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Adaptive Interpolation of Multidimensional Scaling

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Abstract

The recent explosion of publicly available biology gene sequences and chemical compounds offers an unprecedented opportunity for data mining. To make data analysis feasible for such vast volume and high-dimensional scientific data, we apply high performance dimension reduction algorithms. It facilitates the investigation of unknown structures in a three dimensional visualization. Among the known dimension reduction algorithms, we utilize the multidimensional scaling (MDS) algorithm to configure the given high-dimensional or abstract data into a target dimension. However, the MDS algorithm requires large physical memory as well as computational resources. In order to reduce computational complexity and memory requirement effectively, the interpolation method of the MDS was proposed in 2010. With minor trade-off of approximation, the MDS interpolation method enables us to process millions of data points with modest amounts of computation and memory requirement. In this paper, we would like to improve the mapping quality of the MDS interpolation approach by adapting the original dissimilarity based on the ratio between the original dissimilarity and the corresponding mapping distances. Our experimental results illustrate that the quality of interpolated mapping results are improved by adding the adaptation step without runtime loss compared to the original interpolation method. With the proposed adaptive interpolation method, we construct a better configuration of millions of *out-of-sample* data into a target dimension than the previous interpolation method.

Keywords: dimension reduction, multidimensional scaling, interpolation, adaptation

1. Introduction

Due to the advancements in science and technology over the last several decades, every scientific and technical field has generated a huge amount of data as time has passed in the world. We are really in the era of data deluge. In reflection on the data deluge era, data-intensive scientific computing [1] has emerged in the scientific computing fields and it has been attracting more by many people. To analyze those incredible amount of data, many data mining and machine learning algorithms have been developed. Among many data mining and machine learning algorithms that have been invented, we focus on dimension reduction algorithms, which reduce data dimensionality from original high dimension to target dimension, in this paper.

Among the many dimension reduction algorithms which exist, such as principle component analysis (PCA), generative topographic mapping (GTM) [2, 3], self-organizing map (SOM) [4], and multidimensional scaling (MDS) [5, 6],

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we discuss MDS in this paper since it is popular and theoretically strong. The parallelization of MDS algorithm was studied in [7] which aims at utilizing multicore clusters and increasing the computational capability with minimal overhead for the purpose of investigating large data, such as 100,000 data. However, parallelization of an MDS algorithm, whose computational complexity and memory requirement is upto $O(N^2)$ where N is the number of points, is still limited by the memory requirement for huge data, e.g. millions of points, although it utilizes distributed memory environments, such as clusters, for acquiring more memory and computational resources. In this paper, we try to solve the memory-bound problem by interpolation based on pre-configured mappings of the sample data for MDS algorithm, so that we can provide configuration of millions of points in the target space.

This paper is organized as follows. First, we briefly discuss about multidimensional scaling (MDS) in Section 2. The various existed methods of *out-of-sample* approach related to the MDS are explained in Section 3 and Section 4. Then, the proposed adaptive interpolation method is described in Section 5. The quality comparison between interpolated results and full MDS running results and runtime evaluation of those algorithms are shown in Section 6 followed by our conclusion in Section 7.

2. Multidimensional Scaling (MDS)

Multidimensional scaling(MDS) [5, 6] is a general term for the techniques of configuration of the given high dimensional data into a target dimension based on the pairwise proximity information of the data, while each Euclidean distance between two points becomes as similar to the corresponding pairwise dissimilarity as possible. In other words, MDS is a non-linear optimization problem with respect to mapping in the target dimension and original proximity information.

Formally, the pairwise proximity information is given as an $N \times N$ matrix ($\Delta = [\delta_{ij}]$), where N is the number of points and δ_{ij} is the given dissimilarity value between point i and j in the original data space. (1) Symmetric ($\delta_{ij} = \delta_{ji}$), (2) non-negative ($\delta_{ij} \geq 0$), and (3) zero diagonal ($\delta_{ii} = 0$) are the constraints of the dissimilarity matrix Δ . The output of MDS algorithm could be represented as an $N \times L$ configuration matrix (X), where L is the target dimension, and each data point $x_i \in \mathbb{R}^L$ ($i = 1, \dots, N$) resides in i -th rows of X .

