Feature selection for linear SVM with provable guarantees

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We give two provably accurate feature-selection techniques for the linear SVM. The algorithms run in deterministic and randomized time respectively. Our algorithms can be used in an unsupervised or supervised setting. The supervised approach is based on sampling features from support vectors. We prove that the margin in the feature space is preserved to within \( \epsilon \)-relative error of the margin in the full feature space in the worst-case. In the unsupervised setting, we also provide worst-case guarantees of the radius of the minimum enclosing ball, thereby ensuring comparable generalization as in the full feature space in the worst-case. We present extensive experiments on real-world datasets to support our theory and demonstrate that our method is competitive and often better than prior state-of-the-art, for which there are no known provable guarantees.

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1. Introduction

The linear Support Vector Machine (SVM) is a popular classification method [6]. Few theoretical results exist for feature selection with SVMs. Empirically, numerous feature selection techniques work well [14,11]. We present a deterministic and a randomized feature selection technique for the linear SVM with a provable worst-case performance guarantee on the margin. The feature selection is unsupervised if features are selected obliquely to the data labels; otherwise, it is supervised. Our algorithms can be used in an unsupervised or supervised setting. In the unsupervised setting, our algorithm selects a number of features proportional to the rank of the data and preserves both the margin and radius of minimum enclosing ball to within \( \epsilon \)-relative error in the worst-case, thus resolving an open problem posed in Dasgupta et al. (2007) [7]. In the supervised setting, our algorithm selects \( O(\# \text{ support vectors}) \) features using only the set of support vectors, and preserves the margin for the support vectors to within \( \epsilon \)-relative error in the worst-case.

1.1. SVM basics

The training data has \( n \) points \( \mathbf{x}_i \in \mathbb{R}^d \), with respective labels \( y_i \in \{-1, +1\} \) for \( i = 1 \ldots n \). For linearly separable data, the primal SVM learning problem [6] constructs a hyperplane \( \mathbf{w}^* \) which maximizes the geometric margin (the minimum distance of a data point to the hyperplane), while separating the data. For non-separable data the “soft” 1-norm margin is maximized. The dual Lagrangian formulation [6] of the soft 1-norm SVM reduces to the following quadratic program:

\[
\max_{\alpha_i} \sum_{i=1}^{n} \alpha_i - \frac{1}{2} \sum_{i,j=1}^{n} \alpha_i \alpha_j y_i y_j \mathbf{x}_i^T \mathbf{x}_j \\
\text{subject to:} \quad \sum_{i=1}^{n} \alpha_i y_i = 0; \quad 0 \leq \alpha_i \leq C, \quad i = 1 \ldots n.
\]

(1)

The regularizer \( C \) is part of the input and the hyperplane classifier can be constructed from the \( \alpha_i \). The out-of-sample performance is related to the VC-dimension of the resulting “fat”-separator. Assuming that the data lie in a ball of radius \( B \), and that the hypothesis set consists of hyperplanes of width \( \gamma \) (the margin), then the VC-dimension is \( O(B^2/\gamma^2) \) [27]. Thus, by the VC-bound [28], the out-of-sample error is bounded by the in-sample error and a term monotonic in \( B^2/\gamma^2 \).

1.2. Our contributions

Our motivation comes from the fact that all prior feature selection methods for SVM are heuristics with no provable guarantees. We give two provably accurate feature selection techniques for linear SVM in both unsupervised and supervised settings with worst-case performance guarantees on the margin. We use the single set spectral sparsification technique (BSS, for short) [1] as our deterministic algorithm (the algorithm runs in deterministic
time, hence the name “deterministic”) and leverage-score sampling [7] as the randomized algorithm. We give a new simple method of extending unsupervised feature selection to supervised in the context of SVMs by running the unsupervised technique on the support vectors. This allows us to select only ($O$ (# support vectors) features for the deterministic algorithm ($\tilde{O}$ # support vectors) features for the randomized algorithm, where $\tilde{O}$ hides the logarithmic factors) while still preserving the margin on the support vectors. Since the support vectors are a sufficient statistic for learning a linear SVM, preserving the margin on the support vectors should be enough for learning on all the data with the sampled feature set.

More formally, let $\gamma^*$ be the optimal margin for the support vector set (which is also the optimal margin for all the data). The optimal margin $\gamma^*$ is obtained by solving the SVM optimization problem using all the features. For a suitably chosen number of features $r$, let $\gamma_r$ be the optimal margin obtained by solving the SVM optimization problem using the support vectors in the sampled feature space. We prove that the margin is preserved to within $\epsilon$-relative error: $\gamma_r \geq (1 - \epsilon) \gamma^*$. For the deterministic algorithm, the number of features required is $r = \breve{O}(\#\text{support vectors})^2$, whereas the randomized algorithm requires $r = \tilde{O}(\#\text{support vectors})^2$ features to be selected.

In the unsupervised setting, by running our algorithm on all the data, instead of only the support vectors, we get a stronger result statistically, but using more features. The deterministic algorithm requires $O(\rho(\epsilon)^2)$ features to be selected, while the randomized algorithm requires $O\left(\frac{1}{\epsilon^2}\log\left(\frac{r^2}{\epsilon^2}\right)\right)$ features to be selected, where $\rho$ is the rank of the training matrix, $\delta \in (0, 1)$ is the failure probability and $\epsilon \in (0, 1/2]$ is an accuracy parameter. Again, defining $\gamma^*$ as the optimal margin obtained by solving the SVM optimization problem using all the data in the sampled feature space, we prove that $\gamma^* \geq (1 - \epsilon) \gamma^*$. We can now also prove that the data radius is preserved, $\tilde{B} \leq (1 + \epsilon)B^2$. This means that $B_l/\gamma^*$ is preserved to within $\epsilon$-relative error, which means that the generalization error is also preserved to within $\epsilon$-relative error. The rank of the data is the effective dimension of the data, and one can construct this many combinations of pure features that preserve the geometry of the SVM exactly. What makes our result non-trivial is that we select this many pure features and preserve the geometry of the SVM to within $\epsilon$-relative error.

