Sparse multidimensional scaling

using landmark points

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Abstract: In this paper, we discuss a computationally efficient approximation to the classical multidimensional scaling (MDS) algorithm, called Landmark MDS (LMDS), for use when the number of data points is very large. The first step of the algorithm is to run classical MDS to embed a chosen subset of the data, referred to as the ‘landmark points’, in a low-dimensional space. Each remaining data point can be located within this space given knowledge of its distances to the landmark points. We give an elementary and explicit theoretical analysis of this procedure, and demonstrate with examples that LMDS is effective in practical use.

Keywords: visualization, embedding, online algorithms, feature discovery, unsupervised learning

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1 Introduction

Metric-preserving dimensionality reduction has long been an important basic task in data analysis and machine learning. Given suitable metric information (such as similarity or dissimilarity measures) about a collection of $N$ objects, the task is to embed the objects as points in a low-dimensional Euclidean space $\mathbb{R}^k$, while preserving the geometry as faithfully as possible. Low-dimensional metric-preserving embeddings have seen numerous applications – such as visualization, feature construction and selection, compression, and data interpolation – in disciplines ranging across the natural and social sciences. For tutorials and reviews see Shepard [1], Cox and Cox [2], Kruskal and Wish [3], or Borg and Lingoes [4].

The original and still best-known approach to this problem is known as classical multidimensional scaling (MDS). Classical MDS has many appealing features. Its domain of natural applicability can be defined precisely: when the given metric on the input data points truly has a low-dimensional Euclidean structure, classical MDS is guaranteed to find a Euclidean embedding which exactly preserves that metric. It is based on an efficient (polynomial time) matrix algorithm that finds the global optimum of a sensible cost function in closed form. The complexity is approximately $O(kN^2)$, where $N$ is the number of data points and $k$ is the dimension of the embedding.

Unfortunately, when the number of points is very large, this algorithm may be too expensive in practice. This limitation has become more pressing recently, with the increasing availability of very large data sets and the corresponding increasing need for scalable dimensionality reduction algorithms. The bottleneck in classical MDS is the calculation of the top $k$ eigenvalues and eigenvectors of an $N \times N$ matrix derived from the input distance matrix $D$. If the number of points is very large compared to the intrinsic dimensionality—typically, the situation where dimensionality
reduction is called for—there should be a better approach than computing the eigendecomposition of this full matrix.

In this paper we describe a variation on classical MDS which preserves all of the attractive properties but is much more efficient, the cost being essentially linear in the number of data points (Section 2.5). Yet it also achieves the correct solution if the data really have a low-dimensional Euclidean structure; and the algorithm is stable under small perturbations of the input distances so good results are obtained even for noisy data. The required input is an \( n \times N \) submatrix of \( D \), giving the distances between the \( N \) data points and a set of \( n \) distinguished points referred to as “landmarks”. There are two steps; first we apply classical MDS to the landmark points. The second step is a distance-based triangulation procedure, which uses distances to the already-embedded landmark points to determine where the remaining points should go. Our contribution is this two-stage decomposition, and a formulation of the triangulation procedure as a linear problem. Together, these two steps give our procedure for Landmark MDS. However, our analysis of the second step—the linear solution to distance-based triangulation—may be of general interest in itself, because that is a problem we may often wish to solve (for example, in global positioning systems).

Landmark MDS is effectively online with respect to the introduction of new data points. After the landmark points are fixed and the initial calculation carried out, all other points are embedded independently from each other using a fixed linear transformation. A global calculation is necessary only if the embedding coordinates are required to be aligned with the principal axes of the data. There seems to be no reason to do this repeatedly, rather than once or twice as called for.

The Landmark MDS algorithm was introduced by the authors in [5], in the context of finding an efficient approximation to the Isomap algorithm for nonlinear dimensionality reduction suitable for processing large data sets. Related ideas have appeared in the machine learning and pattern
recognition literatures in the contexts of various applications [6–8]. Yet MDS is still often dismissed as too computationally demanding for real-time use with large data sets [9–13]. Our goal in this paper is to make Landmark MDS accessible and appealing to a broad audience, by giving a more thorough and more general treatment than has previously been available, including an extensive theoretical analysis, an investigation of the effects of noise, discussion and explanation of potential pitfalls, and pointers to interesting applications. We also present a number of experiments on synthetic and real data to illustrate and verify our analytical results. The remaining sections are organised as follows. In Section 2 we describe the algorithm and its complexity. In Section 3 we run LMDS on two classes of example data sets. The theoretical analysis is in Section 4, and further applications are discussed briefly in Section 5.

2 Landmark MDS

We are given a collection of \( N \) objects which we wish to embed in the Euclidean space \( \mathbb{R}^k \). Their dissimilarities are represented by an \( N \times N \) matrix \( D_N \). Landmark MDS consists of four steps:

1. Designate a set of \( n \) landmark points.

2. Apply classical MDS to find a \( k \times n \) matrix \( L \) representing an embedding of the \( n \) landmark points in \( \mathbb{R}^k \). As input, use the \( n \times n \) matrix \( D_n \) of distances between pairs of landmark points.

3. Apply distance-based triangulation (described later) to find a \( k \times N \) matrix \( X \) representing an embedding of the \( N \) data points in \( \mathbb{R}^k \). As input, use the \( n \times N \) matrix \( D_{n,N} \) of distances between landmark points and data points. The new coordinates are derived from the squared distances by an affine linear transformation.
4. Recenter the data about their mean, and use PCA to align the principal axes of the newly-embedded data with the coordinate axes, in order of decreasing significance.

Pseudocode for the algorithm is given in Figure 1. Before discussing each step in detail, we make some brief remarks:

- The embedding obtained in Step 3 is consistent with the embedding of the landmark points in Step 2. In other words, \( L \) agrees with the corresponding \( k \times n \) submatrix of \( X \).

- The landmarks may be chosen by any reasonable method. For a \( k \)-dimensional embedding, we require at least \( k + 1 \) landmarks; specifically the affine span of the landmarks (as embedded in Step 2) must be \( k \)-dimensional. For stability reasons it is better to avoid configurations which lie close to some \((k - 1)\)- or lower-dimensional affine subspace. It is generally advisable to choose rather more landmark points than the strict minimum.

- Landmark MDS use only the \( n \times N \) matrix \( D_{n,N} \), and not the full \( N \times N \) matrix \( D_N \). If there is a substantial cost associated with calculating or storing the full matrix, this kind of sparsification can lead to a significant savings. As we discuss later, Landmark MDS substantially improves the performance of Isomap [14] and allows us to embed a sparse similarity graph efficiently.

- The final PCA stage is not essential, but it normalises the output. In addition, if \( k' < k \) then a good embedding in \( \mathbb{R}^{k'} \) is obtained by restricting to the first \( k' \) coordinates of the PCA-aligned embedding. It is better to use a higher value of \( k \) and restrict to \( k' \)-dimensions at the end, than to use a low value of \( k \) throughout.