The evaluation of the constructed configuration is done with respect to the well-known objective functions of MDS, namely STRESS [8] or SSTRESS [9]. Below equations are the definitions of STRESS (1) and SSTRESS (2):

$$\sigma(X) = \sum_{i < j \leq N} w_{ij} (d_{ij}(X) - \delta_{ij})^2 \quad (1)$$

$$\sigma^2(X) = \sum_{i < j \leq N} w_{ij} [(d_{ij}(X))^2 - (\delta_{ij})^2]^2 \quad (2)$$

where $1 \leq i < j \leq N$, d_{ij} is a mapping distance between point i and j , and w_{ij} is a weight value, so $w_{ij} \geq 0$.

3. Related Work

The *out-of-sample* method, which embeds new points with respect to previously configured points, has been actively researched for recent years, and it aims at improving the capability of dimension reduction algorithms by reducing the computational and memory-wide requirement with the trade-off of slightly approximated mapping result.

In a sensor network localization field, when there are only a subset of pairwise distances between sensors and a subset of anchor locations are available, people try to find out the locations of the remaining sensors. For instance, the semi-definite programming relaxation approaches and its extended approaches has been proposed to solve this issue [10]. [11] and [12] proposed out-of-sample extension for the classical multidimensional scaling (CMDS) [13], which is based on spectral decomposition of a symmetric positive semidefinite matrix (or the approximation of positive semidefinite matrix), and the embeddings in the configured space are represented in terms of eigenvalues and eigenvectors of it. [11] projected the new point x onto the principal components, and [12] extends the CMDS algorithm itself to the out-of-sample problem. In [12], the authors describe how to embed one point between the embeddings of the original n objects through modification of the original CMDS equations, which preserves the mappings of the original n objects, with $(n + 1) \times (n + 1)$ matrix A_2 instead of $n \times n$ matrix A_2 , and extends to embedding a number of

points simultaneously by using matrix operations. Recently, a multilevel force-based MDS algorithm was proposed as well [14].

In contrast to applying the out-of-sample problem to CMDs, out-of-sample approach to metric MDS with STRESS criteria of Eq. (1) was proposed by Bae et al. [15], which finds embeddings of approximating to the distance (or dissimilarity) rather than the inner product as in CMDs, with an gradient descent optimization method, called iterative majorizing. The details of the iterative majorizing interpolation approach for the MDS problem [15] is explained in Section 4.

4. Majorizing Interpolation MDS

One of the main limitation of most MDS applications is that it requires $O(N^2)$ memory as well as $O(N^2)$ computation. Thus, though it is possible to run them with small data size without any trouble, it is impossible to execute them with a large number of data due to memory limitation; therefore, this challenge could be considered as being a memory-bound problem. For instance, Scaling by MAjorizing of COmplicated Function (SMACOF) [16, 17], a well-known MDS application via a kind of Expectation-Maximization (EM) [18] approach, uses six $N \times N$ matrices. If $N = 100,000$, then one $N \times N$ matrix of 8-byte double-precision numbers requires 80 GB of main memory, so the algorithm needs to acquire at least 480 GB of memory to store six $N \times N$ matrices. It is possible to run parallel version of SMACOF with MPI on the testbed system in Table 1 with $N = 100,000$. If the data size is increased only twice, however, then SMACOF algorithm should have 1.92 TB of memory, which is bigger than total memory of the system in Table 1 (1.536 TB), so it is impossible to run it within the cluster. Increasing memory size will not be a solution, even though it could increase the runnable number of points. It will encounter the same problem as the data size increases.

To solve this obstacle, Bae et al. developed a simple interpolation approach based on pre-mapped MDS result of the sample of the given data [15]. The interpolation algorithm [15] is similar to k nearest neighbor (k -NN) classification [19], but it approximates new mapping position of the new point based on the positions of k -NN, among pre-mapped subset data, instead of classifying it. For the purpose of deciding new mapping position in relation to the k -NN positions, the iterative majorization method is applied as in the SMACOF [16, 17] algorithm. The algorithm proposed in [15] is called Majorizing Interpolation of MDS (hereafter *MI-MDS*), and the summary of MI-MDS is in this section as below.

