FAST LABEL EMBEDDINGS FOR EXTREMELY LARGE OUTPUT SPACES

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ABSTRACT

Many modern multiclass and multilabel problems are characterized by increasingly large output spaces. For these problems, label embeddings have been shown to be a useful primitive that can improve computational and statistical efficiency. In this work we utilize a correspondence between rank constrained estimation and low dimensional label embeddings that uncovers a fast label embedding algorithm which works in both the multiclass and multilabel settings. The result is a randomized algorithm for partial least squares, whose running time is exponentially faster than naive algorithms. We demonstrate our techniques on two large-scale public datasets, from the Large Scale Hierarchical Text Challenge and the Open Directory Project, where we obtain state of the art results.

1 INTRODUCTION

Recent years have witnessed the emergence of many multiclass and multilabel datasets with increasing number of possible labels, such as ImageNet (Deng et al., 2009) and the Large Scale Hierarchical Text Classification (LSHTC) datasets (Kosmopoulos et al., 2010). One could argue that all problems of vision and language in the wild have extremely large output spaces.

When the number of possible outputs is modest, multiclass and multilabel problems can be dealt with directly (via a max or softmax layer) or with a reduction to binary classification. However, when the output space is large, these strategies are too generic and do not fully exploit some of the common properties that these problems exhibit. For example, often the alternatives in the output space have varying degrees of similarity between them so that typical examples from similar classes tend to be closer\(^1\) to each other than from dissimilar classes. More concretely, classifying an image of a Labrador retriever as a golden retriever is a more benign mistake than classifying it as a rowboat. Shouldn’t these problems then be studied as structured prediction problems, where an algorithm can take advantage of the structure? That would be the case if for every problem there was an unequivocal structure (e.g. a hierarchy) that everyone agreed on and that structure was designed with the goal of being beneficial to a classifier. When this is not the case, we can instead let the algorithm uncover a structure that matches its own capabilities.

In this paper we use label embeddings as the underlying structure that can help us tackle problems with large output spaces, also known as extreme classification problems. Label embeddings can offer improved computational efficiency because the embedding dimension is much smaller than the dimension of the output space. If designed carefully and applied judiciously, embeddings can also offer statistical efficiency because the number of parameters can be greatly reduced without increasing, or even reducing, generalization error.

1.1 CONTRIBUTIONS

We provide a statistical motivation for label embedding by demonstrating that the optimal rank-constrained least squares estimator can be constructed from an optimal unconstrained estimator of an embedding of the labels. In other words, embedding can provide beneficial sample complexity reduction even if computational constraints are not binding.

\(^1\) or more confusable, by machines and humans alike.
We start with a set of parameters. Algorithm 1 shows a recipe for performing randomized PCA. The algorithm is insensitive to the very thorough and more formal discussion see Halko et al. (2011).

**Algorithm 1** Randomized PCA

1: function RPCA(k, X ∈ ℝ^{n×d})
2: (p, q) ← (20, 1)
3: Q ← randn(d, k + p)
4: for i ∈ {1, ..., q} do
5: Ψ ← X^{T}XQ
6: Q ← orthogonalize(Ψ)
7: end for
8: F ← (XQ)^{T}(XQ)
9: (V, Σ^2) ← eig(F, k)
10: V ← QV
11: return (V, Σ)
12: end function

We identify a natural object to define label similarity: the expected outer product of the conditional label probabilities. In particular, in conjunction with a low-rank constraint, this indicates two label embeddings are similar when their conditional probabilities are linearly dependent across the dataset. We apply techniques from randomized linear algebra (Halko et al., 2011) to develop an efficient and scalable algorithm for constructing the embeddings, essentially via a novel randomized algorithm for partial least squares (Geladi & Kowalski, 1986). Intuitively, this technique implicitly decomposes the prediction matrix of a model which would be prohibitively expensive to form explicitly. The first step of our algorithm resembles compressed sensing approaches to extreme classification that use random matrices (Hsu et al., 2009). However our subsequent steps tune the embeddings to the data at hand, providing the opportunity for empirical superiority.

