A Divide-and-Conquer Solver for Kernel Support Vector Machines

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Abstract

The kernel support vector machine (SVM) is one of the most widely used classification methods; however, the amount of computation required becomes the bottleneck when facing millions of samples. In this paper, we propose and analyze a novel divide-and-conquer solver for kernel SVMs (DC-SVM). In the division step, we partition the kernel SVM problem into smaller subproblems by clustering the data, so that each subproblem can be solved independently and efficiently. We show theoretically that the support vectors identified by the subproblem solution are likely to be support vectors of the entire kernel SVM problem, provided that the problem is partitioned appropriately by kernel clustering. In the conquer step, the local solutions from the subproblems are used to initialize a global coordinate descent solver, which converges quickly as suggested by our analysis. By extending this idea, we develop a multilevel Divide-and-Conquer SVM algorithm with adaptive clustering and early prediction strategy, which outperforms state-of-the-art methods in terms of training speed, testing accuracy, and memory usage. As an example, on the covtype dataset with half-a-million samples, DC-SVM is 7 times faster than LIBSVM in obtaining the exact SVM solution (to within $10^{-6}$ relative error) which achieves 96.15% prediction accuracy. Moreover, with our proposed early prediction strategy, DC-SVM achieves about 96% accuracy in only 12 minutes, which is more than 100 times faster than LIBSVM.

1. Introduction

The support vector machine (SVM) (Cortes & Vapnik, 1995) is probably the most widely used classifier in varied machine learning applications. For problems that are not linearly separable, kernel SVM uses a “kernel trick” to implicitly map samples from input space to a high-dimensional feature space, where samples become linearly separable. Due to its importance, optimization methods for kernel SVM have been widely studied (Platt, 1998; Joachims, 1998), and efficient libraries such as LIBSVM (Chang & Lin, 2011) and SVMLight (Joachims, 1998) are well developed. However, the kernel SVM is still hard to scale up when the sample size reaches more than one million instances. The bottleneck stems from the high computational cost and memory requirements of computing and storing the kernel matrix, which in general is not sparse. By approximating the kernel SVM objective function, approximate solvers (Zhang et al., 2012; Le et al., 2013) avoid high computational cost and memory requirement, but suffer in terms of prediction accuracy.

In this paper, we propose a novel divide and conquer approach (DC-SVM) to efficiently solve the kernel SVM problem. DC-SVM achieves faster convergence speed compared to state-of-the-art exact SVM solvers, as well as better prediction accuracy in much less time than approximate solvers. To accomplish this performance, DC-SVM first divides the full problem into smaller subproblems, which can be solved independently and efficiently. We theoretically show that the kernel kmeans algorithm is able to minimize the difference between the solution of subproblems and of the whole problem, and support vectors identified by subproblems are likely to be support vectors of the whole problem. However, running kernel kmeans on the whole dataset is time consuming, so we apply a two-step kernel kmeans procedure to efficiently find the partition. In the conquer step, the local solutions from the subproblems are “glued” together to yield an initial point for the global problem. As suggested by our analysis, the coordinate descent method in the final stage converges quickly to the global optimal.

Empirically, our proposed Divide-and-Conquer Kernel SVM solver can reduce the objective function value much faster than existing SVM solvers. For example, on the covtype dataset with half a million samples, DC-SVM can find a globally optimal solution (to within $10^{-6}$ accuracy) within 3 hours on a single machine with 8 GBytes
RAM, while the state-of-the-art LIBSVM solver takes more than 22 hours to achieve a similarly accurate solution (which yields 96.15% prediction accuracy). More interestingly, due to the closeness of the subproblem solutions to the global solution, we can employ an early prediction approach, using which DC-SVM can obtain high test accuracy extremely quickly. For example, on the Covtype dataset, by using early prediction DC-SVM achieves 96.03% prediction accuracy within 12 minutes, which is more than 100 times faster than LIBSVM (see Figure 3e for more details).

The rest of the paper is outlined as follows. We propose the single-level DC-SVM in Section 3, and extend it to the multilevel version in Section 4. Experimental comparison with other state-of-the-art SVM solvers is shown in Section 5. The relationship between DC-SVM and other methods is discussed in Section 2, and the conclusions are given in Section 6. Extensive experimental comparisons are included in the Appendix.

2. Related Work

Since training SVM requires a large amount of memory, it is natural to apply decomposition methods (Platt, 1998), where each time only a subset of variables are updated. To speedup the decomposition method, (Pérez-Cruz et al., 2004) proposed a double chunking approach to maintain a chunk of important samples, and the shrinking technique (Joachims, 1998) is also widely used to eliminate unimportant samples.