2.1 Step 1: Selecting the landmark points

We suggest two ways of selecting the landmark set:
Input:
\( N \) ← number of data points;
\( \Delta \) ← squared-distance function (two arguments);
n ← desired number of landmark points;
k ← desired number of output dimensions;

Step 1: select landmark points (randomly).
P(1:N) ← randperm(N); // random permutation
\( \ell(\cdot) \) ← P(1:n);

Step 2: classical MDS on landmarks.
for \( (i = 1:n, j = 1:n) \)
\( \Delta_n(i,j) \) ← \( \Delta(\ell(i), \ell(j)) \);
for \( i = (1:n) \)
\( \bar{\mu}_n(i) \) ← mean(\( \Delta_n(i,:))\);
\( \mu \) ← mean(\( \bar{\mu}_n(\cdot) \));
for \( (i = 1:n, j = 1:n) \)
\( B_n(i,j) \) ← \(-\frac{1}{2} (\Delta_n(i,j) - \bar{\mu}_n(i) - \bar{\mu}_n(j) + \mu)\);
[\( V, \lambda \)] ← eigensolve(\( B_n, k \));
// \( V(i,j) \) = i-th component of j-th eigenvector
// \( \lambda(j) \) = j-th eigenvalue, in descending order
\( k_+ \) ← min(k, number of positive eigenvalues);
for \( (i = 1:k_+, j = 1:n) \)
\( L(i,j) \) ← \( V(j,i) \) * sqrt(\( \lambda(i) \));

Step 3: distance-based triangulation.
for \( (i = 1:k_+, j = 1:n) \)
\( L^\#(i,j) \) ← \( V(j,i) / sqrt(\lambda(i)) \);
for \( j = (1:N) \)
\( X(:,j) \) ← \(-L^\#(\cdot,j) * (\Delta(\ell(\cdot),j) - \bar{\mu}_n(\cdot)) / 2\);
// matrix-vector multiplication

Step 4: PCA normalisation (optional).
for \( (i = 1:k_+) \)
\( \bar{X}(i) = \text{mean}(X(i,:)) \);
for \( (j = 1:N) \)
\( X(:,j) \) ← \( X(:,j) - \bar{X}(:,j) \);
[\( U, \kappa \)] ← eigensolve(\( X \ast X^T, k_+ \)); // matrix multiplication
\( X_{\text{pca}} \) ← \( U \ast \bar{X} \); // matrix multiplication

Output:
\( k_+ \) → actual number of output dimensions
\( L \) → \( k_+ \)-dimensional embedding of landmarks
\( X \) → \( k_+ \)-dimensional embedding of data
\( X_{\text{pca}} \) → PCA-normalised \( k_+ \)-dimensional embedding of data

Figure 1: Pseudocode for LMDS.
Input:

\[ N \leftarrow \text{number of data points}; \]
\[ \Delta \leftarrow \text{squared-distance function (two arguments)}; \]
\[ n \leftarrow \text{desired number of landmark points}; \]
\[ s \leftarrow \text{desired number of seed points}; \]

Step 1a: select landmark points by MaxMin.

// choose \( s \) seed points randomly
\[ P(1:N) \leftarrow \text{randperm}(N); \]
\[ \ell(1:s) \leftarrow P(1:s); \]

for \( j = 1:N \)
\[ m(j) \leftarrow \min_{i = 1:s}(\Delta_n(\ell(i), j)); \]

for \( i = s+1:n \)
\[ \ell(i) \leftarrow \text{argmax}_{j = 1:N}(m(j)); \]
\[ m(j) \leftarrow \min(m(j), \Delta_n(\ell(i), j)); \]

Output:

\( \ell \rightarrow \text{list of indices of } n \text{ landmark points} \)

Figure 2: Pseudocode for MaxMin landmark selection, as an alternative to Step 1 of Figure 1.

- Random choice.

- MaxMin (greedy optimisation): landmark points are chosen one at a time, and each new landmark maximises, over all unused data points, the minimum distance to any of the existing landmarks. The first point is chosen arbitrarily.

MaxMin is deterministic after the first landmark point. Sometimes it may be desirable to start MaxMin with \( s \) ‘seed’ points, chosen randomly, for \( s \geq 1 \). At any stage the only information required is the matrix of distances between the existing landmark points and the remaining points, so these entries can be accessed/computed on a need-to-know basis. The cost of using MaxMin instead of random choice amounts to \( O(nN) \) extra operations. See Figure 2.

Random choice works quite well in practice, and we use it for all the examples in this paper. MaxMin has the advantage of being controllable, which may be useful in some contexts where
reproducible results are desired. There is further discussion in Section 4.4.

Remark. At first glance, the choice of landmark points may seem to be a clustering problem. The landmarks can be regarded as a set of \( n \) cluster centers whose distribution should be in general agreement with the distribution of the full data set of \( N \) points. On the other hand, it is expensive to solve an \( n \)-center clustering problem on \( N \) points, and it may require access to the entire distance matrix rather than just a selected set of rows. This is not in the spirit of LMDS, which is meant to be a quick approximation. For this reason we do not endorse the use of full-blooded clustering algorithms. Random choice and MaxMin are cheap and effective.

2.2 Step 2: Classical MDS on landmarks

We recall the procedure, first used in this context by Torgerson [15].

- **Input:** the \( n \times n \) matrix \( \Delta_n \) of squared distances; so \([\Delta_n]_{ij} = [D_n]_{ij}^2\).

- **Construct the mean-centered “inner-product” matrix**

\[
B_n = -\frac{1}{2} H_n \Delta_n H_n
\]  

where \( H_n \) is the mean-centering matrix, defined by \([H_n]_{ij} = \delta_{ij} - 1/n\).

- **Compute** the \( k \) largest positive eigenvalues of \( B_n \) together with an orthonormal set of eigenvectors. Write \( \lambda_i \) for the \( i \)-th largest positive eigenvalue and \( \vec{v}_i \) for the corresponding eigenvector (written as a column vector). Nonpositive eigenvalues are ignored; this includes a zero eigenvalue with eigenvector \( \vec{1} = [1, 1, \ldots, 1]^T \).

- **Output:** the required \( k \)-dimensional embedding vectors \( \vec{\ell}_1, \vec{\ell}_2, \ldots, \vec{\ell}_n \) are given by the columns
of the following\(^1\) matrix:

\[
L_k = \begin{bmatrix}
\sqrt{\lambda_1} \cdot \vec{v}_1^T \\
\sqrt{\lambda_2} \cdot \vec{v}_2^T \\
\vdots \\
\sqrt{\lambda_k} \cdot \vec{v}_k^T
\end{bmatrix}
\]  

(2)

The embedding is automatically mean-centered, so \(\vec{\ell}_1 + \ldots + \vec{\ell}_n = \vec{0}\), or equivalently \(L_k H_n = L_k\). This follows from the orthogonality of eigenvectors, which gives \(\vec{v}_i^T \vec{1} = 0\) for all \(i\).

The justification for this construction lies in the following theorems.

**Theorem 1** (Young and Householder [16]) Suppose \(\Delta_n\) is the Euclidean squared-distance matrix for a collection of data points \(\vec{y}_1, \vec{y}_2, \ldots, \vec{y}_n \in \mathbb{R}^d\). By translating if necessary we can assume that the \(\vec{y}_j\) are mean-centered. In that case the \(i\)-th row of \(L_k\) gives the components of each data point when projected onto the \(i\)-th principal axis of the data (up to an overall \(\pm\) for each row). \(\square\)

The next result is valid for any symmetric distance matrix.

**Theorem 2** (Eckart and Young [17]) The embedding vectors \(\vec{\ell}_i\) have the inner product matrix which best approximates \(B_n\). In other words,

\[
\|L_k^T L_k - B_n\| \leq \|L^T L - B_n\|
\]

for all \(k \times n\) matrices \(L\); this is true in both the Frobenius norm and the \(\ell^2\) operator norm. Equality \(\|L_k^T L_k - B_n\| = 0\) is achieved if and only if \(k \geq p\) and \(B_n\) has no negative eigenvalues. In that case the Euclidean distance matrix of the \(\vec{\ell}_i\) is equal to \(D_n\). \(\square\)

\(^1\)If \(k > p\), then the last \(k - p\) rows are set to zero.
Negative eigenvalues of $B_n$ signify that the original distance matrix $D_n$ is non-Euclidean. In real data, noise in $D_n$ can account for negative eigenvalues of small magnitude. If there are large negative eigenvalues, then no Euclidean embedding can recover $D_n$ even approximately.

Remark. It may seem more natural to minimise (in a suitable norm) the difference between $D_n$ and the distance matrix of the $\vec{\ell}_i$, or the difference between $\Delta_n$ and the squared-distance matrix of the $\vec{\ell}_i$. In fact, both the set of distances and the set of mean-centered inner products uniquely characterise the intrinsic geometry of the data. Zero error in distances is equivalent to zero error in mean-centered inner products. However, because they use different ‘units’, the corresponding optimisation problems give different answers in the case of non-zero error. We use mean-centered inner products since closed-form solutions are unavailable in the other cases.