The MI-MDS algorithm is implemented as follows. We are given N data in a high-dimensional space, say D -dimension, and proximity information ($\Delta = [\delta_{ij}]$) of those data as in Section 2. Among N data, the configuration of the n sample points in L -dimensional space, $\mathbf{x}_1, \dots, \mathbf{x}_n \in \mathbb{R}^L$, called X , are already constructed by an MDS algorithm; here we use SMACOF algorithm. Then, we select k nearest neighbors ($\mathbf{p}_1, \dots, \mathbf{p}_k \in \mathbf{P}$) of the given new point, where \mathbf{P} is defined as a set of k -NNs, among n pre-mapped points with respect to corresponding δ_{ix} , where \mathbf{x} represents the new point. A linear search is used to find k -nearest neighbors among n -sampled data, so that the complexity of finding k -nearest neighbors is $O(n)$ per one interpolated point (here \mathbf{x}). Finally, the new mapping of the given new point $\mathbf{x} \in \mathbb{R}^L$ is calculated based on the pre-mapped position of the selected k -NN and the corresponding proximity information δ_{ix} . The finding new mapping position is considered as a minimization problem of STRESS (3) as similar as normal MDS problem with m points, where $m = k + 1$. However, only one point (\mathbf{x}) is movable among m points, so we can simplify the STRESS equation (3) as follows (Eq. (4)), and we set $w_{ij} = 1$, for $\forall i, j$ in order to simplify.

$$\sigma(X) = \sum_{i < j \leq m} (d_{ij}(X) - \delta_{ij})^2 \quad (3)$$

$$= C + \sum_{i=1}^k d_{ix}^2 - 2 \sum_{i=1}^k \delta_{ix} d_{ix} \quad (4)$$

where δ_{ix} is the original dissimilarity value between \mathbf{p}_i and \mathbf{x} , d_{ix} is the Euclidean distance in L -dimension between \mathbf{p}_i

and \mathbf{x} , and C is constant part. The second term of Eq. (4) can be deployed as following:

$$\sum_{i=1}^k d_{ix}^2 = \|\mathbf{x} - \mathbf{p}_1\|^2 + \cdots + \|\mathbf{x} - \mathbf{p}_k\|^2 \quad (5)$$

$$= k\|\mathbf{x}\|^2 + \sum_{i=1}^k \|\mathbf{p}_i\|^2 - 2\mathbf{x}^t \mathbf{q} \quad (6)$$

where $\mathbf{q}^t = (\sum_{i=1}^k p_{i1}, \dots, \sum_{i=1}^k p_{iL})$ and p_{ij} represents j -th element of \mathbf{p}_i . In order to establish majorizing inequality, we apply *Cauchy-Schwarz* inequality to $-d_{ix}$ of the third term of Eq. (4). Please, refer to chapter 8 in [6] for details of how to apply *Cauchy-Schwarz* inequality to $-d_{ix}$. Since $d_{ix} = \|\mathbf{p}_i - \mathbf{x}\|$, $-d_{ix}$ could have following inequality based on *Cauchy-Schwarz* inequality:

$$-d_{ix} \leq \frac{\sum_{a=1}^L (p_{ia} - x_a)(p_{ia} - z_a)}{d_{iz}} \quad (7)$$

$$= \frac{(\mathbf{p}_i - \mathbf{x})^t (\mathbf{p}_i - \mathbf{z})}{d_{iz}} \quad (8)$$

where $\mathbf{z}^t = (z_1, \dots, z_L)$ and $d_{iz} = \|\mathbf{p}_i - \mathbf{z}\|$. The equality in Eq. (7) occurs if \mathbf{x} and \mathbf{z} are equal. If we apply Eq. (8) to the third term of Eq. (4), then we obtain

$$-\sum_{i=1}^k \delta_{ix} d_{ix} \leq -\sum_{i=1}^k \frac{\delta_{ix}}{d_{iz}} (\mathbf{p}_i - \mathbf{x})^t (\mathbf{p}_i - \mathbf{z}) \quad (9)$$

$$= -\mathbf{x}^t \sum_{i=1}^k \frac{\delta_{ix}}{d_{iz}} (\mathbf{z} - \mathbf{p}_i) + C_\rho \quad (10)$$