To speed up kernel SVM training on large-scale datasets, it is natural to divide the problem into smaller subproblems, and combine the models trained on each partition. (Jacobs et al., 1991) proposed a way to combine models, although in their algorithm subproblems are not trained independently, while (Tresp, 2000) discussed a Bayesian prediction scheme (BCM) for model combination. (Collobert et al., 2002) partition the training dataset arbitrarily in the beginning, and then iteratively refine the partition to obtain an approximate kernel SVM solution. (Kugler et al., 2006) applied the above ideas to solve multi-class problems. (Graf et al., 2005) proposed a multilevel approach (CascadeSVM): they randomly build a partition tree of samples and train the SVM in a “cascade” way: only support vectors in the lower level of the tree are passed to the upper level. However, no earlier method appears to discuss an elegant way to partition the data. In this paper, we theoretically show that kernel k-means minimizes the error of the solution from the subproblems and the global solution. Based on this division step, we propose a simple method to combine locally trained SVM models, and show that the testing performance is better than BCM in terms of both accuracy and time (as presented in Table 1). More importantly, DC-SVM solves the original SVM problem, not just an approximated one. We compare our method with Cascade SVM in the experiments.

Another line of research proposes to reduce the training time by representing the whole dataset using a smaller set of landmark points, and clustering is an effective way to find landmark points (cluster centers). (Moody & Darken, 1989) proposed this idea to train the reduced sized problem with RBF kernel (LTPU); (Pavlov et al., 2000) used a similar idea as a preprocessing of the dataset, while (Yu et al., 2005) further generalized this approach to a hierarchical coarsen-refinement solver for SVM. Based on this idea, the k-means Nyström method (Zhang et al., 2008) was proposed to approximate the kernel matrix using landmark points. (Boley & Cao, 2004) proposed to find samples with similar α values by clustering, so both the clustering goal and training step are quite different from ours. All the above approaches focus on modeling the between-cluster (between-landmark points) relationships. In comparison, our method focuses on preserving the within-cluster relationships at the lower levels and explores the between-cluster information in the upper levels. We compare DC-SVM with LLSSVM (using k-means Nyström) and LTPU in Section 5.

There are many other approximate solvers for the kernel SVM, including kernel approximation approaches (Fine & Scheinberg, 2001; Zhang et al., 2012; Le et al., 2013), greedy basis selection (Keerthi et al., 2006), and online SVM solvers (Bordes et al., 2005). Recently, (Jose et al., 2013) proposed an approximate solver to reduce testing time. They use multiple linear hyperplanes for prediction, so the time complexity for prediction is proportional to the dimensionality instead of number of samples. Therefore they achieve faster prediction but require more training time and have lower prediction accuracy comparing to DC-SVM with early prediction strategy.

3. Divide and Conquer Kernel SVM with a single level

Given a set of instance-label pairs \( \{x_i, y_i\}, i = 1, \ldots, n, x_i \in \mathbb{R}^d \) and \( y_i \in \{1, -1\} \), the main task in training the kernel SVM is to solve the following quadratic optimization problem:

\[
\min_{\alpha} f(\alpha) = \frac{1}{2} \alpha^T Q \alpha - e^T \alpha, \text{ s.t. } 0 \leq \alpha \leq C,
\]

where \( e \) is the vector of all ones; \( C \) is the balancing parameter between loss and regularization in the SVM primal problem; \( \alpha \in \mathbb{R}^n \) is the vector of dual variables; and \( Q \) is an \( n \times n \) matrix with \( Q_{ij} = y_i y_j K(x_i, x_j) \), where \( K(x_i, x_j) \) is the kernel function. Note that, as in (Keerthi et al., 2006; Joachims, 2006), we ignore the “bias” term — indeed, in our experiments reported in Section 5, we did not observe any improvement in test accuracy by including the bias term. Letting \( \alpha^* \) denote the optimal solution of (1), the decision value for a test data \( x \) can be computed by \( \sum_{i=1}^n \alpha_i^* y_i K(x, x_i) \).

We begin by describing the single-level version of our proposed algorithm. The main idea behind our divide and conquer SVM solver (DC-SVM) is to divide the data into
smaller subsets, where each subset can be handled efficiently and independently. The subproblem solutions are then used to initialize a coordinate descent solver for the whole problem. To do this, we first partition the dual variables into $k$ subsets $\{V_1, \ldots, V_k\}$, and then solve the respective subproblems independently

$$\min_{\alpha(c)} \frac{1}{2}(\alpha(c))^T Q(c,c) \alpha(c) - e^T \alpha(c), \text{ s.t. } 0 \leq \alpha(c) \leq C,$$

where $c = 1, \ldots, k$, $\alpha(c)$ denotes the subvector $\{\alpha_p | p \in V_c\}$ and $Q(c,c)$ is the submatrix of $Q$ with row and column indexes $V_c$.

The quadratic programming problem (1) has $n$ variables, and takes at least $O(n^2)$ time to solve in practice (as shown in (Menon, 2009)). By dividing it into $k$ subproblems (2) with equal sizes, the time complexity for solving the subproblems can be dramatically reduced to $O(k \cdot (\frac{n}{k})^2) = O(n^2/k)$. Moreover, the space requirement is also reduced from $O(n^2)$ to $O(n^2/k^2)$.