On the practical side, we provide an efficient heuristic for our supervised feature selection using BSS which allows our algorithm to scale-up to large datasets. While the main focus of this paper is theoretical, we compare both supervised and unsupervised versions of feature selection using single-set spectral sparsification and leverage-score sampling with the corresponding supervised and unsupervised forms of Recursive Feature Elimination (RFE) [14], Linear Programming SVM (LPSVM) [11], uniform sampling and rank-revealing QR factorization (RRQR) [13] based method of column selection. Feature selection based on the single-set spectral sparsification and leverage-score sampling technique is competitive and often better than RFE and LPSVM, and none of the prior art comes with provable performance guarantees in either the supervised or unsupervised setting.

1.3. Related work

All the prior art is heuristic in that there are no performance guarantees; nevertheless, they have been empirically tested against each other. Our algorithm comes with provable bounds, and performs comparably or better in empirical tests. Previous results do not give guarantees in part due to the difficulty of analyzing the behavior of the margin when the data (in particular the features) change. We give a short survey of the prior art: Gu-yon et al. [14] and Rakotomamonjy [24] proposed SVM based criteria to rank features based on the weights. Weston et al. [31] formulated a combinatorial optimization problem to select features by minimizing $B_l/\gamma^2$. Weston et al. [32] used the zero norm to perform error minimization and feature selection in one step. A Newton based method of feature selection using linear programming was given in Fung and Mangasarian [11]. Tan et al. [26] formulated the $\ell_0$-norm Sparse SVM using mixed integer programming. Do et al. [9] proposed margin-radius SVM (MR-SVM) which performs feature selection and ranking by optimizing the radius-margin bound with a scaling factor, and extend this work in Kalousis and Do [15] using a quadratic optimization problem with quadratic constraints. Another line of work includes the doubly regularized Support Vector Machine (DrSVM) [29] which uses a mixture of $\ell_2$-norm and $\ell_0$-norm penalties to solve the SVM optimization problem and perform variable selection. Subsequent works on DrSVM involve reducing the computational bottleneck [30,33]. Gilad-Bachrach et al. [12] formulate the margin as a function of set of features and score to sets of features according to the margin induced. Park et al. [18] studied the Fisher consistency and oracle property of penalized SVM where the dimension of inputs is fixed and showed that their method is able to identify the right model in most cases.

Paul et al. [22,23] used random projections on linear SVM in an unsupervised manner and showed theoretically that the margin and data-radius are preserved. They used linear combinations of features to obtain the provable guarantees and the motivation of their work was to speed-up the SVM running time by using random projections as a pre-processing step. Empirically, they achieved faster running time, but their out-of-sample error was worse than that of full-data. This is different from our work in several aspects. In this work, we select pure features (and not linear combinations of features) in both supervised and unsupervised settings and obtain provable performance guarantees in both cases. We show empirically, that by selecting pure features, we are able to obtain better out-of-sample performance than full-data, which was not possible in the earlier work of [22,23].

BSS and leverage-score sampling have been used to select features for $k$-means [3,4], regularized least-squares classifier [7,20] and ridge regression [21] and for core-set construction for canonical correlation analysis [19]. Our work further expands research into sparsification algorithms for machine learning.

2. Background

2.1. Notation

A, B, … denote matrices and $a, b, …$ denote column vectors; $e_i$ (for all $i = 1, …, n$) is the standard basis, whose dimensionality will be clear from context; and $I_n$ is the $n \times n$ identity matrix. The Singular Value Decomposition (SVD) of a matrix $A \in \mathbb{R}^{n \times d}$ of rank $\rho \leq \min\{n, d\}$ is equal to $A = U \Sigma V^T$, where $U \in \mathbb{R}^{n \times n}$ is an orthogonal matrix containing the left singular vectors, $\Sigma \in \mathbb{R}^{n \times d}$ is a diagonal matrix containing the singular values $\sigma_1 \geq \sigma_2 \geq … \sigma_\rho > 0$, and $V \in \mathbb{R}^{d \times d}$ is a matrix containing the right singular vectors. The spectral norm of $A$ is $\|A\|_2 = \sigma_1$.

2.2. Matrix sampling formalism

Let $A$ be the data matrix consisting of $n$ points and $d$ dimensions, $S \in \mathbb{R}^{d \times r}$ be a matrix such that $AS \in \mathbb{R}^{n \times r}$ contains $r$ columns of $A$ ($S$ is a sampling matrix as it samples $r$ columns of $A$). Let
D ∈ R^{dx} be the diagonal matrix such that ASD ∈ R^{dx} rescales the columns of A that are in AS. We will replace the sampling and rescaling matrices by a single matrix R ∈ R^{dx}, where R = SD first samples and then rescales r columns of A.

Let X be a generic data matrix in d dimensions whose rows are data vectors x_i, and let Y be the diagonal label matrix whose diagonal entries are the labels, y_i = y_j. Let \( \alpha = [\alpha_1, \alpha_2, ..., \alpha_n] \in \mathbb{R}^d \) be the vector of lagrange multipliers to be determined by solving Eq. (2). In matrix form, the SVM dual optimization problem is

\[
\max_{\alpha} \quad \mathbf{Y}^T \alpha - \frac{1}{2} \alpha^T X Y X^T \alpha
\]

subject to: \( \mathbf{Y}^T \alpha = 0; \quad 0 \leq \alpha \leq C. \) (2)

(In the above, 1, 0, C are vectors with the implied constant entry). When the data and label matrices contain all the data, we will emphasize this using the notation \( X^f \in \mathbb{R}^{d \times d}, Y^f \in \mathbb{R}^{d \times d} \). Solving (2) with these full data matrices gives a solution \( \alpha^* \). The data X, for which \( \alpha_i^* > 0 \) are support vectors and we denote by \( X^w \in \mathbb{R}^{pd}, Y^w \in \mathbb{R}^{pd} \) the data and label matrices containing only the p support vectors. Solving (2) with \( (X^w, Y^w) \) result in the same classifier. Let \( \alpha^* \) be the solution to (2) for the support vector data. The optimal separating hyperplane is

\[ w^* = (X^w)^T Y^w \alpha^*, \]

where \( X^w \) is the support vector matrix. The geometric margin is

\[ \gamma^* = 1/\|w^*\|, \quad \|w^*\| = \sum_{i=1}^{n} \alpha_i^*. \]

The radius is \( B = \min_{x} \max_{y} \|x - y\| \).