where C_ρ is a constant. If Eq. (6) and Eq. (10) are applied to Eq. (4), then it could be like following:

$$\sigma(\mathbf{X}) = C + \sum_{i=1}^k d_{ix}^2 - 2 \sum_{i=1}^k \delta_{ix} d_{ix} \quad (11)$$

$$\leq C + k\|\mathbf{x}\|^2 - 2\mathbf{x}^t \mathbf{q} + \sum_{i=1}^k \|\mathbf{p}_i\|^2 - 2\mathbf{x}^t \sum_{i=1}^k \frac{\delta_{ix}}{d_{iz}} (\mathbf{z} - \mathbf{p}_i) + C_\rho \quad (12)$$

$$= \tau(\mathbf{x}, \mathbf{z}) \quad (13)$$

where both C and C_ρ are constants. In the Eq. (13), $\tau(\mathbf{x}, \mathbf{z})$, a quadratic function of \mathbf{x} , is a majorization function of the STRESS. Through setting the derivative of $\tau(\mathbf{x}, \mathbf{z})$ equal to zero, we can obtain minimum of it; that is

$$\nabla \tau(\mathbf{x}, \mathbf{z}) = 2k\mathbf{x} - 2\mathbf{q} - 2 \sum_{i=1}^k \frac{\delta_{ix}}{d_{iz}} (\mathbf{z} - \mathbf{p}_i) = 0 \quad (14)$$

$$\mathbf{x} = \frac{\mathbf{q} + \sum_{i=1}^k \frac{\delta_{ix}}{d_{iz}} (\mathbf{z} - \mathbf{p}_i)}{k} \quad (15)$$

where $\mathbf{q}^t = (\sum_{i=1}^k p_{i1}, \dots, \sum_{i=1}^k p_{iL})$, p_{ij} represents j -th element of \mathbf{p}_i , and k is the number of the nearest neighbors that we selected.

Finally, if we substitute \mathbf{z} with $\mathbf{x}^{[t-1]}$ in Eq. (15), then we generate an iterative majorizing equation like the following:

$$\mathbf{x}^{[t]} = \bar{\mathbf{p}} + \frac{1}{k} \sum_{i=1}^k \frac{\delta_{ix}}{d_{iz}} (\mathbf{x}^{[t-1]} - \mathbf{p}_i) \quad (16)$$

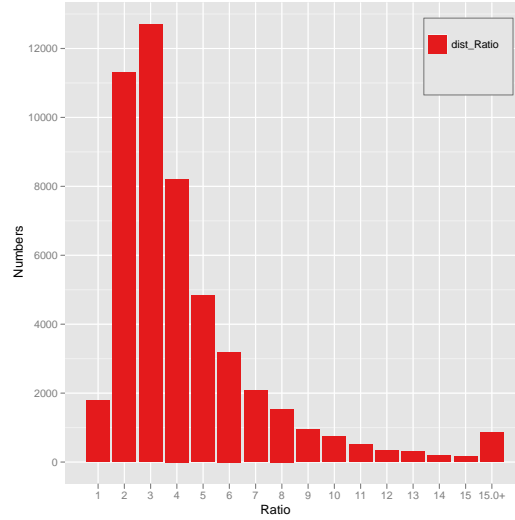


Figure 1: The reciprocal of the distance ratio ($1/r$) between original dissimilarities among k nearest neighbors and the corresponding mapping distances among k nearest neighbors. The tested data here is pubchem data of total 100k and 50k sampled case and $k = 2$.

where $d_{iz} = \|\mathbf{p}_i - \mathbf{x}^{[t-1]}\|$ and $\bar{\mathbf{p}}$ is the average of k -NN's mapping results. Eq. (16) is an iterative equation used to embed newly added point into target-dimensional space, based on pre-mapped positions of k -NN. The iteration stop condition is essentially the same as that of the SMACOF algorithm, which is

$$\Delta\sigma(\mathbf{S}^{[t]}) = \sigma(\mathbf{S}^{[t-1]}) - \sigma(\mathbf{S}^{[t]}) < \varepsilon, \quad (17)$$

where $\mathbf{S}^{[t]} = \mathbf{P} \cup \{\mathbf{x}^{[t]}\}$ and ε is the given threshold value.