2 ALGORITHM DERIVATION

2.1 NOTATION

We denote vectors by lowercase letters x, y etc. and matrices by uppercase letters W, Z etc. The input dimension is denoted by d, the output dimension by c and the embedding dimension by k. For multiclass problems y is a one hot (row) vector (i.e. a vertex of the c−1 unit simplex) while for multilabel problems y is a binary vector (i.e. a vertex of the unit c-cube). For an m × n matrix X ∈ ℝ^{m×n} we use ||X||_F for its Frobenius norm, X^{\dagger} for the pseudoinverse, Π_{X:L} for the projection onto the left singular subspace of X, and X_{1:k} for the matrix resulting by taking the first k columns of X. We use X^{*} to denote a matrix obtained by solving an optimization problem over matrix parameter X. The expectation of a random variable v is denoted by E[v].

2.2 BACKGROUND

In this section we offer an informal discussion of randomized algorithms for approximating the principal components analysis of a data matrix X ∈ ℝ^{n×d} with n examples and d features. For a very thorough and more formal discussion see Halko et al. (2011).

Algorithm 1 shows a recipe for performing randomized PCA. The algorithm is insensitive to the parameters p and q as long as they are large enough (in our experiments we use p = 20 and q = 1). We start with a set of k+p random vectors and use them to probe the range of X^{T}X. Since principal eigenvectors can be thought as “frequent directions” (Liberty [2013]), the range of Ψ will tend to be more aligned with the space spanned by the top eigenvectors of X^{T}X. We compute an orthogonal basis for the range of Ψ and repeat the process q times. This can also be thought as orthogonal (aka subspace) iteration for finding eigenvectors with the caveat that we early stop (i.e., q is small). Once we are done and we have a good approximation for the principal subspace of X^{T}X, we optimize fully over that subspace and back out the solution. The last few steps are cheap because we are only working with a (k+p) × (k+p) matrix and the largest bottleneck is either the computation of Ψ in
Algorithm 2 Rembrandt: Response EMBedding via RANDomized Techniques

1: function REMBRANDT\((k, X \in \mathbb{R}^{n \times d}, Y \in \mathbb{R}^{n \times c})\) \\
2: \( (p, q) \leftarrow (20, 1) \) \hspace{1cm} \triangleright \text{These hyperparameters rarely need adjustment.} \\
3: \( Q \leftarrow \text{randn}(c, k + p) \) \\
4: for \( i \in \{1, \ldots, q\} \) do \hspace{1cm} \triangleright \text{Randomized range finder for } Y^{\top} \Pi_{X,L} Y \\
5: \hspace{1cm} Z \leftarrow \text{arg min}_{Z} ||YQ - XZ||_{F}^{2} \\
6: \hspace{1cm} Q \leftarrow \text{orthogonalize}(Y^{\top}XZ) \\
7: end for \hspace{1cm} \triangleright \text{NB: total of } (q + 1) \text{ data passes, including next line} \\
8: \hspace{1cm} F \leftarrow [Y^{\top}XQ]^{\top} [Y^{\top}XQ] \\
9: \hspace{1cm} (V, \Sigma^{2}) \leftarrow \text{eig}(F, k) \\
10: \hspace{1cm} V \leftarrow QV \\
11: \hspace{1cm} return (V, \Sigma) \\
12: end function

a single machine setting or the orthogonalization step if parallelization is employed. An important observation we use below is that \( X \) or \( X^{\top} X \) need not be available explicitly; to run the algorithm we only need to be able to compute the result of multiplying with \( X^{\top} X \).