Our goal is to study how the SVM performs when run in the sampled feature space. Let X, Y be data and label matrices (such as those above) and R ∈ R^{dx} a sampling and rescaling matrix which selects r columns of X. The transformed dataset into the r selected features is \( X = XR \), and the SVM optimization problem in this feature space becomes

\[
\max_{\hat{\alpha}} \quad \mathbf{Y}^T \hat{\alpha} - \frac{1}{2} \hat{\alpha}^T X Y X^T \hat{\alpha},
\]

subject to: \( \mathbf{Y}^T \hat{\alpha} = 0; \quad 0 \leq \hat{\alpha} \leq C. \) (3)

For the supervised setting, we select features from the support vector matrix and so we set \( X = X^w \) and \( Y = Y^w \) and we select \( r_{sd} \) features using R. For the unsupervised setting, we select features from the full data matrix and so we set \( X = X^f \) and \( Y = Y^f \) and we select \( r_{sd} \) features using R.

3. Feature selection algorithms

As in the previous section, X is a generic data matrix in d dimensions. The feature selection algorithm is easy to state and breaks down into 3 basic steps:

1. Compute the right singular matrix V of X, so \( X = U \Sigma V^T \), where \( V^TV = I \).

2. Use V in the algorithm implied by Lemma 1 or Lemma 3 to construct a sampling and rescaling matrix R ∈ R^{dx} for appropriately chosen r. When using Lemma 1 to construct R, we call our method BSS-feature selection, whereas when using Lemma 3 to construct R, we call it leverage-score based feature selection.

3. Sample and rescale the columns of X to output the feature matrix X = XR.

As stated, our basic algorithm is unsupervised, using only the data matrix X that is the input. If the input data matrix depends on the labels in the training data, then the algorithm becomes supervised.

3.1. Unsupervised feature selection

The input to feature selection is the matrix \( X_{\text{aug}} = \begin{bmatrix} x_v & x_l \end{bmatrix} \), which is the full training data augmented by the vector \( x_v \) at the center of the minimum enclosing ball for the data. The matrix of \( r_{\text{s}} \) selected features is \( X^wR \).

3.2. Supervised feature selection

The input to our feature selection is the support vector matrix \( X^w \) and the matrix of \( r_{s} \) selected features is \( X^wR \). Not only do we select features, but we also shrink the number of data points.

4. Our main tools

In this section, we describe our main tools of feature selection from the numerical linear algebra literature, namely single-set spectral sparsification and leverage-score sampling. Both of these methods select columns from a matrix with orthonormal rows so that the resulting matrix is as close to orthonormal as possible. That is, by using step 2 of feature selection algorithm described in Section 3, we get \( V^TV = RR^T \).

4.1. Single-set spectral sparsification

The BSS algorithm [1] is a deterministic greedy technique that selects columns one at a time. The algorithm samples r columns in deterministic time, hence the name deterministic sampling. Consider the input matrix as a set of d column vectors \( V^f = \{v_1, v_2, ..., v_d\} \) with \( v_i \in \mathbb{R}^d \) (i = 1, ..., d). Given \( \tau > r \), we iterate over \( \tau = 0, 1, 2, ..., r - 1 \). Define the parameters \( L, U, \delta_1 \) and \( \delta_0 \) as follows:

\[
L_1 = \tau - \sqrt{\tau}, \quad \delta_1 = 1, \quad \delta_0 = \left( 1 + \sqrt{\tau} \right) / \left( 1 - \sqrt{\tau} \right), \quad \text{and} \quad U_1 = \delta_0 + \sqrt{\tau}.
\]

For \( U, L \in R \) and \( \Phi \in R^{dx} \) a symmetric positive definite definite matrix with eigenvalues \( \lambda_1, \lambda_2, ..., \lambda_d \), define \( \Phi(L, A) = \sum_{i=1}^{d} \frac{1}{\lambda_i} I \) and \( \Phi(U, A) = \sum_{i=1}^{d} \frac{1}{\lambda_i - U} I \) as the lower and upper potentials respectively. These potential functions measure how far the eigenvalues of \( A \) are from the upper and lower barriers \( \tau \) and \( L \) respectively. We define \( L(v, \delta_1, A, L) = \frac{\Phi'(V(A - \tau L))^{-1}}{V^T(A - \tau L)} + V'(A - \tau L)^{-1} V \).

At every iteration, there exists an index \( i \), and a weight \( t_i > 0 \) such that \( t_i^{-1} \leq L(v, \delta_i, A, L) \), and \( t_i^{-1} \geq U(v, \delta_i, A, U) \). Thus, there will be at most r columns selected after \( r \) iterations. The running time of the algorithm is dominated by the search for an index \( i \), satisfying \( U(v, \delta_0, A, U) \leq L(v, \delta_1, A, L) \) and computing the weight \( t_i \). One needs to compute the upper and lower potentials \( \Phi(U, A) \) and \( \Phi(L, A) \) and hence the eigenvalues of \( A \). Cost per iteration is \( O(r^2) \) and the total cost is \( O(r \tau d^2) \). Thus total running time of the algorithm is \( O(r \tau d^2) \).
O(rdε²). We include the algorithm as Algorithm 1.