2.3 Step 3: Distance-based triangulation

Having embedded the landmark points in $\mathbb{R}^k$, we now compute embedding coordinates for the remaining data points based on their distances from the landmark points. The new coordinates of a point $a$ are obtained by an affine linear transformation of the vector $\vec{\delta}_a$ of its squared distances to the landmark points. First we compute the transformation (which does not depend on $a$).

- Let $\vec{\delta}_1, \vec{\delta}_2, \ldots, \vec{\delta}_n$ denote the columns of $\Delta_n$ (so $\vec{\delta}_i$ is the vector of squared distances from the $i$-th landmark to all the landmarks). Compute the mean $\vec{\delta}_\mu = (\vec{\delta}_1 + \vec{\delta}_2 + \cdots + \vec{\delta}_n)/n$.

- Compute the pseudoinverse transpose $L_k^\#$ of $L_k$. This can be constructed directly from the
eigenvalues and eigenvectors of Step 2, by the following explicit formula:

\[
L_k^\sharp = \begin{bmatrix}
\vec{v}_1^T \sqrt{\lambda_1} \\
\vec{v}_2^T \sqrt{\lambda_2} \\
\vdots \\
\vec{v}_k^T \sqrt{\lambda_k}
\end{bmatrix}
\]

Each data point \(a\) is embedded as follows.

- **Input**: the vector \(\vec{\delta}_a\) of squared distances between the point \(a\) and the \(n\) landmark points.

- **Output**: the embedding vector \(\vec{x}_a\), given by the formula:

\[
\vec{x}_a = -\frac{1}{2} L_k^\sharp (\vec{\delta}_a - \vec{\delta}_\mu)
\]

**Theorem 3** In the situation of Theorem 1, suppose \(\vec{y}_a \in \mathbb{R}^d\) is an additional data point, and \(\vec{\delta}_a\) is its squared-distance vector to the original (landmark) points. Then the vector \(\vec{x}_a\) defined by formula (3) gives the components of \(\vec{y}_a\) when projected onto the first \(k\) principal axes of the landmark points. [By ‘first \(k\) principal axes’ we mean the axes corresponding to the first \(k\) principal components of the set of landmarks.]

The proof of this theorem appears in section 4.1. We conclude this section with a consistency result, valid for any symmetric distance matrix.

**Proposition 4** The new embedding is consistent with the old embedding for the landmark points, meaning that

\[-\frac{1}{2} L_k^\sharp (\vec{\delta}_j - \vec{\delta}_\mu) = \vec{\ell}_j\]
for \( j = 1, 2, \ldots, n \).

**Proof.** Note that \( \vec{\delta}_j - \vec{\delta}_\mu \) is the \( j \)-th column of \( \Delta_n H_n \), and \( \vec{\ell}_j \) is the \( j \)-th column of \( L_k \). Note also that \( L_k^\sharp = L_k^\sharp H_n \) since \( \vec{v}_i^T \vec{1} = 0 \) for all \( i \). Thus what we have to prove is \( -\frac{1}{2} L_k^\sharp H_n \Delta_n H_n = L_k \), that is \( L_k^\sharp B_n = L_k \). For each eigenvector \( \vec{v}_i \) of \( B_n \) we have

\[
\frac{\vec{v}_i^T B_n}{\sqrt{\lambda_i}} = \frac{\lambda_i \vec{v}_i^T}{\sqrt{\lambda_i}} = \sqrt{\lambda_i} \cdot \vec{v}_i^T
\]

so the \( i \)-th row of \( L_k^\sharp B_n \) equals the \( i \)-th row of \( L_k \), for every \( i \). This completes the proof. \( \square \)

### 2.4 Step 4: PCA normalisation

Once the full data set has been embedded in \( \mathbb{R}^k \), it may be appropriate to reorient the axes to reflect the overall distribution, rather than the distribution of the much smaller landmark set. If \( X \) denotes the \( k \times N \) matrix of embedding coordinates, then \( \hat{X} = X H_N \) represents the mean-centered coordinates. We solve the symmetric eigenvalue problem for \( \hat{X} \hat{X}^T = X H_N X^T \) by finding a solution to the equation \( (\hat{X} \hat{X}^T) U = UM \), where \( U \) is orthogonal and \( M \) is diagonal with entries listed in decreasing order. The PCA-normalised coordinates are then given by the matrix \( U^T \hat{X} \).

### 2.5 Sparseness and computational complexity

Landmark MDS leads to savings at three possible bottlenecks in the classical MDS algorithm: storage and calculation of the distance matrix, and the eigenvalue problem. We discuss these in order.

- Storing the distance matrix: Classical MDS requires \( O(N^2) \) storage while LMDS requires \( O(nN) \).

  When \( N \) is very large, this may be decisive in itself.
• Accessing the distance matrix: Assuming a cost of $C$ to compute or access each entry of the distance matrix $D (= D_N$ or $D_{n,N}$), classical MDS requires $O(CN^2)$ as compared to $O(CnN)$ for LMDS. It does sometimes happen that $C$ is large enough to cause a bottleneck here, for example when using a complex metric such as tangent distance [18]. For Euclidean distance in $\mathbb{R}^k$, $C = O(k)$. When LMDS is used to augment the Isomap algorithm [14], as discussed in Section 5.1, each row of the distance matrix gives the answer to a single-source shortest paths problem on a neighbourhood graph. Using Dijkstra’s algorithm the cost per row is $O(\delta N \log N)$, where $\delta$ is the degree of the graph; so the amortised cost per entry is $C = O(\delta \log N)$.

• Finding the embedding: the cost of classical MDS is dominated by the bottleneck eigenvalue problem on a full symmetric $N \times N$ matrix, which costs $O(N^3)$ using typical iterative methods and assuming rapid convergence. Given the distance matrix, the dominant costs of running Landmark MDS are as follows: $O(n^3)$ for the $n \times n$ eigenvalue problem in Step 2, and $O(knN)$ for the distance-based triangulation in Step 3. The remaining two steps are dominated by this: Step 1 costs $O(nN)$ if random choice is replaced by the more expensive MaxMin; and the PCA normalisation at Step 4 costs $O(k^2N)$.

In summary, classical MDS requires $O(N^2)$ space and $O(CN^2 + N^3)$ time, while LMDS requires $O(nN)$ space and $O(CnN + knN + n^3)$ time.

Remark. Note the distinction between local sparsity, which is characteristic of the optimisation problems in LLE, Hessian Eigenmaps, and similar; versus global or rectangular sparsity, as used in LMDS and Landmark Isomap. In both cases there is a square matrix whose rows (and corresponding columns) are identified with points in a Euclidean space. In a local problem, non-zero entries occur in the matrix only when the row and column involved correspond to nearby points in the space.
In contrast, classical MDS deals with a global problem involving a full matrix. To sparsify the problem, we choose a small subset of the points which is nonetheless globally representative of the whole data set, and work with the resulting rectangular submatrix. In a different mathematical context, locally sparse matrices are discussed in [19], where a general concept of geometric sparsity is developed.

3 Examples

3.1 Example: noisy grid

To illustrate the noise-sensitivity of Landmark MDS, we consider the example of a regular grid subjected to varying degrees of noise and using different numbers of landmark points (Figure 3). The initial data set consists of 600 data points in a rectangular grid of aspect ratio 1.5. To add noise to the data, the Euclidean distance matrix was multiplied entrywise by independent random variables distributed as \( \exp(N(0, \sigma)) \) and then symmetrized. The noise levels \( \nu \% \) referred to in the figure are defined by \( \nu = 100(\exp(\sigma) - 1) \). This modified distance matrix was then passed to LMDS with varying numbers of landmark points, chosen randomly but consistently across the different noise levels. PCA normalisation was used as a final step. Figure 4 plots the 2-dimensional embeddings output by LMDS using a colour spectrum to indicate the \( x \)-coordinate of the original configuration.