2.3 Rank-Constrained Estimation and Embedding

We begin with a setting superficially unrelated to label embedding. Suppose we seek an optimal squared loss predictor of a high-cardinality target vector \( y \in \mathbb{R}^{c} \) which is linear in a high dimensional feature vector \( x \in \mathbb{R}^{d} \). Due to sample complexity concerns, we impose a low-rank constraint on the weight matrix. In matrix form,
\[
W^{*} = \arg \min_{W \in \mathbb{R}^{d \times c} : \text{rank}(W) \leq k} \| Y - XW \|_{F}^{2},
\]
where \( Y \in \mathbb{R}^{n \times c} \) and \( X \in \mathbb{R}^{n \times d} \) are the target and design matrices respectively. This is a special case of a more general problem studied by Friedland & Torokhti (2007), specializing their result yields the solution \( W^{*} = X^{\dagger}(\Pi_{X,L} Y)_{k} \), where \( \Pi_{X,L} \) projects onto the left singular subspace of \( X \), and \((\cdot)_{k}\) denotes optimal Frobenius norm rank-\( k \) approximation, which can be computed via SVD. The expression for \( W^{*} \) can be written in terms of the SVD \( \Pi_{X,L} Y = U \Sigma V^{\top} \), which, after simple algebra, yields \( W^{*} = (X^{\dagger}(YV_{1;k})) V_{1;k}^{\top} \). This is equivalent to the following procedure:

1. \( YV_{1;k} \): Project \( Y \) down to \( k \) dimensions using the top right singular vectors of \( \Pi_{X,L} Y \).
2. \( X^{\dagger}(YV_{1;k}) \): Least squares fit the projected labels using \( X \) and predict them.
3. \( (X^{\dagger}(YV_{1;k})) V_{1;k}^{\top} \): Map predictions to the original output space, using the transpose of the top right singular vectors of \( \Pi_{X,L} Y \).

This motivates the use of the right singular vectors of \( \Pi_{X,L} Y \) as a label embedding. The \( \Pi_{X,L} Y \) term can be demystified: it corresponds to the predictions of the optimal unconstrained model,
\[
Z^{*} = \arg \min_{Z \in \mathbb{R}^{d \times c}} \| Y - XZ \|_{F}^{2},
\]
\( \Pi_{X,L} Y = XZ^{*} \overset{\text{def}}{=} \hat{Y} \).

The right singular vectors \( V \) of \( \Pi_{X,L} Y \) are therefore the eigenvectors of \( \hat{Y}^{\top} \hat{Y} \), i.e., the matrix formed by the sum of outer products of the optimal unconstrained model’s predictions on each example. Note that actually computing and materializing \( Z^{*} \in \mathbb{R}^{d \times c} \) would be expensive; a key aspect of the randomized algorithm is that we get the same result while avoiding this intermediate.

Because squared loss is a proper scoring rule it is minimized at the conditional mean, in the limit of infinite training data \((n \to \infty)\) and sufficient model flexibility (so that \( \hat{y} = \mathbb{E}[y|x] \)) we have that
\[
\frac{1}{n} \hat{Y}^{\top} \hat{Y} \overset{\text{a.s.}}{\to} \mathbb{E}[\mathbb{E}[y|x]^{\top} \mathbb{E}[y|x]]
\]
\(2\) if \( X = U_{X} \Sigma_{X} V_{X}^{\top} \) is the SVD of \( X \), then \( \Pi_{X,L} = U_{X} U_{X}^{\top} \).
by the strong law of large numbers. An embedding based upon the eigendecomposition of \( \mathbb{E}[y|x] \mathbb{E}[^{T}y|x] \) is not practically actionable, but does provide valuable insights. For example, the principal label space transformation of \cite{TaiLin2012} is an eigendecomposition of the empirical label covariance \( Y^\top Y \). This can be a reasonable approximation to \( \mathbb{E}[y|x] \mathbb{E}[^{T}y|x] \) in the multilabel case, as this corresponds to the above procedure when there is no approximation and estimation error. However, for multiclass (or multilabel where most examples have at most one nonzero component), the low-rank constraint alone cannot produce good generalization if the input representation is sufficiently flexible; the eigendecomposition of the prediction covariance will merely select a basis for the \( k \) most frequent labels due to the absence of empirical cooccurrence statistics. Under these conditions we must further regularize (i.e., tradeoff variance for bias) beyond the low-rank constraint, so that \( \hat{Y} \) better approximates \( \mathbb{E}[Y|X] \) rather than the observed \( Y \). Our procedure admits tuning the bias-variance tradeoff via choice of model (features) used in line \( 5 \) of Algorithm \( 2 \).