**Algorithm 1.** Single-set spectral sparsification [1].

**Input:** Vf = [v₁, v₂, …, vₙ] ∈ R⁺ⁿᵈ with vᵢ ∈ Rᵈ and r > ε.

**Output:** Matrices S ∈ Rᵈᵗ, D ∈ Rᵈᵗ. 
1. Initialize A₀ = 0ᵈᵗ, S = 0ᵈᵗ, D = 0ᵈᵗ.
2. Set constants δ₁ = 1 and δ₀ = (1 + √ε/𝑡) / (1 − √ε/𝑡).
3. for r = 0 to r − 1 do
   - Let Lᵣ = r − √ε/𝑡; Uᵣ = δ₀(r + √ε/𝑡).
   - Pick index i ∈ {1, 2, ..., d} and number tᵢ > 0, such that
     \( L_i = \mathcal{U}(vᵢ, δᵢ, Aᵢ, Uᵢ) \leq \mathcal{L}(vᵢ, δᵢ, Aᵢ, Lᵢ). \)
   - Let \( tᵢ ← 1 / (\mathcal{U}(vᵢ, δᵢ, Aᵢ, Uᵢ)) \).
   - Update \( Aᵢ₊₁ = Aᵢ + tᵢvvᵢᵀ \); set \( Sᵢ₊₁ = 1 \) and \( Dᵢ₊₁ = 1 / √tᵢ. \)
4. end for
5. Multiply all the weights in D by \( \sqrt{r/3} \).
6. Return S and D.

**Algorithm 2.** Leverage-score sampling [25].

**Input:** Vf = [v₁, v₂, …, vₙ] ∈ R⁺ⁿᵈ with vᵢ ∈ Rᵈ and r > ε, \( ε ∈ (0, 1/2), δ ∈ (0, 1/2). \)

**Output:** Matrices S ∈ Rᵈᵗ, D ∈ Rᵈᵗ. 
1. Compute the normalized leverage scores \( pᵢ = \|vᵢ\|² / δ, \) for \( i = 1, 2, ..., d. \)
2. Initialize S = 0ᵈᵗ, D = 0ᵈᵗ.
3. Fix a sampling parameter \( r = O(\sqrt{\log(1/(δ, 2)})). \) Then in r random i.i.d. trials, keep the i-th feature with probability \( pᵢ \) and multiply it by the factor \( 1/√pᵢ. \) The time complexity is dominated by the time to compute the SVD of \( X. \) The i-th leverage-score characterizes the importance of the i-th feature. These scores form a probability distribution over the columns of \( X. \) since \( \sumᵢ pᵢ = 1. \) We include the algorithm as Algorithm 2.

We present the following lemma for the single-set spectral sparsification algorithm.

**Lemma 1** (BSS [1]). Given \( V ∈ R⁺ⁿᵈ \) satisfying \( VVᵀ = Iᵩ \) and \( r > ε, \) we can deterministically construct sampling and rescaling matrices \( S ∈ Rᵈᵗ \) and \( D ∈ Rᵈᵗ \) with \( R = SD, \) such that, for all \( Y ∈ Rᵈ \):
\[
(1 − √ε/𝑡) \|VVₚY\|₂ \leq \|VVᵢYᵢ\|₂ \leq (1 + √ε/𝑡) \|VVᵢYᵢ\|₂.
\]

We now present a slightly modified version of Lemma 1 for our theorems.

**Lemma 2.** Given \( V ∈ R⁺ⁿᵈ \) satisfying \( VVᵀ = Iᵩ \) and \( r > ε, \) we can deterministically construct sampling and rescaling matrices \( S ∈ Rᵈᵗ \) and \( D ∈ Rᵈᵗ \) such that for \( R = SD, \) \( \|VVᵢVᵢ − VᵢRᵢVᵢ\|₀ ≤ 3√ε/𝑡. \)

**Proof.** From Lemma 1, it follows,
\[
σ₁(VᵢRᵢVᵢ) ≥ (1 − √ε/𝑡)², \quad σ₁(VᵢRᵢVᵢ) ≤ (1 + √ε/𝑡)².
\]
Thus, \( λᵢ \max \left( VᵢVᵢ − VᵢRᵢVᵢ \right) \leq \left( 1 − (1 + √ε/𝑡)² \right) ≤ 2√ε/𝑡. \)

Similarly, \( λᵢ \min \left( VᵢVᵢ − VᵢRᵢVᵢ \right) \geq \left( 1 + (1 − √ε/𝑡)² \right) ≥ 3√ε/𝑡. \)
Combining these two results, we have \( \|VVᵢVᵢ − VᵢRᵢVᵢ\|₁ ≤ 3√ε/𝑡. \)

Note: Let \( e = 3√ε/𝑡. \) It is possible to set an upper bound on \( e \) by setting the value of \( r. \) We will assume \( e ∈ (0, 1/2). \)

4.1.1. **Leverage-score sampling**

Our randomized feature selection method is based on importance sampling or the so-called leverage-score sampling [17]. Let \( V ∈ R⁺ⁿᵈ \) be the matrix containing the top-\( ε \) right singular vectors of the training set \( X, \) where \( ε \) is the rank of \( X. \) Compute the normalized leverage scores \( pᵢ \) as follows,
\[
pᵢ = \frac{\|vᵢ\|²}{δ}, \quad \text{for } i = 1, 2, ..., d.
\]
Fix a sampling parameter \( r = O(\sqrt{\log(1/(δ, 2)})). \) Then in r random i.i.d. trials, keep the i-th feature with probability \( pᵢ \) and multiply it by the factor \( 1/√pᵢ. \) The time complexity is dominated by the time to compute the SVD of \( X. \) The i-th leverage-score characterizes the importance of the i-th feature. These scores form a probability distribution over the columns of \( X. \) since \( \sumᵢ pᵢ = 1. \) We include the algorithm as Algorithm 2.

**Lemma 3** ([25]). Let \( ε ∈ (0, 1/2) \) be an accuracy parameter. Given \( V ∈ R⁺ⁿᵈ \) satisfying \( VVᵀ = Iᵩ, \) let \( pᵢ \) be as Eq. (4) and let \( r = O(\sqrt{\log(1/(δ, 2)})). \) Then with probability at least 0.99, \( \|VVᵢVᵢ − VᵢRᵢVᵢ\|₁ ≤ ε. \)

5. **Theoretical analysis**

Our feature selection algorithms are relatively simple and we show that running the linear SVM in the feature space results in a classifier with provably comparable margin to the SVM classifier obtained from the full feature space. Our main results are in Theorems 1 and 3. We state the theorems for BSS, but similar theorems can be stated for leverage-score sampling. Since the feature selection algorithm using leverage-score sampling is a randomized algorithm, the theorems related to margin and geometry preservation with leverage-score sampling hold with certain probability. BSS is a deterministic feature selection algorithm, hence there is no failure probability involved in the theorems.