As predicted by Theorem 3, LMDS works perfectly in the absence of noise (left column), right down to 3 landmark points. When there is noise (columns 2–4), more landmarks means better results. Classical MDS itself (top row, equivalent to LMDS with 600 landmarks) is highly robust to noise up to at least the 10% level. With 50 or 10 landmarks, the overall quality of the embeddings is still quite high, though there is noticeable deterioration at finer scales and at the extremities.
Figure 3: Sensitivity to noise. In this example Landmark MDS is applied to a set of 600 points arranged in a grid, varying the number of landmark points and the amount of noise in the distance matrix. With 600 landmarks (top row) LMDS is equivalent to classical MDS. The bottom two rows compare what happens when three landmarks are well distributed (fourth row) versus almost degenerate (fifth row).
In all three cases (600, 50, 10 landmarks) the embedding is accurate at a coarse resolution. We discuss the dependence of fine-resolution noise on the number of landmarks $n$ in Section 4.4.

The remaining two cases show what happens when we use the bare minimum number of landmarks, namely 3, for a 2-dimensional embedding. The situation is degenerate if the three points are collinear, and in that case there is no 2-dimensional embedding at all. We contrast what happens when the three landmarks are *almost* degenerate (fifth row) and when they are well-distributed, or far from degenerate (fourth row).

Perhaps surprisingly for only three landmark points, the results in the well-distributed case are moderately acceptable; at least at the lower noise levels. However, there is a systematic distortion, which we call *foreshortening*, which is most easily observed in the case with 5% noise. The overall shape of the data appears to be a parallelogram rather than a rectangle. This can be ascribed to a uniform diminution of all coordinate values in a diagonal direction corresponding to the second coordinate function of the initial LMDS embedding (before PCA normalisation). We give an explanation of foreshortening in Section 4.4.

In the almost-degenerate case the results are much worse. There is a foreshortening effect, as before, in the perpendicular direction to the principal axis of the three points. However, this is dominated by a *noise amplification* effect in the same direction. Noise amplification is easily explained. When the landmark set is almost degenerate, the second eigenvalue $\lambda_2$ of $B_n$ is close to zero, and hence the $1/\sqrt{\lambda_2}$ factor causes the second row of $L^2$ to be large. If all the distances are true then this causes no problem, but any noise is amplified by this factor in the second component of the initial LMDS embedding.

Figure 4 shows the same data coloured by the $y$-coordinate of the original configuration. Evidently, in the almost-degenerate case at the higher noise levels, we have lost the original $y$-coordinate altogether. With 2% noise there remains some visible correlation. Still, both of the systematic
distortions—foreshortening and noise amplification—can be seen at work.

3.2 Example: digits from the MNIST database

Landmark MDS is particularly useful when dealing with large data sets for which the cost of classical MDS may be rather high. In this section we apply both algorithms to digit sets taken from the MNIST database of handwritten digits [20]. We find that it is considerably cheaper to run LMDS with 200 landmark points than to run classical MDS; but in this case the results turn out to be of comparable quality.

Each digit set in the MNIST database consists of about 6000 instances of a given numeral (‘0’ to ‘9’) scanned in as 28 × 28 pixel grayscale images from actual handwritten ZIP codes. These data sets are natural candidates for dimensionality reduction, since the images themselves are high-dimensional but much of the variation in the data can be explained by a small number of
Figure 5: Comparing classical MDS (left) and Landmark MDS with 200 landmarks (right) on a set of 5923 digits ‘0’ from the MNIST database.

Figure 6: Applying classical MDS (left) and Landmark MDS with 200 landmarks (right) to a set of 5851 digits ‘8’ from the MNIST database.
natural parameters such as left-right slope, height, and loop size. The relationship between these parameters and the images is highly non-linear, which would suggest the use of Isomap, LLE or some other NLDR algorithm. However, the simpler linear methods can sometimes detect one or two salient degrees of variation, and this turns out to be the case here.

For each digit, we adopted the following procedure. Every $28 \times 28$ grayscale image was rasterised to a vector of length 784 of intensity values in the range $0 \leq s \leq 1$. From the approximately 6000 images, 200 were chosen randomly to be landmark points. Using the Euclidean metric in $\mathbb{R}^{784}$, both classical MDS and Landmark MDS (with the chosen 200 landmarks) were applied to obtain a 2-dimensional embedding. A selection of the results is shown in Figures 5, 6 and 7, with classical MDS in the left column compared with Landmark MDS in the right column. Red highlights mark the points in each embedding closest to the vertices of a superimposed regular grid; the images corresponding to these points are displayed below in a grid, as a visual aid.
The ten digits are not equally suitable candidates for classical MDS. Some of the digit sets do exhibit a clear correlation between the first coordinate and any natural variation parameter. For other digits such as ‘1’, there is too much non-linearity to find a consistent correspondence. In the best cases, ‘0’ and ‘8’, there are identifiable features corresponding to each of the first two coordinates. For ‘0’ (Figure 5), the left–right axis corresponds to area and eccentricity of the oval: smaller, more eccentric ovals on the left; and larger, rounder figures on the right. Variation along the other direction corresponds to the angle of the major axis of the ‘0’, running from vertical (at the top) to 45° clockwise (at the bottom). In the case of the ‘8’ data, the left–right axis corresponds to the angle of the major axis of the ‘8’; and the top–bottom direction correlates with ‘weight’: dark, heavily written characters at the top, and lighter characters at the bottom. For both digits, it is evident that Landmark MDS performs comparably well with classical MDS (subject to the caveat below); certainly the same trends can be seen in the LMDS results.

In contrast to ‘0’ and ‘8’, the space of digits ‘1’ seems to be significantly non-linear; the 2-dimensional embedding coordinates produced by classical MDS (Figure 7, left) fail to correlate consistently with discernible features. Indeed, the curved shape of the data distribution gives fair warning of this, and strongly suggests the use of non-linear techniques. Landmark MDS (right) produces very similar output, confirming that its diagnostic value is comparable to classical MDS.

Finally we compare running times for these calculations, under a straightforward MATLAB implementation. To the nearest 0.1 s, classical MDS on the ‘8’ data took 160.6 s to calculate the distance matrix and 57.2 s to calculate the embedding, and a total of 217.9 s. Landmark MDS took 1.9 s to calculate the reduced distance matrix and 0.4 s for the embedding calculation, making 2.2 s in total. There was a similar ‘improvement factor’ of about 100 for each of the ten digits. These results support our claim that Landmark MDS is an effective, computationally efficient substitute for classical MDS in a range of real-world data analysis situations, both linear and non-linear.
Caveat. We have taken care to show typical results for these experiments with 200 randomly-chosen landmarks. For some of the digits where the data space is highly nonlinear, such as ‘8’, it happened once or twice in every ten runs that the two-dimensional embedding given by LMDS differed significantly from classical MDS in the second coordinate. This raises the question of how to choose good landmarks (and how many of them). We discuss this question in the analysis section.

4 Analysis

We now discuss the theoretical performance of LMDS in terms of the following question: If classical MDS is applied to an all-pairs distance matrix $\Delta_N$ and LMDS is applied to a landmark-to-point rectangular submatrix $\Delta_{n,N}$ then are the two embeddings the same? Here ‘the same’ means ‘congruent up to a rigid transformation’.

We answer this in three parts. First, if there is an exact Euclidean embedding preserving $\Delta_N$, we verify that LMDS recovers this configuration exactly provided that all the data points are contained in the affine span of the landmark points. If the landmarks span a lower-dimensional affine subspace, then LMDS recovers the orthogonal projection of the points onto this subspace.