### 2.4 Rembrandt

Our proposal is Rembrandt, described in Algorithm \( 2 \). In the previous section, we motivated the use of the top right singular space of \( \Pi_{X,Y} \) as a label embedding, or equivalently, the top principal components of \( Y^\top \Pi_{X,Y} \) (leveraging the fact that the projection is idempotent). Using randomized techniques, we can decompose this matrix without explicitly forming it, because we can compute the product of \( \Pi_{X,Y} \) with another matrix \( Q \),

\[
Z^* = \arg \min_{Z \in \mathbb{R}^{d \times (k+p)}} \| YQ - XZ \|^2_F.
\]

(3)

Algorithm \( 2 \) is a specialization of randomized PCA to this particular form of the matrix multiplication operator. Starting from a random label embedding which satisfies the conditions for randomized PCA (e.g., a Gaussian random matrix), the algorithm first fits the embedding, outer products the embedding with the labels, orthogonizes and repeats for some number of iterations. Then a final exact eigendecomposition is used to remove the additional dimensions of the embedding that were added to improve convergence. Note that the optimization here is over \( \mathbb{R}^{d \times (k+p)} \), not \( \mathbb{R}^{d \times c} \), although the result is equivalent; this is the main computational advantage of our technique.

The connection to compressed sensing approaches to extreme classification is now clear, as the random sensing matrix corresponds to the starting point of the iterations in Algorithm \( 2 \). In other words, compressed sensing corresponds to Algorithm \( 2 \) with \( q = 0 \) and \( p = 0 \), which results in a whitened random projection of the labels as the embedding. Additional iterations \( (q > 0) \) and oversampling \( (p > 0) \) improve the approximation of the top eigenspace, hence the potential for improved performance. However when the model is sufficiently flexible, an embedding matrix which ignores the training data might be superior to one which overfits the training data.

Equation (3) is inexpensive to compute. The matrix vector product \( YQ \) is a sparse matrix-vector product so complexity \( O(nsk) \) depends only on the average (label) sparsity per example \( s \) and the embedding dimension \( k \), and is independent of the number of classes \( c \). The fit is done in the embedding space and therefore is independent of the number of classes \( c \), and the outer product with the predicted embedding is again a sparse product with complexity \( O(nsk) \). The orthogonalization step is \( O(k^2) \), but this is amortized over the data set and essentially irrelevant as long as \( n \gg c \). While random projection theory suggests \( k \) should grow logarithmically with \( c \), this is only a mild dependence on the number of classes.

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**Algorithm 3** Loss is either log loss (multiclass) or independent log loss per class (multilabel).

```
1: function DECODERTRAIN(k, X ∈ \mathbb{R}^{n \times d}, Y ∈ \mathbb{R}^{n \times c}, \phi)
2: \( (R, \sim) \leftarrow \text{Rembrandt} (k, X, Y) \)
3: \( W^* \leftarrow \arg \min_{W} \| YR - XD \|^2_F \)
4: \( Y \leftarrow \phi(XW^*) \) \hspace{1cm} \( \triangleright \phi \) is an optional random feature map
5: \( Q^* \leftarrow \arg \min_{Q} \text{loss}(Y, \hat{Y}Q) \) \hspace{1cm} \( \triangleright \) early-stopped, see text
6: return \( (W^*, Q^*) \)
7: end function
```
3 Related Work

Low-dimensional dense embeddings of sparse high-cardinality output spaces have been leveraged extensively in the literature, due to their beneficial impact on multiple algorithmic desiderata. As this work emphasizes, there are potential statistical (i.e., regularization) benefits to label embeddings. Another common motivation is to mitigate space or time complexity at training or inference time. Finally, embeddings can be part of a strategy for zero-shot learning (Palatucci et al., 2009), i.e., designing a classifier which is extensible in the output space.