After computing all the subproblem solutions, we concatenate them to form an approximate solution for the whole problem $\bar{\alpha} = [\alpha(1), \ldots, \alpha(k)]$, where $\alpha(c)$ is the optimal solution for the $c$-th subproblem. In the conquer step, $\bar{\alpha}$ is used to initialize the solver for the whole problem. We show that this procedure achieves faster convergence due to the following reasons: (1) $\bar{\alpha}$ is close to the optimal solution for the whole problem $\alpha^*$, so the solver only requires a few iterations to converge (see Theorem 1); (2) the set of support vectors of the subproblems is expected to be close to the set of support vectors of the whole problem (see Theorem 2). Hence, the coordinate descent solver for the whole problem converges very quickly.

**Divide Step.** We now discuss in detail how to divide problem (1) into subproblems. In order for our proposed method to be efficient, we require $\bar{\alpha}$ to be close to or the optimal solution of the original problem $\alpha^*$. In the following, we derive a bound on $\|\alpha^* - \bar{\alpha}\|_2$ by first showing that $\bar{\alpha}$ is the optimal solution of (1) with an approximate kernel.

**Lemma 1.** $\bar{\alpha}$ is the optimal solution of (1) with kernel function $K(x_i, x_j)$ replaced by

$$\tilde{K}(x_i, x_j) = I(\pi(x_i), \pi(x_j))K(x_i, x_j),$$

where $\pi(x_i)$ is the cluster that $x_i$ belongs to; $I(a, b) = 1$ iff $a = b$ and $I(a, b) = 0$ otherwise.

Based on the above lemma, we are able to bound $\|\alpha^* - \bar{\alpha}\|_2$ by the sum of between-cluster kernel values:

**Theorem 1.** Given data points $\{(x_i, y_i)\}_{i=1}^n$ with labels $y_i \in \{1, -1\}$ and a partition indicator $\{\pi(x_1), \ldots, \pi(x_n)\}$,

$$0 \leq f(\bar{\alpha}) - f(\alpha^*) \leq (1/2)C^2D(\pi),$$

where $f(\alpha)$ is the objective function in (1) and $D(\pi) = \sum_{i,j: \pi(x_i) \neq \pi(x_j)} \|K(x_i, x_j)\|$. Furthermore, $\|\alpha^* - \bar{\alpha}\|_2 \leq C^2D(\pi)/\sigma_n$ where $\sigma_n$ is the smallest eigenvalue of the kernel matrix.

The proof is provided in Appendix 8.2. In order to minimize $\|\alpha^* - \bar{\alpha}\|$, we want to find a partition with small $D(\pi)$. Moreover, a balanced partition is preferred to achieve faster training speed. This can be done by the kernel kmeans algorithm, which aims to minimize the off-diagonal values of the kernel matrix with a balancing normalization.

We now show that the bound derived in Theorem 1 is reasonably tight in practice. On a subset (10000 instances) of the covtype data, we try different numbers of clusters $k = 8, 16, 32, 64, 128$; for each $k$, we use kernel kmeans to obtain the data partition $\{V_1, \ldots, V_k\}$, and then compute $C^2D(\pi)/2$ (the right hand side of (4)) and $f(\bar{\alpha}) - f(\alpha^*)$ (the left hand side of (4)). The results are presented in Figure 1. The left panel shows the bound (in red) and the difference in objectives $f(\bar{\alpha}) - f(\alpha^*)$ in absolute scale, while the right panel shows these values in a log scale. Figure 1 shows that the bound is quite close to the difference in objectives in an absolute sense (the red and blue curves nearly overlap), especially compared to the difference in objectives when the data is partitioned randomly (this also shows effectiveness of the kernel kmeans procedure). Thus, our data partitioning scheme and subsequent solution of the subproblems leads to good approximations to the global kernel SVM problem.

![Figure 1](image)

**Figure 1:** Demonstration of the bound in Theorem 1 – our data partitioning scheme leads to good approximations to the global solution $\alpha^*$. The left plot is on an absolute scale, while the right one is on a logarithmic scale.

However, kernel kmeans has $O(n^2d)$ time complexity, which is too expensive for large-scale problems. Therefore we consider a simple two-step kernel kmeans approach as in (Ghitta et al., 2011). The two-step kernel kmeans algorithm first runs kernel kmeans on $m$ randomly sampled data points ($m \ll n$) to construct cluster centers in the kernel space. Based on these centers, each data point computes its distance to cluster centers and decides which cluster it belongs to. The algorithm has time complexity $O(mnd)$ and space complexity $O(m^2)$. In our implementation we just use random initialization for kernel kmeans, and observe good performance in practice.