Second, we study the perturbation properties of LMDS. We give first-order perturbation formulas for classical MDS and LMDS and identify the parameters which lead to poor stability. Under reasonable conditions, LMDS and classical MDS both behave well with respect to small perturbations of the input. As a consequence if the distance matrix is approximately Euclidean, then LMDS and classical MDS will give approximately the same embedding.

Third, we give an example of a non-Euclidean data set for which the two algorithms give quite different results. It is therefore wrong to assume that LMDS is always a good approximation to
classical MDS. More precisely, if a data set is not well suited to classical MDS, then Landmark MDS may give unpredictable results. On the other hand, for data sets where the use of classical MDS is appropriate, the preceding results imply that Landmark MDS is a good approximation.

4.1 The Euclidean case

When classical MDS is applied to a distance matrix originating from a set of Euclidean data, the resulting \( k \)-dimensional embedding recovers the projection of the data onto its first \( k \) mean-centered principal axes. This is the content of Theorem 1. Note that if the affine span of the data is \( p \)-dimensional, then there are exactly \( p \) non-zero principal components. Thus the configuration is recovered perfectly up to isometry when \( k = p \); nothing is gained by taking \( k > p \).

Similarly, Theorem 3 implies that when Landmark MDS is applied to a set of distances originating from Euclidean data, the resulting \( k \)-dimensional embedding recovers the projection of the data points onto the first \( k \) mean-centered principal axes of the designated landmark points. If the landmark points span a \( p \)-dimensional affine subspace, then by taking \( k = p \) we recover up to an isometry the projection of the data onto this subspace. Since the affine span of \( n \) points is at most \((n - 1)\)-dimensional, we need at least \( k + 1 \) landmark points in general position for a \( k \)-dimensional embedding.

In the remainder of this section we prove Theorem 3. First we fix our notation. Given a collection of landmark points \( \vec{y}_1, \vec{y}_2, \ldots, \vec{y}_n \in \mathbb{R}^m \) let \( \Delta_n \) denote their squared-distance matrix, with columns \( \vec{\delta}_j \) and column mean \( \vec{\delta}_\mu \). By translation we may assume that the landmarks are mean centered, or equivalently \( YH_n = Y \) where \( Y = [\vec{y}_1, \vec{y}_2, \ldots, \vec{y}_n] \). The principal axes are given by the eigenvectors of \( YY^T \); let \( \vec{p}_i \) denote a unit eigenvector for the \( i \)-th positive eigenvalue \( \lambda'_i \). By Theorem 1 we have the relation

\[
\sqrt{\lambda'_i} \vec{v}_i^T = \vec{p}_i^T Y
\]
where \( \vec{v}_i \) (chosen with the appropriate sign) and \( \lambda_i \) are defined as in classical MDS. Then

\[
\lambda_i = \sqrt{\lambda_i \vec{v}_i^T \vec{v}_i} = \vec{p}_i^T YY^T \vec{p}_i = \lambda'_i \vec{p}_i^T \vec{p}_i = \lambda'_i
\]

so we can write \( \lambda_i \) for \( \lambda'_i \).

Proof of Theorem 3. We must show that the \( i \)-th component of \( -\frac{1}{2} L_k^2 (\vec{\delta}_a - \vec{\delta}_\mu) \) is equal to the projection of \( \vec{y}_a \) onto the \( i \)-th principal axis. That is:

\[
-\frac{1}{2} \vec{v}_i^T \sqrt{\lambda_i} (\vec{\delta}_a - \vec{\delta}_\mu) = \vec{p}_i^T \vec{y}_a
\]  

(5)

First we compute the \( j \)-th entries of \( \vec{\delta}_a \) and \( \vec{\delta}_\mu \):

\[
[\vec{\delta}_a]_j = |\vec{y}_a|^2 - 2\vec{y}_j^T \vec{y}_a + |\vec{y}_j|^2
\]

\[
[\vec{\delta}_\mu]_j = \frac{1}{n} \sum_{k=1}^{n} |\vec{y}_k|^2 - 2\vec{y}_j^T \left( \frac{1}{n} \sum_{k=1}^{n} \vec{y}_k \right) + |\vec{y}_j|^2
\]

The left-hand terms are independent of \( j \), and the middle term on the second row is zero since the \( \vec{y}_k \) are mean centered. Thus we can subtract to derive the vector equation:

\[
\vec{\delta}_a - \vec{\delta}_\mu = c\vec{1} - 2Y^T \vec{y}_a
\]

where \( c = |\vec{y}_a|^2 - \frac{1}{n} \sum_{k=1}^{n} |\vec{y}_k|^2 \) is a scalar. Since \( \vec{v}_i^T \vec{1} = 0 \) we can drop the term \( c\vec{1} \) when substituting this formula into the left-hand side of (5).

Using (4) and this last observation, equation (5) becomes equivalent to

\[
\vec{p}_i^T YY^T \vec{y}_a = \lambda_i \vec{p}_i^T \vec{y}_a
\]
which is plainly true since $\vec{p}_i$ is the $i$-th eigenvector of $YY^T$ with eigenvalue $\lambda_i$. 

### 4.2 The approximately Euclidean case

The discussion in the previous section assumes that we have ideal data whose metric structure is exactly Euclidean. In practice it is more common for the distance matrix to be approximately but not exactly Euclidean. This may be due to noise, but it also happens systematically in applications such as Isomap, where larger entries in the distance matrix are extrapolated from smaller entries, resulting in a matrix that is at best an approximation to any underlying Euclidean structure.

In this section we study perturbation theory for classical MDS and Landmark MDS: given a small change in the distance matrix, what are the corresponding changes in the resulting $k$-dimensional embeddings? For example, what is the effect of perturbing the entries of the distance matrix away from an exactly Euclidean situation? If the answer is “there is not much change” then the results of the previous section apply more generally, with small error, to approximately Euclidean situations.

It turns out that classical MDS and Landmark MDS are stable up to orthogonal transformations, provided that $\lambda_k$ and $(\lambda_k - \lambda_{k+1})$ are not too small, where $\lambda_j$ denotes the $j$-th largest eigenvalue of the inner-product matrix $B_n$. The key technical result is the following theorem.

**Theorem 5** Let $\Delta_n$ be the squared-distance matrix of the set of landmark points; let $B_n$ be the corresponding inner-product matrix with eigenvalues $\lambda_1 \geq \ldots \geq \lambda_n$ and $\lambda_k > \max(0, \lambda_{k+1})$ for some $k$; and let $L_k, L_k^\dagger$ be the $k$-dimensional embedding matrix from classical MDS together with its pseudoinverse transpose. Consider a perturbation $\hat{\Delta}_n = \Delta_n + t\phi + O(t^2)$. Then there are
perturbations
\[
\hat{B}_n = B_n + t\beta + O(t^2)
\]
\[
\hat{L}_k = L_k + t\chi + O(t^2)
\]
\[
\hat{L}_k^\# = L_k^\# + t\psi + O(t^2)
\]
and an orthogonal matrix \( Q = Q(t) \) such that \( \hat{B}_n, Q\hat{L}_k \) and \( Q\hat{L}_k^\# \) are the corresponding matrices derived from \( \hat{\Delta}_n \). Moreover there are bounds

\[
\|\beta\| \leq \frac{\|\phi\|}{2}
\]
\[
\|\chi\| \leq \left\{ \frac{1}{4\lambda_k^{1/2}} + \frac{\lambda_1^{1/2}}{2(\lambda_k - \lambda_{k+1})} \right\} \|\phi\|
\]
\[
\|\psi\| \leq \left\{ \frac{1}{4\lambda_k^{3/2}} + \frac{1}{2\lambda_k^{1/2}(\lambda_k - \lambda_{k+1})} \right\} \|\phi\|
\]
in the Frobenius norm.

These are bounds on the first-order behaviour of the error terms, following the tradition of Sibson [21]. These first-order estimates make it clear how the perturbation theory for classical and Landmark MDS depends on the eigenvalues \( \lambda_1, \lambda_k \) and \( \lambda_{k+1} \). To obtain absolute bounds one would apply techniques of the kind used in [22], at considerably greater effort.