Hsu et al. (2009), motivated by advances in compressed sensing, utilized a random embedding of the labels along with greedy sparse decoding strategy. For the multilabel case, Tai & Lin (2012) construct a low-dimensional embedding using principal components on the empirical label covariance, which they utilize along with a greedy sparse decoding strategy. In both of these works, the motivation for embeddings was primarily statistical benefit. Conversely, Weston et al. (2011) motivate their ranking-loss optimized embeddings solely by computational considerations of inference time and space complexity.

Multiple authors leverage side information about the classes, such as a taxonomy or graph, in order to learn a label representation which is felicitous for classification, e.g., when composed with online learning (Dekel et al., 2004); Bayesian learning (DeCoro et al., 2007); support vector machines (Bennett & Nguyen, 2009); and decision tree ensembles (Schietgat et al., 2010). Our embedding approach neither requires nor exploits such side information, and is therefore applicable to different scenarios, but is potentially suboptimal when side information is present. However, our embeddings can be complementary to such techniques when side information is not present, as some approaches condense side information into a similarity matrix between classes, e.g., the sub-linear inference approach of Cissé et al. (2012) and the large margin approach of Weinberger & Chapelle (2009). Our embeddings provide a low-rank similarity matrix between classes in factored form, i.e., represented in \( O(ke) \) rather than \( O(c^2) \) space, which can be composed with these techniques. Analogously, Bengio et al. (2010) utilize a surrogate classifier rather than side information to define a similarity matrix between classes; our procedure can efficiently produce a similarity matrix which can ease the computational burden of this portion of their procedure.

Another intriguing use of side information about the classes is to enable zero-shot learning. To this end, several authors have exploited the textual nature of classes in image annotation to learn an embedding over the classes which generalizes to novel classes, e.g., Frome et al. (2013) and Socher et al. (2013). Our embedding technique does not address this problem.

Gopal & Yang (2013) focus nearly exclusively on the statistical benefit of incorporating label structure by overcoming the space and time complexity of large-scale one-against-all classification via distributed training and inference. Specifically, they utilize side information about the classes to regularize a set of one-against-all classifiers towards each other. This leads to state-of-the-art predictive performance, but the resulting model has high space complexity, e.g., terabytes of parameters for the LSHTC (Kaggle, 2014) dataset we utilize in section 4.3. This necessitates distributed learning and distributed inference, the latter being a more serious objection in practice. In contrast, our embedding technique mitigates space complexity and avoids model parallelism.

Our objective in equation (1) corresponds exactly to partial least squares (Geladi & Kowalski, 1986), and an alternative interpretation of Algorithm 2 is a randomized algorithm for PLS. Unsurprisingly, supervised dimensionality reduction techniques such as PLS can be much better than unsupervised dimensionality reduction techniques such as PCA regression in the discriminative setting if the features vary in ways irrelevant to the classification task (Barker & Rayens, 2003).

Two other classical procedures for supervised dimensionality reduction are Fisher Linear Discriminant (Rao, 1948) and Canonical Correlation Analysis (Hotelling, 1936). For multiclass problems these two techniques yield the same result (Bartlett, 1938; Barker & Rayens, 2003), although for multilabel problems they are distinct. Indeed, extension of FLD to the multilabel case is a relatively recent development (Wang et al., 2010) whose straightforward implementation does not appear to be computationally viable for large number of classes. CCA and PLS are highly related, as CCA maximizes latent correlation and PLS maximizes latent covariance (Barker & Rayens, 2003). Furthermore, CCA produces equivalent results to PLS if the features are whitened (Sun et al., 2009). Therefore, there is no statistical reason to prefer CCA to PLS in this context.
Table 1: Data sets used for experimentation and associated times to compute an embedding.

| Dataset  | Type    | Modality | Examples | Features | Classes | Rembrandt k | Time (sec) |
|----------|---------|----------|----------|----------|---------|-------------|------------|
| ALOI     | Multiclass | Vision   | 108K     | 128      | 1000    | 50          | 4          |
| ODP      | Multiclass | Text     | ~1.5M    | ~0.5M    | ~100K   | 300         | 6,530      |
| LSHTC    | Multilabel | Text     | ~2.4M    | ~1.6M    | ~325K   | 500         | 8,006      |

Regarding computational considerations, scalable CCA algorithms are available [Lu & Foster, 2014; Mineiro & Karampatziakis, 2014], but it remains open how to specialize them to this context to leverage the equivalent of equation (3); whereas, if CCA is desired, Algorithm 2 can be utilized in conjunction with whitening pre-processing.

