The $\lambda$-path algorithm may need to randomly move the tube looking for valid points during the execution, while the $\epsilon$-path algorithm can be explored more easily.
- A sparse regression function can be obtained from the $\epsilon$-path algorithm after very few iterations. From the experiments, we notice that the regression function is not very sensitive to the $\lambda$ value. Thus, it is very efficient to explore the 2-D solution space of $\epsilon$-SVR by executing the $\epsilon$-path algorithm several times with different $\lambda$ values.

Therefore, in practical applications, we recommend applying the $\epsilon$-path algorithm for some predetermined $\lambda$ values.

This paper discusses two solution path algorithms, either of which is with respect to only one hyperparameter while the other hyperparameters are fixed. When the dimensionality of the solution space is one, every solution can be explored as the solution path is traced in the direction of either increasing or decreasing the hyperparameter value. However, many models contain multiple hyperparameters. When the dimensionality of the solution space is larger than one, it becomes difficult to explore all possible solutions using the path-following approach. As in the SVR problem, the solution path is piecewise linear with respect to either $\epsilon$ or $\lambda$. For any given $\lambda$ value, every solution along the $\epsilon$ path can be computed as $\epsilon$ changes. This is similar to the case where $\lambda$ is free but $\epsilon$ is fixed. Although these two kinds of path algorithms can be interchangeably used, it is challenging to integrate two 1-D path algorithms to explore the entire 2-D solution space $(\lambda, \epsilon)$. One solution to overcome this difficulty is to execute the 1-D path algorithm several times with different values of another hyperparameter, such as our recommendation for the SVR problem. The problem becomes more severe if more hyperparameters are involved. The attempt to explore every solution in the high-dimensional solution space is infeasible. On the other hand, it is not necessary to explore the whole solution space, because it may involve a great deal of unnecessary computation. We are often interested in just a small portion of the solution space in which there exist some solutions.
that can generalize well. One possible approach to this problem is to take the performance on the validation data into account. The path of solutions can be traced according to the direction of decreasing the prediction error estimated from the validation data. In each iteration, the new solution from the path-following algorithm will provide better performance than in the last iteration. Thus, a locally optimal solution in the high-dimensional solution space can be found efficiently.

APPENDIX

We define some notations used in the paper

\[ \alpha_R = (\alpha_i) \quad \forall i \in \mathcal{E}_R \]  
\[ y_R = (y_i) \quad \forall i \in \mathcal{E}_R \]  
\[ \alpha_L = (\alpha^L_i) \quad \forall i \in \mathcal{E}_L \]  
\[ y_L = (y_i) \quad \forall i \in \mathcal{E}_L \]  
\[ \alpha^*_R = (\alpha^*_i) \quad \forall i \in \mathcal{E}_R \]  
\[ \alpha^*_L = (\alpha^*_i) \quad \forall i \in \mathcal{E}_L \]  
\[ K = \begin{bmatrix} K(x_i, x_j) \end{bmatrix}_{i,j} \quad \forall i, j \in \mathcal{E}_R \cup \mathcal{E}_L \]  
\[ K_{RL} = \begin{bmatrix} K(x_i, x_j) \end{bmatrix}_{i,j} \quad \forall i \in \mathcal{E}_R \cup \mathcal{E}_L, \quad j \in \mathcal{R} \cup \mathcal{L} \]  
\[ \mathbf{1}_R = (1)_{p_1 \times 1} \]  
\[ \mathbf{1}_L = (1)_{p_2 \times 1} \]  

and

\[ K^a = \begin{bmatrix} 0 & \mathbf{1}^T \\ \mathbf{1} & K_L \end{bmatrix} \]  
\[ g_L = \begin{bmatrix} y_L \\ 0 \end{bmatrix} \]  
\[ 1^a = \begin{bmatrix} 0 \\ \mathbf{1}_R \\ -\mathbf{1}_L \end{bmatrix} \]  
\[ K^b = \begin{bmatrix} 0 & \mathbf{1}^T \\ 0 & K_{RL} \end{bmatrix} \]  

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