5.1. **Margin is preserved by supervised feature selection**

Theorem 1 says that you get comparable margin from solving the SVM on the support vectors (equivalently all the data) and from solving the SVM on support vectors in a feature space with only \( O(\text{support vectors}) \) features.

**Theorem 1.** Given \( ε ∈ (0, 1/2), \) perform supervised BSS-feature selection on \( Xᵩ \) with \( nᵩ = O(1/ε²), \) to obtain the feature sampling and rescaling matrix \( R. \) Let \( γᵢ \) and \( γᵢ² \) be the margins obtained by solving the SVM dual (2) with \( (Xᵩ, Yᵩ) \) and \( (XᵩR, Yᵩ) \) respectively. Then, \( γᵢ² ≥ (1 − ε)γᵢ². \)

**Proof.** Let \( Xᵩ ∈ R⁺ⁿᵈ, Yᵩ ∈ R⁺ⁿ \) be the feature matrix and class labels of the training set (as defined in Section 2) and let
\[ \alpha^* = [\alpha_1^*, \alpha_2^*, \ldots, \alpha_p^*] \in \mathbb{R}^p \] be the vector achieving the optimal solution for the problem of Eq. (2). Then,

\[ Z_{\text{opt}} = \sum_{j=1}^{n} \alpha_j - \frac{1}{2} \alpha^{\text{T}} Y^\Sigma Y^{\text{T}} \alpha^* \]  

(5)

Let \( p \leq n \) be the support vectors with \( \alpha_j > 0 \). Let \( \alpha^* = [\alpha_1^*, \alpha_2^*, \ldots, \alpha_p^*] \in \mathbb{R}^p \) be the vector achieving the optimal solution for the problem of Eq. (5). Let \( X^w \in \mathbb{R}^{d \times d} \), \( Y^w \in \mathbb{R}^{p \times p} \) be the support vector matrix and the corresponding labels respectively. Let \( X^w = \mathbb{U} \Sigma \mathbb{V} \), and \( E = \mathbb{V} \Sigma \mathbb{V}^\text{T} - \mathbb{V} \mathbb{R} \mathbb{R}^\text{T} \mathbb{V} \). Then, we can write Eq. (5) in terms of support vectors as,

\[ Z_{\text{opt}} = \sum_{i=1}^{p} \alpha_i^* - \frac{1}{2} \alpha^{\text{T}} \Sigma \Sigma \mathbb{V} \mathbb{V}^\text{T} \Sigma \Sigma \mathbb{V} \mathbb{V}^\text{T} \alpha^* \]

\[ = \sum_{i=1}^{p} \alpha_i^* - \frac{1}{2} \alpha^{\text{T}} \Sigma \Sigma \mathbb{V} \mathbb{V}^\text{T} \Sigma \Sigma \mathbb{V} \mathbb{V}^\text{T} \alpha^* \]

\[ \geq \sum_{i=1}^{p} \alpha_i^* - \frac{1}{2} \alpha^{\text{T}} \Sigma \Sigma \mathbb{V} \mathbb{V}^\text{T} \Sigma \Sigma \mathbb{V} \mathbb{V}^\text{T} \alpha^* \]

\[ = \sum_{i=1}^{p} \alpha_i^* - \frac{1}{2} \alpha^{\text{T}} \Sigma \Sigma \mathbb{V} \mathbb{V}^\text{T} \Sigma \Sigma \mathbb{V} \mathbb{V}^\text{T} \alpha^* \]

\[ \geq \sum_{i=1}^{p} \alpha_i^* - \frac{1}{2} \alpha^{\text{T}} \Sigma \Sigma \mathbb{V} \mathbb{V}^\text{T} \Sigma \Sigma \mathbb{V} \mathbb{V}^\text{T} \alpha^* \]

\[ = \sum_{i=1}^{p} \alpha_i^* - \frac{1}{2} \alpha^{\text{T}} \Sigma \Sigma \mathbb{V} \mathbb{V}^\text{T} \Sigma \Sigma \mathbb{V} \mathbb{V}^\text{T} \alpha^* \]

\[ \geq \sum_{i=1}^{p} \alpha_i^* - \frac{1}{2} \alpha^{\text{T}} \Sigma \Sigma \mathbb{V} \mathbb{V}^\text{T} \Sigma \Sigma \mathbb{V} \mathbb{V}^\text{T} \alpha^* \]

\[ Z_{\text{opt}} \geq Z_{\text{opt}} - \frac{1}{2} ||E||_2^2 ||\alpha^{\text{T}} \Sigma \Sigma \mathbb{V} \mathbb{V}^\text{T} \alpha||_2^2 \]

(10)

We now proceed to bound the second term in the right-hand side of the above equation. Towards that end, we bound the difference:

\[ ||\alpha^{\text{T}} \Sigma \Sigma \mathbb{V} \mathbb{V}^\text{T} \alpha||_2^2 \]

\[ \leq \frac{1}{2} ||Q||_2 ||E||_2 ||Q||_2 \]

\[ = \frac{1}{2} ||E||_2 ||Q||_2^2 \]

\[ \leq \frac{1}{2} ||E||_2 ||\alpha^{\text{T}} \Sigma \Sigma \mathbb{V} \mathbb{V}^\text{T} \alpha||_2^2 \]

\[ \leq \frac{1}{2} ||E||_2 ||\alpha^{\text{T}} \Sigma \Sigma \mathbb{V} \mathbb{V}^\text{T} \alpha||_2^2 \]

(9)

Combining Eqs. (8) and (9), we get

\[ Z_{\text{opt}} \geq Z_{\text{opt}} - \frac{1}{2} ||E||_2 ||\alpha^{\text{T}} \Sigma \Sigma \mathbb{V} \mathbb{V}^\text{T} \alpha||_2^2 \]

(10)

We now proceed to bound the second term in the right-hand side of the above equation. Towards that end, we bound the difference:
prove that $\gamma^2 \geq (1 - \epsilon)\gamma^2$. This proof is almost identical to the proof of Theorem 1 (replacing $(X^W, Y^W)$ with $(X^W, Y^R)$), and so we omit it. Second, we prove that $B^2 \leq (1 + \epsilon)B^2$. We give the result (with proof) as Theorem 5. The theorem follows by combining these two results.