The time complexity of the MI-MDS algorithm [15] to find the mapping of one interpolated point is $O(k)$ on the basis of Eq. (16), if we assume that the number of iterations of finding one interpolated mapping is very small. Since finding nearest neighbors takes $O(n)$ and mapping via MI-MDS requires $O(k)$ for one interpolated point, the overall time complexity to find mappings of overall out-of-sample points ($N-n$ points) via the MI-MDS algorithm is $O(kn(N-n)) \approx O(n(N-n))$, due to the fact that k is usually negligible compared to n or N .

The process of the overall out-of-sample MDS with a large dataset could be summarized by the following steps: (1) Sampling; (2) Running MDS with sample data; and (3) Interpolating the remain data points based on the mapping results of the sample data.

5. Adaptive Interpolation of MDS

An intrinsic assumption of the MI-MDS [15] is that the mapping distances of k -NNs might be highly likely similar to the original dissimilarities of k -NNs. In real life, it is unlikely happened because the MDS results, which is used for prior mapping of sampled data, are usually produced with positive normalized STRESS values, which represent normalized errors of the constructing mappings in the target dimension. There is a possibility, however, that the smaller original dissimilarities will be represented in the smaller distances and the larger ones will be represented in the larger distances by the full MDS mappings of sampled data with regard to the distribution of the original dissimilarities and the distribution of the mapping distances of sampled data in target space. In other words, the mappings within a small region might be mapped in the target space within a proportional distances of the original dissimilarities by the full MDS running of the sampled data. On a basis of the above phenomenon, in this paper, we would like to propose an adaptive interpolation method which aims at improving the mapping quality compared to the MI-MDS method.

5.1. Mapping Distance and Original Dissimilarity Comparison

At first, we investigated the distance ratio ($r = \bar{d}_{ij}/\bar{\delta}_{ij}$) between the average of mapping distances among k -NNs (\bar{d}_{ij}) and the average of the corresponding original dissimilarities among k -NNs ($\bar{\delta}_{ij}$) of each interpolated point based

on the prior mappings the of sampled data and the corresponding pubchem data. The explanation of the pubchem data is in Section 6. Fig. 1 shows the distribution of the reciprocal of the distance ratio of 100,000 pubchem data with sampling 50,000 when $k = 2$.

Among about 50,000 cases, over 96% of the reciprocal of the distance ratio ($1/r$) is larger than 1.0 and around 75% of that is in between 1.0 and 5.0. Interestingly, the cases of larger than 15.0 occur around 1,000 times. Those are the cases that the mapping distances (d_{ij}) are much smaller than the corresponding original dissimilarities (δ_{ij}). We exclude the case of $\delta_{ij} = 0.0$ in Fig. 1. Based on Fig. 1, we think that the adaptation of the original dissimilarities (δ_{ix}) between the interpolated point (\mathbf{x}) and the k -NNs (\mathbf{p}_i , where $1 \leq i \leq k$) can be helpful to get better interpolation mapping result.

5.2. Adaptive Original Dissimilarity of k Nearest Neighbors of an Interpolated Point

In the above section, we found that there is some difference between the prior mapping distances and the corresponding original dissimilarities of k -NNs. Since the interpolation method generates the mapping of each interpolated point based on the relation to the prior mappings of k -NNs of the point, it will be better to adjust the given original dissimilarities (δ_{ix}) between the interpolated point (\mathbf{x}) and its k -NNs based on the distance ratio (r) of its k -NNs. Thus, we propose an adaptive interpolation of MDS which constructs a mapping of a new point based on the adapted dissimilarities.