A key facet of our proposed divide and conquer algorithm...
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is that the set of support vectors from the subproblems \( S := \{i \mid \alpha_i > 0\} \), where \( \alpha_i \) is the \( i \)-th element of \( \alpha \), is very close to that of the whole problem \( S := \{i \mid \alpha^*_i > 0\} \).

Letting \( \bar{f}(\alpha) \) denote the objective function of (1) with kernel \( K \) defined in (3), the following theorem shows that when \( \alpha_i = 0 \) (\( x_i \) is not a support vector of the subproblem) and \( \nabla_i \bar{f}(\alpha) \) is large enough, then \( x_i \) will not be a support vector of the whole problem.

**Theorem 2.** For any \( i \in \{1, \ldots, n\} \), if \( \alpha_i = 0 \) and

\[
\nabla_i \bar{f}(\alpha) > CD(\pi)(1 + \sqrt{nK_{\max}/\sigma_nD(\pi)}),
\]

where \( K_{\max} = \max K(x_i, x_i) \), then \( x_i \) will not be a support vector of the whole problem, i.e., \( \alpha^*_i = 0 \).

The proof is given in Appendix 8.3. In practice also, we observe that DC-SVM can identify the set of support vectors of the whole problem very quickly. Figure 2 demonstrates that DC-SVM identifies support vectors much faster than the shrinking strategy implemented in LIBSVM (Chang & Lin, 2011) (we discuss these results in more detail in Section 4).

**Conquer Step.** After computing \( \alpha \) from the subproblems, we use \( \alpha \) to initialize the solver for the whole problem. In principle, we can use any SVM solver in our divide and conquer framework, but we focus on using the coordinate descent method as in LIBSVM to solve the whole problem. The main idea is to update one variable at a time, and always choosing the \( \alpha_i \) with the largest gradient value to update. The benefit of applying coordinate descent is that we can avoid a lot of unnecessary access to the kernel matrix entries if \( \alpha_i \) never changes from zero to nonzero. Since \( \alpha \)'s are close to \( \alpha^* \), the \( \alpha \)-values for most vectors that are not support vectors will not become nonzero, and so the algorithm converges quickly.

4. **Divide and Conquer SVM with multiple levels**

There is a trade-off in choosing the number of clusters \( k \) for a single-level DC-SVM with only one divide and conquer step. When \( k \) is small, the subproblems have similar sizes as the original problem, so we will not gain much speedup. On the other hand, when we increase \( k \), time complexity for solving subproblems can be reduced, but the resulting \( \alpha \) can be quite different from \( \alpha^* \) according to Theorem 1, so the conquer step will be slow. Therefore, we propose to run DC-SVM with multiple levels to further reduce the time for solving the subproblems, and meanwhile still obtain \( \alpha \) values that are close to \( \alpha^* \).

In multilevel DC-SVM, at the \( l \)-th level, we partition the whole dataset into \( k^l \) clusters \( \{V_1^{(l)}, \ldots, V_k^{(l)}\} \), and solve those \( k^l \) subproblems independently to get \( \alpha^{(l)} \). In order to solve each subproblem efficiently, we use the solutions from the lower level \( \alpha^{(l+1)} \) to initialize the solver at the \( l \)-th level, so each level requires very few iterations. This allows us to use small values of \( k \), for example, we use \( k = 4 \) for all the experiments. In the following, we discuss more insights to further speed up our procedure.

**Adaptive Clustering.** The two-step kernel kmeans approach has time complexity \( O(nmd) \), so the number of samples \( m \) cannot be too large. In our implementation we use \( m = 1000 \). When the data set is very large, the performance of two-step kernel kmeans may not be good because we sample only a few data points. This will influence the performance of DC-SVM.

To improve the clustering for DC-SVM, we propose the following adaptive clustering approach. The main idea is to explore the sparsity of \( \alpha \) in the SVM problem, and sample

![Figure 2: Our multilevel DC-SVM algorithm computes support vectors for subproblems during the “conquer” phase. The above plots show that DC-SVM identifies support vectors more accurately (Figure 2a, 2b, 2e, 2f) than cascade SVM, and more quickly than the shrinking strategy in LIBSVM.](image-url)
from the set of support vectors to perform two-step kernel kmeans. Suppose we are at the l-th level, and the current set of support vectors is defined by $\tilde{S} = \{i \mid \alpha_i > 0\}$. Suppose the set of support vectors for the final solution is given by $S^* = \{i \mid \alpha_i^* > 0\}$. We can define the sum of off-diagonal elements on $\tilde{S} \cup S^*$ as $D_{\tilde{S} \cup S^*}(\pi) = \sum_{i,j \in \tilde{S} \cup S^*} \pi(x_i) \neq \pi(x_j) |K(x_i, x_j)|$. The following theorem shows that we can refine the bound in Theorem 1:

**Theorem 3.** Given data points $x_1, \ldots, x_n$ and a partition \{$V_1, \ldots, V_k$\} with indicators $\pi$,

$$0 \leq f(\tilde{\alpha}) - f(\alpha^*) \leq (1/2)C^2D_{\tilde{S} \cup S^*}(\pi).$$

Furthermore, $||\alpha^* - \tilde{\alpha}||_2^2 \leq C^2D_{\tilde{S} \cup S^*}(\pi)/\sigma_n$.