We now state a similar theorem for leverage-score sampling.

**Theorem 4.** Given $\epsilon \in (0, 1/2]$, perform unsupervised leverage-score based feature selection on the full data $X^W$ with $t_2 = O\left(\frac{\epsilon}{\epsilon^2}\right)$, where $\rho = \text{rank}(X^W)$, to obtain the feature sampling and rescaling matrix $R$. Let $\gamma^W$ and $\gamma^R$ be the margins obtained by solving the SVM dual (2) with $(X^W, Y^W)$ and $(X^W, Y^R)$ respectively; and, let $B$ and $\tilde{B}$ be the radii for the data matrices $X^W$ and $X^W R$ respectively. Then with probability at least 0.99,

$$B^2 \leq (1 + \epsilon)B^2.$$

**5.3. Proof that the data radius is preserved by unsupervised BSS-feature selection**

**Theorem 5.** Let $t_2 = O\left(\frac{n_{\text{fl}}}{\epsilon^2}\right)$, where $\epsilon > 0$ is an accuracy parameter, $n$ is the number of training points and $t_2$ is the number of features selected. Let $B$ be the radius of the minimum ball enclosing all points in the full-dimensional space, and let $\tilde{B}$ be the radius of the ball enclosing all points in the sampled subspace obtained by using BSS in an unsupervised manner. For $R$ as in Lemma 2, $B^2 \leq (1 + \epsilon)B^2$.

**Proof.** We consider the matrix $X^W \in \mathbb{R}^{n \times d}$ whose first $n$ rows are the rows of $X^W$ and whose last row is the vector $x_n^W$; here $x_n$ denotes the center of the minimum radius ball enclosing all $n$ points. Then, the SV of $X_n$ is equal to $x_n = U_n \Sigma_n V_n$, where $U_n \in \mathbb{R}^{n \times \min(n, d)}$, $\Sigma_n \in \mathbb{R}^{\min(n, d) \times \min(n, d)}$, and $V \in \mathbb{R}^{d \times \min(n, d)}$. Here $\rho_n$ is the rank of the matrix $X_n$ and clearly $\rho_n \leq n + 1$. (Recall that $\rho$ is the rank of the matrix $X^W$.) Let $B$ be the radius of the minimal radius ball enclosing all points in the original space. Then, for any $i = 1, \ldots, n$,

$$B^2 \geq \|x_i - x_n^W\|^2 = \|e_i - e_n^W\|^2 \|x_i\|^2. \quad (14)$$

Now consider the matrix $X_n R$ and notice that

$$\left\|e_i - e_{n+1}\right\|^2 \|x_i\|^2 = \left\|e_i - e_{n+1}\right\|^2 \|x_i\|^2 \|X_n R\|^2 = \left\|e_i - e_{n+1}\right\| \|x_i - x_n^W R\|^2 \|e_i - e_{n+1}\| \|x_i\|^2 \|X_n R\|^2$$

$$\leq \|x_i\| \|e_i - e_{n+1}\| \|U_n \Sigma_n V_n\|^2 \|e_i - e_{n+1}\| \|x_i\| \|X_n R\|^2 \|e_i - e_{n+1}\| \|x_i\|^2 \|X_n R\|^2$$

$$\leq \|x_i\| \|e_i - e_{n+1}\| \|U_n \Sigma_n V_n\|^2 \|e_i - e_{n+1}\| \|x_i\| \|X_n R\|^2 \|e_i - e_{n+1}\| \|x_i\|^2 \|X_n R\|^2.$$

In the above, we let $E = E \in \mathbb{R}^{d \times d}$ be the matrix that satisfies $V_i V_i^T = V_i V_i^T E$, and we also used $V_i V_i^T = I$. Now consider the ball whose center is the $(n + 1)$-th row of the matrix $X_n R$ (essentially, the center of the minimal radius enclosing ball for the original points in the sampled space). Let $i = \arg \max_{n + 1, \ldots, n} \|e_i - e_{n+1}\|^2 \|x_i\|^2$; then, using the above bound and Eq. (14), we get

$$\left\|e_i - e_{n+1}\right\|^2 \|x_i\|^2 \leq \left(1 + \|E\|\right)\left\|e_i - e_{n+1}\right\|^2 \|x_i\|^2 \leq \left(1 + \|E\|\right)\left\|e_i - e_{n+1}\right\|^2 \|x_i\|^2.$$
we repeat the sampling process five times.

6.1.3. Recursive feature elimination

Recursive Feature Elimination (RFE), [14] tries to find the best subset of features which leads to the largest margin of class separation using SVM. At each iteration, the algorithm greedily removes the feature that decreases the margin the least, until the required number of features remain. At each step, it computes the weight vector and removes the feature with smallest weight. RFE is computationally expensive for high-dimensional datasets. Therefore, at each iteration, multiple features are removed to avoid the computational bottleneck.

6.1.4. LPSVM

The feature selection problem for SVM can be formulated in the form of a linear program. LPSVM [11] uses a fast Newton method to solve this problem and obtains a sparse solution of the weight vector, which is used to select the features.

6.2. BSS implementation issues

At every iteration, there can be multiple columns which satisfy the condition,

\[ \mathcal{U}(\mathbf{v}, \mathbf{d}, \mathbf{A}, \mathbf{U}) \leq \mathcal{L}(\mathbf{v}, \mathbf{d}, \mathbf{A}, \mathbf{L}) \].

Batson et al. [1] suggest picking any column which satisfies this constraint. Selecting a column naively leaves out important features required for classification. Therefore, we choose the column \( \mathbf{v} \) which has not been selected in previous iterations and whose Euclidean norm is highest among the candidate set. Columns with zero Euclidean norm never get selected by the algorithm.

In our implementation, we do not use the data center as one of the inputs (since computing the center involves solving a quadratic program).