Text is one the common input domains over which large-scale multiclass and multilabel problems are defined. There has been substantial recent work on text embeddings, e.g., word2vec [Mikolov et al., 2013], which (empirically) provide analogous statistical and computational benefits despite being unsupervised. The text embedding technique of Lebret & Collobert (2013) is a particularly interesting comparison because it is a variant of Hellinger PCA which leverages sequential information. This suggests that unsupervised dimensionality reduction approaches can work well when additional structure of the input domain is incorporated, in this case by modeling word burstiness with the square root nonlinearity [Jégou et al., 2009] and word order via decomposing neighborhood statistics. Nonetheless Lebret & Collobert (2013) note that when maximum statistical performance is desired, the embeddings must be fine-tuned to the particular task, i.e., supervised dimensionality reduction is required.

Another plausible regularization technique which mitigates inference space and time complexity is L1 regularization [Lokhorst, 1999]. One reason to prefer low-rank regularization to L1 regularization is if the prediction covariance of equation (2) is well-modeled by a low-rank matrix.

4 Experiments

The goal of these experiments is to demonstrate the computational viability and statistical benefits of the embedding algorithm, not to advocate for a particular classification algorithm per se. Although we utilize classification tasks for demonstration, we will compose our embedding with convenient classification algorithms and focus our attention on the impact of the embedding on the result.

In table 1 we present some statistics about the datasets we use in this section as well as times required to compute an embedding for the dataset. Unless otherwise indicated, all timings presented in the experiments section are for a Matlab implementation running on a standard desktop, which has dual 3.2Ghz Xeon E5-1650 CPU and 48Gb of RAM.

4.1 ALOI

ALOI is a color image collection of one-thousand small objects recorded for scientific purposes [Geusebroek et al., 2005]. The number of classes in this data set does not qualify as extreme by current standards, but we begin with it as it will facilitate comparison with techniques which in our other experiments are intractable on a single machine. For these experiments we will consider test classification accuracy utilizing the same train-test split and features from Choromanska & Langford (2014). Specifically there is a fixed train-test split of 90:10 for all experiments and the representation is linear in 128 raw pixel values.

Algorithm 2 produces an embedding matrix whose transpose is a squared-loss optimal decoder. In practice, optimizing the decode matrix for logistic loss as described in Algorithm 3 gives much better results. This is by far the most computationally demanding step in this experiment, e.g., it takes 4 seconds to compute the embedding but 300 seconds to perform the logistic regression. Fortunately the number of features (i.e., embedding dimensionality) for this logistic regression is modest so the second order techniques of Agarwal et al. (2014) are applicable (in particular, their Algorithm 1 with a simple modification to include acceleration [Nesterov, 1983; Beck & Teboulle, 2009]). We determine the number of fit iterations for the logistic regression by extracting a hold-out set from
the training set and monitoring held-out loss. We do not use a random feature map, i.e., $\phi$ in line 4 of Algorithm 3 is the identity function.

Table 2: ALOI results. $k = 50$ for all embedding strategies.

| Method       | RE + LR | PCA + LR | CS + LR | LR   | OAA  | LT   |
|--------------|---------|----------|---------|------|------|------|
| Test Error   | 9.7%    | 9.7%     | 10.8%   | 10.8%| 11.5%| 16.5%|

We compare to several different strategies in table 2. OAA is the one-against-all reduction of multiclass to binary. LR is a standard logistic regression, i.e., learning directly from the original features. Both of these options are intractable on a single machine for our other data sets. We also compare against Lomtree (LT), which has training and test time complexity logarithmic in the number of classes (Choromanska & Langford, 2014). Both OAA and LT are provided by the Vowpal Wabbit (Langford, 2007) machine learning tool.