The proof is given in Appendix 8.4. The above observations suggest that if we know the set of support vectors $\tilde{S}$ and $S^*$, $||\alpha^* - \tilde{\alpha}||$ only depends on whether we can obtain a good partition of $\tilde{S} \cup S^*$. Therefore, we can sample $m$ points from $\tilde{S} \cup S^*$ instead of the whole dataset to perform the clustering. The performance of two-step kernel kmeans depends on the sampling rate; we enhance the sampling rate from $m/n$ to $m/(|\tilde{S} \cup S^*|)$. As a result, the performance significantly improves when $|\tilde{S} \cup S^*| \ll n$.

In practice we do not know $S^*$ or $\tilde{S}$ before solving the problem. However, both Theorem 2 and experiments shown in Figure 2 suggest that we have a good guess of support vectors even at the bottom level. Therefore, we can use the lower level support vectors as a good guess of the upper level support vectors. More specifically, after computing $\tilde{\alpha}^l$ from level $l$, we can use its support vector set $\tilde{S}^l := \{i \mid \tilde{\alpha}_i^l > 0\}$ to run two-step kernel kmeans for finding the clusters at the $(l-1)$-th level. Using this strategy, we obtain progressively better partitioning as we approach the original problem at the top level.

### Early identification of support vectors.

We first run LIBSVM to obtain the final set of support vectors, and then run DC-SVM with various numbers of clusters $4^3, 4^4, \ldots, 4^6$ (corresponding to level 5, 4, $\ldots$, 0 for multi-level DC-SVM). We show the precision and recall for the upper level support vectors. More specifically, after computing the lower level support vectors as a good guess of the upper level support vectors, we use the lower level support vectors as a good guess of the upper level support vectors.

The details of the experiments are given in Appendix 8.4. The above observations show that we can refine the initialization by solving the SVM problem induced by all support vectors at the first level.

### Early prediction based on the $l$-th level solution.

Computing the exact kernel SVM solution can be quite time consuming, so it is important to obtain a good model using limited time and memory. We now propose a way to efficiently predict the label of unknown instances using the lower-level models $\alpha^l$. We will see in the experiments that prediction using $\alpha^l$ from a lower level $l$ already can achieve near-optimal testing performance.

When the $l$-th level solution $\alpha^l$ is computed, a naive way to predict a new instance $x$’s label $\hat{y}$ is:

$$\hat{y} = \text{sign} \left( \sum_{i=1}^{n} y_i \alpha_i^l K(x, x_i) \right).$$

Another way to combine the models trained from $k$ clusters is to use the probabilistic framework proposed in the Bayesian Committee Machine (BCM) (Tresp, 2000). However, as we show below, both these methods do not give good prediction accuracy when the number of clusters is large.

Instead, we propose the following early prediction strategy. From Lemma 1, $\alpha$ is the optimal solution to the SVM dual problem (1) on the whole dataset with the approximated kernel $K$ defined in (3). Therefore, we propose to use the same kernel function $K$ in the testing phase, which leads to the prediction

$$\sum_{c=1}^{k} \sum_{i \in \mathcal{V}_c} y_i \alpha_i K(x, x_i) = \sum_{i \in \mathcal{V}_{\pi(x)}} y_i \alpha_i K(x, x_i),$$

where $\pi(x)$ can be computed by finding the nearest cluster center. Therefore, the testing procedure for early prediction is: (1) find the nearest cluster that $x$ belongs to, and then (2) use the model trained by data within that cluster to compute the decision value.

We compare this method with prediction by (5) and BCM in Table 1. The results show that our proposed testing scheme is better in terms of test accuracy. We also compare average testing time per instance in Table 1, and our proposed method is much more efficient as we only evaluate $K(x, x_i)$ for all $x_i$ in the same cluster as $x$, thus reducing the testing time from $O(|\tilde{S}|d)$ to $O(|\tilde{S}|d/k)$, where $\tilde{S}$ is the set of support vectors.