6.3. Experiments on supervised feature selection

6.3.1. Synthetic data

We generate synthetic data as described in Bhattacharyya [2], where we control the number of relevant features in the dataset. The dataset has \( n \) data-points and \( d \) features. The class label \( y_i \) of each data-point was randomly chosen to be 1 or -1 with equal probability. The first \( k \) features of each data-point \( x_i \) are the relevant features and are drawn from \( \mathcal{N}(0, 1) \) distribution, where \( \mathcal{N}(\mu, \sigma^2) \) is a normal distribution mean \( \mu \) and variance \( \sigma^2 \) and \( j \) varies from 1 to \( k \). The remaining \( (d - k) \) features are chosen from a \( \mathcal{N}(0, 1) \) distribution and are noisy features. By construction, among the first \( k \) features, the \( k \)th feature has the most discriminatory power, followed by \( (k - 1) \)th feature and so on. We set \( n \) to 200 and \( d \) to 1000. We set \( k \) to 40 and 50 and ran two sets of experiments. We set the value of \( r_j \), i.e. the number of features selected, to 30 and 40 for all experiments. We performed ten-fold cross-validation and repeated it ten times. We used LIBSVM with default settings and set \( C = 1 \). We compared with the other methods. The mean out-of-sample error was 0 for all methods for both \( k = 40 \) and \( k = 50 \). Table 1 shows the set of five most frequently selected features for each method for one such synthetic dataset. The top features picked up by the different methods are the relevant features by construction and also have good discriminatory power. This shows that supervised BSS and leverage-score sampling are as good as any other method in terms of feature selection. We repeated our experiments on ten different synthetic datasets and each time, the five most frequently selected features were from the set of relevant features. Thus, by selecting only 3–4% of all features, we show that we are able to obtain the most discriminatory features along with good out-of-sample error using BSS and leverage-score sampling.

6.3.2. TechTC-300

For our first real dataset, we use 49 datasets of TechTC-300 [8] which contain binary classification tasks. Each data matrix consists of 150–280 documents (the rows of the data matrix), and each document is described with respect to 10,000–50,000 words (features are columns of the matrix). We removed all words with at most four letters from the datasets. We set the parameter \( C = 1 \) in LIBSVM and used default settings. We tried different values of \( C \) for the full-dataset and the out-of-sample error averaged over 49 TechTC-300 documents did not change much, so we report the results of \( C = 1 \). We set the number of features to 300, 400 and 500. Fig. 1 shows the out-of-sample error for the 49 datasets for \( r_1 = 300, 400 \) and 500. For the supervised feature selection, BSS is comparable to RFE and leverage-score sampling and better than RRQR, LPSVM, full-data and uniform sampling in terms of out-of-sample error. For LPSVM, the number of selected features averaged over 49 datasets was greater than 500, but it performed worse than BSS and leverage-score sampling. Leverage-score sampling is comparable to BSS and better than RRQR, LPSVM, full-data and uniform sampling and slightly worse than RFE. The experiments on full-data give worse performance in terms of out-of-sample error than the established and proposed methods. This seems to suggest the full-data with all the features are overfitting the model and by removing irrelevant features, we are reducing overfitting and obtaining better performance.

We list the most frequently occurring words selected by supervised BSS and leverage-score for the \( r_1 = 300 \) case for five TechTC-300 datasets over 100 training sets. Table 2 shows the names of the five TechTC-300 document-term matrices. The words shown in Table 3 were selected in all cross-validation experiments for these five datasets. The words are closely related to the categories to which the documents belong, which shows that BSS and leverage-score sampling select important features from the support-vector matrix. For example, for the document-pair (ii), where the documents belong to the category of “Arts:Music:Styles:Opera” and “US:Navy:Decommissioned Attack Submarines”, the BSS algorithm selects submarine, hullnumber, opera, tickets and leverage-score sampling selects hullnumber, opera, music, tickets which are closely related to the two classes. Thus, we see that using only 2–4% of all features we are able to obtain good out-of-sample error.

### Table 1

<table>
<thead>
<tr>
<th>( r_j = 30 )</th>
<th>( r_j = 40 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( k=40 )</td>
<td>( k=50 )</td>
</tr>
<tr>
<td>BSS</td>
<td>40, 39, 34, 36, 37</td>
</tr>
<tr>
<td>Lvg</td>
<td>40, 39, 37, 36, 34</td>
</tr>
<tr>
<td>RFE</td>
<td>40, 39, 38, 37, 36</td>
</tr>
<tr>
<td>LPSVM</td>
<td>40, 39, 38, 37, 36</td>
</tr>
<tr>
<td>RRQR</td>
<td>40, 30, 29, 28, 27</td>
</tr>
</tbody>
</table>

6.4. Experiments on unsupervised feature selection

For the unsupervised feature selection case, we performed experiments on the same 49 TechTC-300 datasets and set \( r_2 \) to 300, 400 and 500. We include the results for different values of \( r_2 \) in Fig. 2. For LPSVM, the number of selected features averaged over
49 datasets was close to 300. In the unsupervised case, BSS and leverage-score sampling are comparable to each other and also comparable to the other methods RRQR, LPSVM and RFE. These methods are better than random feature selection and full-data without feature selection. This shows that unsupervised BSS and leverage-score sampling are competitive feature selection algorithms. The full-data with all features are overfitting the model and by using feature selection methods, we are removing irrelevant features to reduce overfitting and obtaining better performance. Supervised feature selection is comparable to unsupervised feature selection for BSS, Leverage-score sampling and RFE, while unsupervised RRQR and LPSVM are better than their supervised versions. Running BSS (or leverage-score sampling) on the support-vector set is equivalent to running BSS (or leverage-score sampling) on the training data. However, RRQR and LPSVM are primarily used as unsupervised feature selection techniques and so they perform well in that setting. RFE is a heuristic based on SVM and running RFE on the support-vectors is equivalent to running RFE on the training data.