**Corollary 6** Let \( \delta_a = \vec{\delta}_a + t\vec{\zeta}_a + O(t^2) \) be a perturbation family for the vector of squared distances between the point \( a \) and the landmark points of Theorem 5. Then the perturbation family \( \hat{x}_a = \vec{x}_a - t(L_k^\# \vec{\zeta}_a + \psi \vec{\delta}_a) / 2 + O(t^2) \) agrees with the family of embedding vectors for \( a \) given by applying LMDS, up to a rotation and translation which depend on \( t \) but are independent of \( a \).

**Proof.** This is immediate from Theorem 5 and equation (3); the rotation is by \( Q \) and the translation deals with the term \(-(1/2)L_k^\# \vec{\delta}_\mu\) and its perturbation. \( \square \)
Corollary 6 establishes the overall stability of Landmark MDS in the following sense: the \( k \)-dimensional embedding given by a particular set of data closely approximates the \( k \)-dimensional embedding given by an approximation to that set of data, up to a rigid motion, provided the eigenvalue condition is satisfied for the landmark points. If we follow up with a PCA normalisation step, then the qualification “up to a rigid motion” is moot, in that the results of PCA remain essentially unchanged after a rigid motion is applied. The only ambiguity is that each coordinate is only well-defined up to a ± sign, and that principal components having the similar eigenvalues may become exchanged after perturbation; there is no real way round this. (Note the distinction between the eigenvalues of \( B_n \)—which are defined by reference to the landmark points only—and the eigenvalues at the PCA normalisation—which depend on the distribution of the entire data set at that time.)

The stability of classical MDS follows from Theorem 5, since \( \tilde{L}_k \) is the embedding matrix when all the points are regarded as landmark points.

Remark. Our approach differs from Sibson’s analysis of classical MDS [21] in the following way. Sibson compares the embedding matrix \( L_k \) directly with the embedding \( \tilde{L}_k \) for a nearby perturbed problem, estimating the error \( \| L_k - \tilde{L}_k \| \). In contrast we permit an orthogonal change of coordinates to improve the fit; essentially we are estimating \( \| L_k - Q\tilde{L}_k \| \) for a suitable orthogonal matrix \( Q \). Sibson’s approach gives poor bounds if any pair of eigenvalues is not well separated, since the individual ranked eigenvectors are required to be stable. In our case it is good enough for the subspace spanned by the the highest \( k \) eigenvectors to be stable; the final PCA normalisation step depends only on this subspace and not on the set of basis vectors which span it.

We now return to our main theorem. The proof is somewhat technical.

Proof of Theorem 5. First, the transformation \( \Delta_n \mapsto B_n \) is linear and has norm 1/2, giving us \( \beta \)
and the bound on its norm immediately.

Let \( V_k = [\vec{v}_1 \vec{v}_2 \ldots \vec{v}_k] \) be the matrix whose columns are the first \( k \) eigenvectors of \( B_n \) and let \( V_\perp = [\vec{v}_{k+1} \vec{v}_{k+2} \ldots \vec{v}_n] \) be the matrix containing the remaining eigenvectors. Let \( \Lambda_k \) be the diagonal matrix of eigenvalues for \( V_k \), defined \( \Lambda_k = V_k^T B_n V_k \). We can also define \( \Lambda_\perp = V_\perp^T B_n V_\perp \). Then \( L_k = \frac{\Lambda_k^{1/2} V_T}{V_k} \) and \( L_k^\sharp = \frac{\Lambda_k^{-1/2} V_T}{V_k} \) by definition. The columns of \( V_k \) span a subspace \( E^k \). Suppose we have a matrix \( W_k = [\vec{w}_1 \vec{w}_2 \ldots \vec{w}_k] \) whose columns form another orthonormal basis of \( E^k \). Then we can write \( W_k = V_k Q \) for some uniquely determined \( k \times k \) orthogonal matrix \( Q \).

**Lemma 7** We have the equations:

\[
L_k = Q(W_k^T B_n W_k)^{1/2} W_k^T
\]

\[
L_k^\sharp = Q(W_k^T B_n W_k)^{-1/2} W_k^T
\]

**Proof.** The formulae are derived by making substitutions into \( L_k = \frac{\Lambda_k^{1/2} V_T}{V_k} \) and \( L_k^\sharp = \frac{\Lambda_k^{-1/2} V_T}{V_k} \), and using \( \Lambda_k = V_k^T B_n V_k \).

\[\square\]

We apply the lemma not just to \( B_n, V_k \) and \( W_k \), but also to their perturbation families. We will define a perturbation family \( \hat{W}_k \) of orthonormal bases for the perturbed subspaces \( \hat{E}^k \) and set \( \hat{L}_k = (\hat{W}_k^T \hat{B}_n \hat{W}_k)^{1/2} \hat{W}_k^T \) and \( \hat{L}_k^\sharp = (\hat{W}_k^T \hat{B}_n \hat{W}_k)^{-1/2} \hat{W}_k^T \). The family \( \hat{W}_k \) will be more controllable than the naturally defined family \( \hat{V}_k \) whose columns give the first \( k \) eigenvectors in order. By the lemma, we recover the “correct” embedding matrices as \( Q \hat{L}_k \) and \( Q \hat{L}_k^\sharp \).

The family \( \hat{W}_k \) and a complementary family \( \hat{W}_\perp \) of orthonormal vectors take the following form, where \( \Theta \) is an \((n - k) \times k\) matrix:

\[
\hat{W}_k = V_k + tV_\perp \Theta + O(t^2)
\]

\[
\hat{W}_\perp = V_\perp - tV_k \Theta^T + O(t^2)
\]
For any choice of $\Theta$, it is easy to check to first order that the families $\hat{W}_k$ and $\hat{W}_\perp$ are orthonormal and mutually orthogonal. By considering a suitable exponential family of rotations, we can take families where this holds in fact, and not just to first order.

We now determine $\Theta$. The column space of $\hat{W}_k$ must equal the column space of $\hat{V}_k$, and so must be an invariant subspace of $\hat{B}_n$. This leads to the condition $\hat{W}_\perp^T \hat{B}_n \hat{W}_k = 0$. On expansion, we find that the zeroth-order part $V_\perp^T B_n V_k = 0$ is automatically true, so we are left with the first-order part of the equation

$$\Theta V_k^T B_n V_k - V_\perp^T B_n V_\perp \Theta = V_\perp^T \beta V_k$$

which is equivalent to $\Theta \Lambda_k - \Lambda_\perp \Theta = V_\perp^T \beta V_k$ and has the following componentwise solution:

$$\Theta_{ij} = \frac{[V_\perp^T \beta V_k]_{ij}}{\lambda_j - \lambda_{k+i}} \quad (6)$$

Since $\min(|\lambda_j - \lambda_{k+i}|) = \lambda_k - \lambda_{k+1}$, it follows that $\|\Theta\| \leq \|\beta\|/(\lambda_k - \lambda_{k+1})$.