### Refine solution before solving the whole problem.

Before training the final model at the top level using the whole dataset, we can refine the initialization by solving the SVM problem induced by all support vectors at the first level.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>$k = 50$</th>
<th>$k = 100$</th>
<th>$k = 50$</th>
<th>$k = 100$</th>
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<td>89.5% / 3.3ms</td>
<td>94.6% / 2.4ms</td>
<td>92.7% / 2.5ms</td>
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<tr>
<td>covtype</td>
<td>96.1% / .4ms</td>
<td>92.6% / 2.4ms</td>
<td>92.6% / 2.4ms</td>
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</table>

*Table 1: Comparing prediction methods using a lower level model. Our proposed early prediction strategy is better in terms of prediction accuracy and testing time per sample (time given in milliseconds).*
level below the final level. As proved in Theorem 2, the support vectors of lower level models are likely to be the support vectors of the whole model, so this will give a more accurate solution, and only requires us to solve a problem with $O(|S^{(1)}|)$ samples, where $S^{(1)}$ is the set of support vectors at the first level. Our final algorithm is given in Algorithm 1.

Algorithm 1 Divide and Conquer SVM

Input: Training data $\{(x_i, y_i)\}_{i=1}^n$, balancing parameter $C$, kernel function.

Output: The SVM dual solution $\alpha$.

for $l = 1, \ldots, \maxI$ do

Set number of clusters in the current level $k_l = k_l^\text{init}$;  

if $l = 1$ then

Sample $m$ points $\{x_1, \ldots, x_m\}$ from the whole training set;

else

Sample $m$ points $\{x_1, \ldots, x_m\}$ from $\{x_i \mid c_i^{(l+1)} > 0\}$;

end

Run kernel kmeans on $\{x_1, \ldots, x_m\}$ to get cluster centers $c_1, \ldots, c_m$;

Obtain partition $V_1, \ldots, V_{k_l}$ for all data points;

for $c = 1, \ldots, k_l$ do

Obtain $\alpha_c^{(l)}$ by solving SVM for the data in the $c$-th cluster $V_c$ with $\alpha_c^{(l+1)}$ as the initial point ($\alpha_c^{l+1}$ is set to 0);

end

Refine solution: Compute $\alpha^{(0)}$ by solving SVM on $\{x_i \mid c_i^{(1)} \neq 0\}$ using $\alpha^{(1)}$ as the initial point;

Solve SVM on the whole data using $\alpha^{(0)}$ as the initial point;

5. Experimental Results

We now compare our proposed algorithm with other SVM solvers. All the experiments are conducted on an Intel 2.66GHz CPU with 8G RAM. We use 7 benchmark datasets as shown in Table 3. The cifar dataset can be downloaded from http://www.cs.toronto.edu/~kriz/cifar.html, and other datasets can be downloaded from http://www.csie.ntu.edu.tw/

Table 2: Comparison on real datasets.

<table>
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<th>$C = 8, \gamma = 2^{16}$</th>
<th>$C = 512, \gamma = 2^{64}$</th>
<th>$C = 32, \gamma = 2$</th>
<th>$C = 8, \gamma = 2^{16}$</th>
<th>$C = 512, \gamma = 2^{64}$</th>
<th>$C = 256, \gamma = 0.5$</th>
<th>$C = 512, \gamma = 2^{16}$</th>
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<td>time(s) acc(%)</td>
<td>time(s) acc(%)</td>
<td>time(s) acc(%)</td>
<td>time(s) acc(%)</td>
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<td>1977 87.02</td>
<td>261 94.9</td>
<td>672 96.12</td>
<td>670 99.13</td>
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<td>10287 99.85</td>
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<tr>
<td>DC-SVM</td>
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<td>10485 99.28</td>
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<td>83631 96.15</td>
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<td>3514 93.2</td>
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<td>6700 92.13</td>
<td>171400 99.85</td>
<td>171400 99.85</td>
</tr>
<tr>
<td>CascadesVM</td>
<td>17.1 98.08</td>
<td>6148 86.8</td>
<td>849 93.0</td>
<td>5600 89.51</td>
<td>3515 98.1</td>
<td>1155 91.2</td>
<td>64151 99.83</td>
<td>64151 99.83</td>
</tr>
<tr>
<td>LLSVM</td>
<td>38 98.23</td>
<td>9745 86.5</td>
<td>1212 92.8</td>
<td>4544 84.21</td>
<td>2853 97.74</td>
<td>3015 91.5</td>
<td>63121 97.64</td>
<td>63121 97.64</td>
</tr>
<tr>
<td>FastFood</td>
<td>87 95.95</td>
<td>13357 80.3</td>
<td>851 91.6</td>
<td>8550 80.1</td>
<td>5563 96.47</td>
<td>2191 91.6</td>
<td>14917 96.5</td>
<td>14917 96.5</td>
</tr>
<tr>
<td>SpSVM</td>
<td>20 94.92</td>
<td>21335 85.6</td>
<td>3121 90.4</td>
<td>15113 83.37</td>
<td>6235 95.3</td>
<td>5124 90.5</td>
<td>121563 96.3</td>
<td>121563 96.3</td>
</tr>
<tr>
<td>LTPU</td>
<td>248 96.64</td>
<td>17418 85.16</td>
<td>1695 92.0</td>
<td>11532 83.25</td>
<td>4005 96.12</td>
<td>5100 92.1</td>
<td>105210 97.82</td>
<td>105210 97.82</td>
</tr>
</tbody>
</table>