The remaining techniques are variants of Algorithm 3 using different embedding strategies. PCA + LR refers to logistic regression after first projecting the features onto their top principal components. CS + LR refers to logistic regression on a label embedding which is a random Gaussian matrix suitable for compressed sensing. Finally RE + LR is Rembrandt composed with logistic regression. These techniques were all implemented in Matlab.

Interestingly, OAA underperforms the full logistic regression. Rembrandt combined with logistic regression outperforms logistic regression, suggesting a beneficial effect from low-rank regularization. Compressed sensing is able to match the performance of the full logistic regression while being computationally more tractable, but underperforms Rembrandt. Lomtree has the worst prediction performance but the lowest computational overhead when the number of classes is large.

At $k = 50$, there is no difference in quality between using the Rembrandt (label) embedding and the PCA (feature) embedding. This is not surprising considering the effective rank of the covariance matrix of ALOI is 70. For small embedding dimensionalities, however, PCA underperforms Rembrandt as indicated in Figure 1a. For larger numbers of output classes, where the embedding dimension will be a small fraction of the number of classes by computational necessity, we anticipate PCA regression will not be competitive.

Note that, in addition to better statistical performance, all of the “embedding + LR” approaches have lower space complexity $O(k(c+d))$ than direct logistic regression $O(cd)$. For ALOI the savings are modest (255600 bytes vs. 516000 bytes) because the input dimensionality is only $d = 128$, but for larger problems the space savings are necessary for feasible implementation on a single commodity computer. Inference time on ALOI is identical for embedding and direct approaches in practice (both achieving $\approx 170k$ examples/sec).
Table 3: ODP results. \( k = 300 \) for all embedding strategies.

| Method   | RE + LR | CS + LR | PCA + LR | LT     |
|----------|---------|---------|----------|--------|
| Test Error | 86.91%  | 88.01%  | 90.37%   | 93.46% |

4.2 ODP

The Open Directory Project \[\text{DMOZ} \ (2014)\] is a public human-edited directory of the web which was processed by \[\text{Bennett \\& Nguyen} \ (2009)\] into a multiclass data set. For these experiments we will consider test classification error utilizing the same train-test split, features, and labels from \[\text{Choromanska \\& Langford} \ (2014)\]. Specifically there is a fixed train-test split of 2:1 for all experiments, the representation of document is a bag of words, and the unique class assignment for each document is the most specific category associated with the document.

The procedures are the same as in the previous experiment, except that we do not compare to OAA or full logistic regression due to intractability on a single machine.

The combination of Rembrandt and logistic regression result is, to the best of our knowledge, the best published result on this dataset. PCA logistic regression has a performance gap compared to Rembrandt and logistic regression. The poor performance of PCA logistic regression is doubly unsurprising, both for general reasons previously discussed, and due to the fact that covariance matrices of text data typically have a long plateau of weak spectral decay. In other words, for text problems projection dimensions quickly become nearly equivalent in terms of input reconstruction error, and common words and word combinations are not discriminative. In contrast, Rembrandt leverages the spectral properties of the prediction covariance of equation (2), rather than the spectral properties of the input features.

Finally, we remark the following: although inference (i.e., finding the maximum output) is linear in the number of classes, the constant factors are favorable due to modern vectorized processors, and therefore proceeds at \( \approx 1700 \) examples/sec for the embedding based approaches.

4.3 LSHTC

The Large Scale Hierarchical Text Classification Challenge (version 4) was a public competition involving multilabel classification of documents into approximately 300,000 categories \[\text{Kaggle} \ (2014)\]. The training and test files are available from the Kaggle platform. The features are bag of words representations of each document.