Next we expand the following crucial term:

$$\hat{W}_k^T \hat{B}_n \hat{W}_k = (V_k^T + t\Theta^T V_k^T)(B_n + t\beta)(V_k + tV_\perp \Theta) + O(t^2)$$

$$= V_k^T B_n V_k + t (\Theta^T V_\perp^T B_n V_k + V_k^T \beta V_k + V_k^T B_n V_\perp \Theta) + O(t^2)$$

$$= \Lambda_k + t(V_k^T \beta V_k) + O(t^2)$$

In order to take a square root, note that $(\Lambda_k + t\nu + O(t^2))^{1/2} = \Lambda_k^{1/2} + t\eta + O(t^2)$ is equivalent to $\Lambda_k^{1/2}/\eta + \eta \Lambda_k^{1/2} = \nu$ and so $(\hat{W}_k^T \hat{B}_n \hat{W}_k)^{1/2} = \Lambda_k^{1/2} + t\eta + O(t^2)$ where:

$$\eta_{ij} = \frac{[V_\perp^T \beta V_k]_{ij}}{\lambda_i^{1/2} + \lambda_j^{1/2}} \quad (7)$$
It follows that $\|\eta\| \leq \|\beta\|/2\lambda_k^{1/2}$. This at last yields:

\[
\hat{L}_k = (\hat{W}_k^T \hat{B}_n \hat{W}_k)^{1/2} W_k^T \\
= \Lambda_k^{1/2} V_k^T + t(\eta V_k^T + \Lambda_k^{1/2} \Theta^T V_k^T) + O(t^2) \\
= L_k + t\chi + O(t^2)
\]

where $\|\chi\| \leq \|\eta V_k^T\| + \|\Lambda_k^{1/2} \Theta^T V_k^T\| = \|\eta\| + \|\Lambda_k^{1/2} \Theta^T\| \leq \|\eta\| + \lambda_k^{1/2}\|\Theta\|$. This gives the bound in the theorem.

To compute the inverse square root, $(\Lambda_k + t\nu + O(t^2))^{-1/2} = \Lambda_k^{-1/2} + t\xi + O(t^2)$ is equivalent to $\Lambda_k^{-1/2} \xi + \xi \Lambda_k^{-1/2} = -\Lambda_k^{-1} \nu \Lambda_k^{-1}$ and so $(\hat{W}_k^T \hat{B}_n \hat{W}_k)^{-1/2} = \Lambda_k^{-1/2} + t\xi + O(t^2)$ where:

\[
\xi_{ij} = -\frac{[V_k^T \beta V_k]_{ij}}{\lambda_i \lambda_j (\lambda_i^{-1/2} + \lambda_j^{-1/2})} \tag{8}
\]

It follows that $\|\xi\| \leq \|\beta\|/2\lambda_k^{3/2}$. Finally:

\[
\hat{L}_k^\sharp = (\hat{W}_k^T \hat{B}_n \hat{W}_k)^{-1/2} W_k^T \\
= \Lambda_k^{-1/2} V_k^T + t(\xi V_k^T + \Lambda_k^{-1/2} \Theta^T V_k^T) + O(t^2) \\
= L_k^\sharp + t\psi + O(t^2)
\]

where $\|\psi\| \leq \|\xi V_k^T\| + \|\Lambda_k^{-1/2} \Theta^T V_k^T\| = \|\xi\| + \|\Lambda_k^{-1/2} \Theta^T\| \leq \|\xi\| + \lambda_k^{-1/2}\|\Theta\|$. This gives the bound stated in the theorem, and completes the proof. $\square$
4.3 A non-Euclidean counterexample

As we have seen, Landmark MDS is a good approximation to classical MDS when the distance matrix $D$ is low-rank Euclidean, or approximately so. For data with non-Euclidean geometry, the two algorithms may give quite different results, as demonstrated by the following counterexample.

Consider 50 points evenly spaced around a unit circle, and let $d$ denote the arc-length metric. This is quite different from the Euclidean chord-length metric $d'$. When $x$ and $y$ are nearby points then $d(x, y) \approx d'(x, y)$, but when $x$ and $y$ are diametrically opposite then $d(x, y) = \pi$ whereas $d'(x, y) = 2$. Let $D$ and $D'$ denote the corresponding distance matrices. Since $D'$ is Euclidean, classical MDS recovers the original 2-dimensional unit circle and Landmark MDS successfully finds the same embedding (up to a rigid motion) provided at least 3 landmarks are used. On the other hand $D$ is highly non-Euclidean: its inner-product matrix $B$ has positive and negative eigenvalues, occurring in identical pairs. Figure 8 (left) shows the first 10 eigenvalues, ranked by magnitude. Using the first two eigenvalues only, classical MDS embedding appears to give the correct result: a two dimensional embedding of a circle (centre). However this neglects much of the geometry in $D$, as expressed in the remaining eigenvectors. When we run Landmark MDS with 5 evenly-spaced landmarks, we get something unexpected (right). The five landmarks are symmetrically arrayed, but the apparent geometry of the remaining points is quite different. In particular, the full symmetry of the original distance matrix has been lost.
The moral is that in the non-Euclidean setting the low-dimensional embeddings given by classical MDS and Landmark MDS need not approximate each other particularly well. To be fair, these are situations where the use of classical MDS may not be all that appropriate. For data sets which are well-suited to classical MDS, with a clear $k$-dimensional metric embedding and small eigenvalues for the remaining ‘noise’ coordinates, we emphasise that Landmark MDS does indeed give a good approximation to classical MDS.

Remark. Genuinely non-Euclidean data sets such as the circle example in Section 4.3 cannot sensibly be treated as ‘Euclidean but noisy’. One possibility is to think of such a data set as living in Minkowski space, with imaginary (‘timelike’) dimensions alongside real (‘spacelike’) dimensions, and carry out a similar analysis to the Euclidean case. Goldfarb [6] lays out some of the foundations for this point of view. We do not pursue the idea here.

4.4 Choosing a good landmark set

What constitutes a good set of landmark points? From the example of the noisy grid, we know that there is better accuracy with more landmarks; and that there are systematic errors in the output when the landmarks are in degenerate configuration (lying close to a low-dimensional subspace such as a line).

The situation can be described in the following way. Suppose first that we have a mean-centered Euclidean configuration of $N$ points in $\mathbb{R}^d$, represented by a $d \times N$ matrix $Z$. The first $k$ principal components are obtained by projection onto the top $k$ eigenvectors of the covariance matrix $C_Z = (1/N)ZZ^T$. Let $E_k^Z$ denote the subspace spanned by these top $k$ eigenvectors. A set of $n$ landmark points is represented by a submatrix $Y$, and has (mean-centered) covariance matrix $C_Y = (1/n)YY^T$ and an associated subspace $E_k^Y$ spanned by the top $k$ eigenvectors of $C_Y$.

We now seek a $k$-dimensional embedding. Running classical MDS is equivalent to projecting
the points onto the subspace $E_Z^k$, with axes being the eigenvectors of $C_Z$ in order. Running LMDS with the given landmark points is equivalent to projecting the data onto $E_Y^k$, and then realigning the data according to the new principal components of the projected data. Informally, we can say:

**Proposition 8** For Euclidean data, Landmark MDS succeeds when the subspace $E_Y^k$ is close to the subspace $E_Z^k$, and more specifically when the data has a similar covariance structure when projected onto each of the two subspaces.

More generally, if $j \leq k$ one can restrict the $k$-dimensional embeddings produced by classical MDS and LMDS to the first $j$ coordinates. Landmark MDS succeeds when the corresponding $j$-dimensional subspaces of $E_Y^k$ and $E_Z^k$ are close and have similar covariance structure for the data. The remaining dimensions of $E_Y^k$ and $E_Z^k$ need not agree closely.

This gives an argument in favour of choosing landmark points randomly from the given data set; with enough landmark points the covariance matrix of the sample is a good approximation to the covariance matrix of the data. Provided that the $k$-th and $(k+1)$-st eigenvalues are sufficiently well separated (or the $j$-th and $(j+1)$-st, if restricting), the corresponding subspaces will be close and Landmark MDS will succeed.

When the covariance matrix of the landmark points satisfies the separation condition and is close to the true covariance matrix of the full data, we may still observe noise at fine scales. The variance of this noise behaves as $\sim n^{-1}$, where $n$ is the number of landmarks. The back-of-the-envelope calculation runs as follows: $\|L_k\| \sim n^{1/2}$ so $\|L_k^\sharp\| \sim n^{-1}\|L_k\|$. For a given point $a$ there are $n$ entries in $\vec{\delta}_a$. If each entry independently is subject to a certain amount of noise, then the discrepancy in the embedding of $a$ takes the form $n^{-1} \cdot (n$ independent random variables), which has variance $\sim n^{-1}$ as claimed.