Table 3: Dataset statistics

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Number of training samples</th>
<th>Number of testing samples</th>
<th>d</th>
</tr>
</thead>
<tbody>
<tr>
<td>icnn1</td>
<td>49990</td>
<td>91701</td>
<td>22</td>
</tr>
<tr>
<td>cifar</td>
<td>50000</td>
<td>10000</td>
<td>3072</td>
</tr>
<tr>
<td>census</td>
<td>159,619</td>
<td>39,904</td>
<td>409</td>
</tr>
<tr>
<td>covtype</td>
<td>464,810</td>
<td>116,202</td>
<td>54</td>
</tr>
<tr>
<td>webspam</td>
<td>280,000</td>
<td>70,000</td>
<td>254</td>
</tr>
<tr>
<td>kddcup99</td>
<td>4,898,431</td>
<td>311,029</td>
<td>125</td>
</tr>
<tr>
<td>mnist8m</td>
<td>8,000,000</td>
<td>100,000</td>
<td>784</td>
</tr>
</tbody>
</table>

"cjlin/libsvmtools/datasets" or the UCI data repository. We use the raw data without scaling for two image datasets cifar and mnist8m, while features in all the other datasets are linearly scaled to $[0, 1]$. mnist8m is a digital recognition dataset with 10 numbers, we follow the procedure in (Zhang et al., 2012) to transform it into a binary classification problem by classifying round digits and non-round digits. Similarly, we transform cifar into a binary classification problem by classifying animals and non-animals. We use a random 80%-20% split for covtype, webspam, kddcup99, a random 80%-20% split for mnist8m (used in the original paper (Loosli et al., 2007)), and the original training/testing split for icnn1 and cifar.

Competing Methods: We include the following exact SVM solvers (LIBSVM, CascadeSVM), approximate SVM solvers (SpSVM, LLSVM, FastFood, LTPU), and online SVM (LaSVM) in our comparison:

1. LIBSVM: the implementation in the LIBSVM library (Chang & Lin, 2011) with a small modification to handle SVM without the bias term – we observe that LIBSVM has similar test accuracy with/without bias. We also include the results for using LIBSVM with random 1/5 subsamples on each dataset in Table 2.
2. Cascade SVM: we implement cascade SVM (Graf et al., 2005) using LIBSVM as the base solver.
3. SpSVM: Greedy basis selection for nonlinear SVM (Keerthi et al., 2006).
4. LLSVM: improved Nyström method for nonlinear SVM by (Wang et al., 2011).
5. FastFood: use random Fourier features to approximate the kernel function (Le et al., 2013). We solve the resulting linear SVM problem by the dual coordinate descent solver in LIBLINEAR.
6. LTPU: Locally-Tuned Processing Units proposed in (Moody & Darken, 1989). We set $\gamma$ equal to the
**Parameter Setting:** We first consider the RBF kernel
\[ K(x_i, x_j) = \exp(-\gamma \|x_i - x_j\|^2) \]
and kernel parameter \( \gamma \) by 5-fold cross validation on a grid of points:
\[ C = [2^{-10}, 2^{-9}, \ldots, 2^{10}] \]
and \( \gamma = [2^{-10}, \ldots, 2^{10}] \) for ijcnn1, census, covtype, webspam, and kddcup99. The average distance between samples for un-scaled image datasets mnist8m and cifar is much larger than other datasets, so we test them on smaller \( \gamma \)'s; \( \gamma = [2^{-30}, 2^{-29}, \ldots, 2^{-10}] \). Regarding the parameters for DC-SVM, we use 5 levels (\( l_{\max} = 4 \)) and \( k = 4 \), so the five levels have 1, 4, 16, 64 and 256 clusters respectively. For DC-SVM (early), we stop at the level with 64 clusters. The following are parameter settings for other methods in Table 2: the rank is set to be 3000 in LLSVM; number of Fourier features is 3000 in Fastfood\(^1\); number of clusters is 3000 in LTPU; number of basis vectors is 200 in SpSVM; the tolerance in the stopping condition for LIBSVM and DC-SVM is set to \( 10^{-3} \) (the default setting of LIBSVM); for LaSVM we set the number of passes to be 1; for CascadeSVM we output the results after the first round.

**Experimental Results with RBF kernel:** Table 2 presents time taken and test accuracies. Experimental results show that the early prediction approach in DC-SVM achieves near-optimal test performance. By going to the top level (handling the whole problem), DC-SVM achieves better test performance but needs more time. Table 2 only gives the comparison on one setting; it is natural to ask, for example, about the performance of LIBSVM with a looser stopping condition, or Fastfood with varied number of Fourier features. Therefore, for each algorithm we change the parameter settings and present the detailed experimental results in Figure 3 and Figure 5 in Appendix.