### Table 2
A subset of the TechTC matrices of our study.

<table>
<thead>
<tr>
<th>id1</th>
<th>id2</th>
</tr>
</thead>
<tbody>
<tr>
<td>(i) Arts: Music: Styles: Opera</td>
<td>Arts: Education: Language: Reading Instructions</td>
</tr>
<tr>
<td>(iii) US: Michigan: Travel &amp; Tourism</td>
<td>Recreation: Sailing Clubs: UK</td>
</tr>
<tr>
<td>(iv) US: Michigan: Travel &amp; Tourism</td>
<td>Science: Chemistry: Analytical: Products</td>
</tr>
<tr>
<td>(v) US: Colorado: Localities: Boulder</td>
<td>Europe: Ireland: Dublin: Localities</td>
</tr>
</tbody>
</table>

### Table 3
Frequently occurring terms of the five TechTC-300 datasets of Table 2 selected by supervised BSS and Leverage-score sampling.

<table>
<thead>
<tr>
<th>BSS</th>
<th>Leverage-score sampling</th>
</tr>
</thead>
<tbody>
<tr>
<td>(i) Reading, education, opera, frame</td>
<td>Reading, opera, frame, spacer</td>
</tr>
<tr>
<td>(ii) Submarine, hullnumber, opera, tickets</td>
<td>Hullnumber, opera, music, tickets</td>
</tr>
<tr>
<td>(iii) Michigan, vacation, yacht, sailing</td>
<td>Sailing, yacht, michigan, vacation</td>
</tr>
<tr>
<td>(iv) Chemical, michigan, environmental, asbestos</td>
<td>Travel, vacation, michigan, environmental</td>
</tr>
<tr>
<td>(v) Ireland, dublin, swords, boulder, colorado</td>
<td>Ireland, boulder, swords, school, grade</td>
</tr>
</tbody>
</table>

Fig. 1. Plots of out-of-sample error (eout) of Supervised BSS and leverage-score compared with other methods for 49 TechTC-300 documents averaged over ten ten-fold cross validation experiments. Vertical bars represent standard deviation.

Fig. 2. Plots of out-of-sample error (eout) of Unsupervised BSS and leverage-score compared with other methods for 49 TechTC-300 documents averaged over ten ten-fold cross validation experiments. Vertical bars represent standard deviation.
6.5. Approximate BSS

We describe a heuristic to make supervised BSS scalable to large-scale datasets. For datasets with large number of support vectors, we premultiply the support vector matrix \( X \) with a random gaussian matrix \( G \in \mathbb{R}^{d \times p} \) to obtain \( \hat{X} = GX \) and then use BSS to select features from the right singular vectors of \( \hat{X} \). The right singular vectors of \( \hat{X} \) closely approximates the right singular vectors of \( X \). Hence the columns selected from \( \hat{X} \) will be approximately same as the columns selected from \( X \). We include the algorithm as Algorithm 3. We performed experiments on a subset of Reuters Corpus dataset, namely reuters-CCAT, which contains binary classification task. We used the L2-regularized L2-loss SVM formulation in the dual form in LIBLINEAR and set the value of \( C \) to 10. We experimented with different values of \( C \) on the full-dataset, and since there was small change in classification accuracy among the different values of \( C \), we chose \( C=10 \) for our experiments. We pre-multiplied the support vector matrix with a random gaussian matrix of size \( t \times p \), where \( p \) is the number of support vectors and \( t \) was set to 128 and 256. We repeated our experiments five times using five different random gaussian matrices to get around the randomness. We set the value of \( r_1 \) in BSS to 1024 and 2048. LPSVM selects 1898 features for CCAT. Table 4 shows the results of our experiments. We observe that the out-of-sample error using approx-BSS is close to that of RRQR and comparable to RFE, LPSVM and full-data. The out-of-sample error of approx-BSS decreases with an increase in the value of \( t \). This shows that we get a good approximation of the right singular vectors of the support vector matrix with an increase in number of projections.

Algorithm 3. Approximate BSS.

**Input:** Support vector matrix \( X \in \mathbb{R}^{d \times p} \), \( t \), \( r \).

**Output:** Matrices \( S \in \mathbb{R}^{d \times r} \), \( D \in \mathbb{R}^{r \times r} \).

1. Generate a random Gaussian matrix, \( G \in \mathbb{R}^{d \times p} \).
2. Compute \( \hat{X} = GX \).
3. Compute right singular vectors \( \text{VoF} \hat{X} \) using SVD.
4. Run Algorithm 1 using \( V \) and \( r \) as inputs and get matrices \( S \) and \( D \) as outputs.
5. Return \( S \) and \( D \).

7. Conclusions

Our simple method of extending an unsupervised feature selection method into a supervised one for SVM not only has a provable guarantee, but also works well empirically: BSS and leverage-score sampling are comparable and often better than prior state-of-the-art feature selection methods for SVM, and those methods do not come with guarantees.

Our supervised sparsification algorithms only preserve the margin for the support vectors in the feature space. We do not make any claims about the margin of the full data in the feature space constructed from the support vectors. This appears challenging and it would be interesting to see progress made in this direction: can one choose 0(\#support vectors) features for the full data set and obtain provable guarantees on the margin and data radius? There have been recent advances in approximate leverage-scores for large-scale datasets. A possible future work in this direction would be to see if those algorithms indeed work well with SVMs.

**Conflict of interest**

None declared.

**Acknowledgments**

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**References**


**Table 4** Results of Approximate BSS. CCAT (train/test): (23,149/781,265), d=47,236. Mean and standard deviation (in parenthesis) of out-of-sample error (eout). Eout of full-data is 8.66±0.54.

<table>
<thead>
<tr>
<th>Eout</th>
<th>( r_1 )</th>
<th>BSS (t=128)</th>
<th>BSS (t=256)</th>
<th>RRQR</th>
<th>RFE</th>
<th>LPSVM</th>
</tr>
</thead>
<tbody>
<tr>
<td>CCAT</td>
<td>1024</td>
<td>10.53 (0.59)</td>
<td>10.35 (0.64)</td>
<td>9.97 (0.62)</td>
<td>8.92 (0.57)</td>
<td>9.97 (0.55)</td>
</tr>
<tr>
<td>CCAT</td>
<td>2048</td>
<td>11.13 (0.66)</td>
<td>10.63 (0.62)</td>
<td>10.04 (0.66)</td>
<td>8.56 (0.54)</td>
<td>9.97 (0.55)</td>
</tr>
</tbody>
</table>


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