When the covariance matrix of the landmark points has poorly separated $k$-th and $(k+1)$-st
Figure 9: Foreshortening. Points sampled noisily from the black line become closer together after projection onto the LMDS principal axis generated by the two red landmark points (left). The effect is much less pronounced with one extra landmark point (right).

Eigenvalues, then with noisy data we run into the two systematic problems discussed in Section 3.1: foreshortening and noise amplification. Noise amplification was explained earlier. Foreshortening is explained in the left panel of Figure 9. Data have been sampled with noise from the solid black line. Two landmark points are chosen; they span the broken red line. The projection from the original subspace (black) onto the estimated subspace (red) diminishes distances by a constant linear factor, the cosine of the angle between the lines. Since the angle here is quite large, this observation holds approximately for data close to the original subspace (and not just on it).

It is clear that there is much less of a problem if the two landmarks are better separated; for example, taking the two extreme points on the left and the right. This is evidence in favour of the MaxMin approach, which by design rapidly seeks out extreme, well-separated landmarks. On the other hand, there is a corresponding danger of selecting outlier points. And, in fairness, random choice will eventually perform well if enough points are chosen; adding even one more landmark point in this example makes a big difference (right panel of Figure 9). However, “enough” may be too many in some cases. The user must decide from experience which technique to choose for a given data set.
4.4.1 Cross-validation

As a precaution against the kind of bad luck arising from a poor landmark configuration, it may be wise to cross-validate by running LMDS several times and computing correlations between the respective coordinates. Those trials which closely approximate the (uncomputed) classical MDS embedding will closely correlate with each other. The occasional ‘bad’ embedding is easily detected by its comparatively low average correlation with the remaining trials.

More generally, this leads to a method for determining the ideal number of landmark points. Run Landmark MDS repeatedly in batches, where each batch consists of several trials with the same number of landmarks chosen randomly, and compute correlations between the trials. This process is repeated, with an increasing number of landmarks, until there are consistent correlations across (most of) the trials in a batch. This is still much cheaper than running full MDS on a large set.

5 Discussion

We briefly describe some of the more important applications of Landmark MDS, and mention some of its connections with related work.

5.1 Landmark Isomap

The initial motivation for Landmark MDS was the need to speed up the Isomap algorithm [14] for nonlinear dimensionality reduction (NLDR). Isomap constructs a low-dimensional embedding of of nonlinear data set observed in a high-dimensional space. There are three steps: (i) a neighbourhood relation is determined in the form of a sparse graph, with edges connecting data point pairs that are deemed to be “close” (for example, $x, y$ are neighbours iff $x$ is one of the $p$ nearest neighbours of $y$.
and vice versa); (ii) a new metric is defined: the distance between a pair of data points is the length of the shortest path between them, with edges weighted by length in the observation space; (iii) classical MDS is applied to the new metric find the low-dimensional embedding. This procedure emphasises the local structure of the data, but at the same time it globalises this information so that the embedding can be found using a global optimisation (here classical MDS). Highly non-linear data sets can be successfully linearised in this way, for example, a sample of images of a face under varying pose and lighting conditions, or a sample of images of a hand as the fingers and wrist move [14].

For a large data set of \( N \) points the main bottleneck is the classical MDS stage, which reduces to an \( O(N^3) \) eigenvalue problem on a dense \( N \times N \) matrix. Storage of the full distance matrix is also an issue. Other well-known NLDR algorithms such as Locally Linear Embedding [23] and Laplacian Eigenmaps [24] involve eigenvalue problems on sparse \( N \times N \) matrices, so they do not suffer from either of these issues. Landmark MDS solves both problems, since it only requires an \( n \times N \) submatrix of the full distance matrix, and the eigenvalue problem is \( O(n^3) \). Using Dijkstra’s algorithm, the shortest-paths distance can be calculated row-by-row, so there are no extraneous calculations. If the neighbourhood graph is given, the cost of running Isomap with \( N \) points and \( p \) nearest neighbours is \( O(N^3 + pN^2 \log N) \) as opposed to \( O(pmN \log N + knN + n^3) \) for Landmark Isomap with \( n \) landmarks in \( k \) dimensions, a huge savings. The cost of building the \( p \)-nearest-neighbours graph—the same in both cases—softens the advantage slightly. Standard methods have complexity \( O(N^2) \), but there are more sophisticated techniques [25, 26] which give an expected bound of \( c^{O(1)}(pN + N \log N \log \log N) \) for data sets with an expansion coefficient of \( c \). For data sampled randomly from a \( k \)-dimensional submanifold of Euclidean space \( \mathbb{R}^d \), the expansion coefficient is essentially \( c = 2^k \).
5.2 Sparsely observed distance matrices

Isomap imposes a sparsification of the original distance metric of the data, by using only the distance information for pairs of neighbouring points. A different situation occurs for large data sets when the metric information is inherently sparsely observed, for instance when the metric comes from human judgments of similarity [7] or association [27]. This leads to the problem of multidimensional scaling with incomplete information – how does one find a good low-dimensional representation of data from a sparse set of dissimilarity coefficients? One possible answer is to estimate the missing coefficients using shortest paths in the weighted graph having an edge for every observed comparison, with weight equal to the dissimilarity coefficient; and then use classical MDS. This is formally analogous to Isomap, except that the graph is not a neighbourhood graph. Steyvers et al [27] use this approach to estimate the missing coefficients in a matrix of similarities generated by human test subjects. Platt [7] shows that good results can be obtained by combining this method with LMDS; this avoids having to deal with the full dissimilarity matrix.

5.3 Nyström method

Landmark MDS is similar in spirit to the well-known Nyström method, which finds approximate solutions to a positive semi-definite symmetric eigenvalue problem using just a few of the columns of the matrix, and it is natural to suspect that they are closely related. In fact, Bengio et al. [8] establish a precise connection, using a Nyström-like approach to derive formulae equivalent to Step 3 of LMDS for the purposes of projecting new points into a pre-existing MDS solution. Bengio et al. also derive Nyström-based out-of-sample extensions to other dimensionality reduction algorithms, including Locally Linear Embedding [10,23] and Laplacian Eigenmaps [24] (where the initial eigenvalue problem is not as acutely expensive as it is for Isomap). Our work here shows that this approach provides a rigorous basis for efficient and accurate landmark-based approximations to the
full MDS computation on large data sets.

5.4 FastMap

An early example of cheaply approximating classical MDS comes from Faloutsos and Lin [9], with their FastMap algorithm. The idea is to construct a $k$-dimensional embedding one dimension at a time. To determine an axis, two points are selected from the data and the distances of all points to these two points are used to find the first embedding coordinate. The distance matrix is then modified to take the first dimension into account, and the process is repeated, with a new axis, to find second and subsequent coordinates. In fact, FastMap is exactly an iterated form of LMDS in the simplest case of 2 landmarks.

An inherent weakness of FastMap is that each projection axis is specified by only two points, so the alignment of these axes with the true principal axes of the data is unstable at best. In contrast, Landmark MDS makes use of as many landmarks as provided, even if the required embedding is much lower dimensional. Redundancy brings stability; and LMDS is not that much more expensive than FastMap to run. In tests carried out by Platt [7], LMDS gives better performance than FastMap.

Conclusion

Dimensionality reduction and effective visualization of the intrinsic structure of a data set constitute a major challenge in statistics, machine learning, information retrieval, and knowledge discovery. Classical multidimensional scaling has proved to be an excellent approach to these problems, either by itself, or as part of a larger scheme dealing with cases where the data are nonlinear or incomplete.

However, modern data sets present a challenge not well addressed by the classical technique of MDS: how does one deal with large volumes of data? This paper has addressed that challenge,
describing an approximation to the algorithm which works tractably with very large data sets, and providing a theoretical analysis that explains when and where our more efficient methods can be trusted to give results faithful to the output of classical MDS.

Given its simplicity, efficiency and accuracy, Landmark MDS should be an important part of the next generation data analyst’s toolkit.

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References