Figure 3 shows convergence results with time – in 3a, 3b, 3c the relative error on the y-axis is defined as \( \frac{f(\alpha) - f(\alpha^*)}{|f(\alpha^*)|} \), where \( \alpha^* \) is computed by running LIBSVM with \( 10^{-8} \) accuracy. Online and approximate solvers are not included in this comparison as they do not solve

\(^1\)In Fastfood we control the number of blocks so that number of Fourier features is close to 3000 for each dataset.
the exact kernel SVM problem. We observe that DC-SVM achieves faster convergence in objective function compared with the state-of-the-art exact SVM solvers. Moreover, DC-SVM is also able to achieve superior test accuracy in lesser training time as compared with approximate solvers. Figure 3d, 3e, 3f compare the efficiency in achieving different testing accuracies. We can see that DC-SVM consistently achieves more than 50 fold speedup while achieving the same test accuracy with LIBSVM.

**Experimental Results with varying values of C, γ:** As shown in Theorem 1 the quality of approximation depends on $D(\pi)$, which is strongly related to the kernel parameters. In the RBF kernel, when $\gamma$ is large, a large portion of kernel entries will be close to 0, and $D(\pi)$ will be small so that $\bar{\alpha}$ is a good initial point for the top level. On the other hand, when $\gamma$ is small, $\bar{\alpha}$ may not be close to the optimal solution. To test the performance of DC-SVM under different parameters, we conduct the comparison on a wide range of parameters $(C = [2^{-10}, 2^{-6}, 2^1, 2^6, 2^{10}], \gamma = [2^{-10}, 2^{-6}, 2^1, 2^6, 2^{10}])$. The results on the ijcnn1, covtype, webspam, and census datasets are shown in Tables 7, 8, 9, 10. We observe that even when $\gamma$ is small, DC-SVM is still 1-2 times faster than LIBSVM: among all the 100 settings, DC-SVM is faster on 96/100 settings. The reason is that even when $\bar{\alpha}$ is not so close to $\alpha$, using $\bar{\alpha}$ as the initial point is still better than initialization with a random or zero vector. On the other hand, DC-SVM (early) is extremely fast, and achieves almost the same or even better accuracy when $\gamma$ is small (as it uses an approximated kernel). In Figure 6, 8, 7, 9 we plot the performance of DC-SVM and LIBSVM under various $C$ and $\gamma$ values, the results indicate that DC-SVM (early) is more robust to parameters. Note that DC-SVM (early) can be viewed as solving SVM with a different kernel $K$, which focuses on “within-cluster” information, and there is no reason to believe that the global kernel $K$ always yields better test accuracy than $\hat{K}$. The accumulated runtimes are shown in Table 4.

**Experimental Results with polynomial kernel:** To show that DC-SVM is efficient for different types of kernels, we further conduct experiments on covtype and webspam datasets for the degree-3 polynomial kernel $K(x_i, x_j) = (\eta + \gamma x_i^T x_j)^3$. For the polynomial kernel, the parameters chosen by cross validation are $C = 2, \gamma = 1$ for covtype, and $C = 8, \gamma = 16$ for webspam. We set $\eta = 0$, which is the default setting in LIBSVM. Figures 4a and 4c compare the training speed of DC-SVM and LIBSVM for reducing the objective function value and Figures 4b and 4d show the testing accuracy compared with LIBSVM and LaSVM. Since LL SVM, FastFood and LPTU are developed for shift-invariant kernels, we do not include them in our comparison. We can see that when using the polynomial kernel, our algorithm is more than 100 times faster than LIBSVM and LaSVM. One main reason for such large improvement is that it is hard for LIBSVM and LaSVM to identify the right set of support vectors when using the polynomial kernel. As shown in Figure 2, LIBSVM cannot identify 20% of the support vectors in $10^5$ seconds, while DC-SVM has a very good guess of the support vectors even at the bottom level, where number of clusters is 256. In Appendix 8.5 we show that the clustering step only takes a small portion of the time taken by DC-SVM.

**6. Conclusions**

In this paper, we have proposed a novel divide and conquer algorithm for solving kernel SVMs (DC-SVM). Our algorithm divides the problem into smaller subproblems that can be solved independently and efficiently. We show that the subproblem solutions are close to that of the original problem, which motivates us to “glue” solutions from subproblems in order to efficiently solve the original kernel SVM problem. Using this, we also incorporate an early prediction strategy into our algorithm. We report extensive experiments to demonstrate that DC-SVM significantly outperforms state-of-the-art exact and approximate solvers for nonlinear kernel SVM on large-scale datasets. The code for DC-SVM is available at http://www.cs.utexas.edu/~cjhsieh/dcsvm.

**7. Acknowledgements**

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A Divide-and-Conquer Solver for Kernel Support Vector Machines

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