4.3.1 Embedding Quality Assessment

| Method | Most fraternal | CS | PLST | Rembrandt |
|--------|----------------|----|------|-----------|
| Sibling Fraction | 0.32% | 3.08% | 19.65% | 23.61% |

A representation of a DAG hierarchy associated with the classes is also available. We used this to assess the quality of various embedding strategies independent of classification performance. In particular, we computed the fraction of class embeddings whose nearest neighbor was also a sibling in the DAG, as shown in Table[1]. “Most fraternal” refers to an embedding which arranges for every category’s nearest neighbor in the embedding to be the node in the DAG with the most siblings, i.e., the constant predictor baseline for this task. PLST \[\text{Tai \\& Lin} \ (2012)\] has performance close to Rembrandt according to this metric, so the 3.2 average nonzero classes per example is apparently enough for the approximation underlying PLST to be reasonable.

4.3.2 Empirical Label Covariance Spectrum

Our embedding approach is based upon a low-rank assumption for the (unobservable) prediction covariance of equation [4]. Because LSHTC is a multi-label dataset, we can use the empirical label covariance as a proxy to investigate the spectral properties of the prediction covariance and test our
assumption. We used Algorithm 1 (i.e., two pass randomized PCA) to estimate the spectrum of the empirical label covariance, shown in Figure 1b. The spectrum decays modestly and suggests that an embedding dimension of $k \approx 1000$ or more might be necessary for good classification performance.

4.3.3 Classification Performance

We built an end-to-end classifier using an approximate kernelized variant of Algorithm 3 where we processed the embeddings with Random Fourier Features (Rahimi & Recht, 2007), i.e., in line 4 of Algorithm 3 we use a random cosine feature map for $\phi$. We found Cauchy distributed random vectors, corresponding to the Laplacian kernel, gave good results. We used 4,000 random features and tuned the kernel bandwidth via cross-validation on the training set.

The LSHTC competition metric is macro-averaged F1, which emphasizes performance on rare classes. However, we are using a multilabel classification algorithm which maximizes accuracy of predictions without regard to the importance of rare classes. Therefore we compare with published results of Prabhu & Varma (2014), who report example-averaged precision-at-$k$ on the label ordering induced for each example. To facilitate comparison we do a 75:25 train-test split of the public training set, which is the same proportions as in their experiments (albeit a different split).

Based upon the previous spectral analysis, we anticipate a large embedding dimension is required for best results. With our current implementation, up to the limit of available memory in our desktop machine ($k = 800$) we found increasing embedding dimensionality improved performance.

“RE ($k = \ldots$) + ILR” corresponds for Rembrandt coupled with independent (kernel) logistic regression, i.e., Algorithm 3. LPSR-NB is the Label Partitioning by Sub-linear Ranking algorithm of Weston et al. (2013) composed with a Naive Bayes base learner, as reported in Prabhu & Varma (2014), where they also introduce and report precision for the multilabel tree learning algorithm FastXML. Inference for our best model proceeds at $\approx 60$ examples/sec, substantially slower than for ODP, due to the larger output space, larger embedding dimensionality, and the use of random Fourier features.

5 Discussion

In this paper we identify a correspondence between rank constrained regression and label embedding, and we exploit that correspondence along with randomized matrix decomposition techniques to develop a fast label embedding algorithm.

To facilitate analysis and implementation, we focused on linear prediction, which is equivalent to a simple neural network architecture with a single linear hidden layer bottleneck. Because linear predictors perform well for text classification, we obtained excellent experimental results, but more sophistication is required for tasks where deep architectures are state-of-the-art. Although the analysis presented herein would not strictly be applicable, it is plausible that replacing line 5 in Algorithm 2 with an optimization over a deep architecture could yield good embeddings. This would be computationally beneficial as reducing the number of outputs (i.e., predicting embeddings rather than labels) would mitigate space constraints for GPU training.

Our technique leverages the (putative) low-rank structure of the prediction covariance of equation 2. For some problems a low-rank plus sparse assumption might be more appropriate. In such cases combining our technique with L1 regularization, e.g., on a classification residual or on separately regularized direct connections from the original inputs, might yield superior results.

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