The proposed adaptive interpolation of MDS (hereafter called *AI-MDS*) will interpolate points based on the prior mappings of the sampled data in terms of the adaptive dissimilarities between interpolated points and k -NNs. Those adaptive dissimilarities ($\widehat{\delta}_{ix}$) are calculated by multiplying the original dissimilarities (δ_{ix}) and the distance ratio ($r = d_{ij}/\bar{d}_{ij}$) which is defined in Section 5.1, as shown in Eq. (18).

$$\widehat{\delta}_{ix} = \delta_{ix} \cdot \frac{\bar{d}_k}{\bar{\delta}_k} \quad (18)$$

Respectively, AI-MDS substitutes Eq. (16) of the iterative majorizing interpolation process with Eq. (19), by exchanging δ_{ix} with $\widehat{\delta}_{ix}$.

$$\mathbf{x}^{[t]} = \bar{\mathbf{p}} + \frac{1}{k} \sum_{i=1}^k \frac{\widehat{\delta}_{ix}}{d_{iz}} (\mathbf{x}^{[t-1]} - \mathbf{p}_i). \quad (19)$$

The summary of the proposed AI-MDS algorithm for interpolation of a new data, say \mathbf{x} , in relation to pre-mapping result of the sample data is described in Alg. 1. Note that the algorithm uses $\bar{\mathbf{p}}$ as an initial mapping of the new point $\mathbf{x}^{[0]}$ unless initialization with $\bar{\mathbf{p}}$ makes $d_{ix} = 0$, since the mapping is based on the k -NN. $\bar{\mathbf{p}}$ makes $d_{ix} = 0$, if and only if all the mapping positions of the k -NNs are on the same position. If $\bar{\mathbf{p}}$ makes $d_{ix} = 0$ ($i = 1, \dots, k$), then we generate a random variation from the $\bar{\mathbf{p}}$ point with the average distance of δ_{ix} as an initial position of $\mathbf{x}^{[0]}$.

5.3. Parallelization of MDS Interpolation Algorithms

Suppose that, among N points, the mapping results of n sample points in the target dimension, say L -dimension, are given so that we could use those pre-mapped results of n points via MDS interpolation algorithms which are described above to embed the remaining points ($M = N - n$). Though interpolation approach is very fast algorithm, i.e. $O(Mn)$, implementing parallel MDS interpolation algorithms is essential, since the out-of-sample size can be still huge, like millions. In addition, most of clusters are now in forms of multicore-clusters, so we are using hybrid-model parallelism, which combine processes and threads together as used in [20, 1].

In contrast to the original MDS algorithm in which the mapping of a point is influenced by the other points, interpolated points are totally independent one another, except selected k -NN in the MDS interpolation algorithms, and the independency of among interpolated points makes the MDS interpolation algorithm to be pleasingly-parallel. In other words, there must be minimum communication overhead. Also, load-balance can be achieved by using modular calculation to assign interpolated points to each parallel unit, either between processes or between threads, as the number of assigned points are different at most one. Thus, we can parallelize MDS interpolation algorithms via not only the traditional MPI but also the emerging MapReduce [21, 22] runtimes.

Algorithm 1 Adaptive Interpolation of MDS algorithm

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1: Find  $k$ -NN: find  $k$  nearest neighbors  $p_i \in P$   $i = 1, \dots, k$  of a new data  $x$  based on  $\delta_{ix}$ .
2: Gather mapping results in target dimension of the  $k$ -NN.
3: Calculate  $\bar{p}$ , the average of pre-mapped results of  $p_i \in P$ .
4: Calculate  $\delta_{ix}$ , by Eq. (18).
5: Generate initial mapping of  $x$ , called  $x^{[0]}$ , either  $\bar{p}$  or a random variation from  $\bar{p}$  point.
6: Compute  $\sigma(S^{[0]})$ , where  $S^{[0]} = P \cup \{x^{[0]}\}$ .
7: while  $t = 0$  or  $(\Delta\sigma(S^{[t]}) > \varepsilon$  and  $t \leq \text{MAX\_ITER})$  do
8:   increase  $t$  by one.
9:   Compute  $x^{[t]}$  by Eq. (19).
10:  Compute  $\sigma(S^{[t]})$ .
11: end while
12: return  $x^{[t]}$ ;

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Table 1: Compute cluster systems used for the performance analysis

# Nodes	32
CPU	Intel Xeon E7450 2.4 GHz
# CPU / # Cores per node	4 / 24
Total Cores	768
Memory per node	48 GB
Network	20 Gbps Infiniband
Operating System	Windows Server 2008 HPC Edition (Service Pack 2) - 64 bit

6. Analysis of Experimental Results**6.1. Experimental Environment**

In this section, we provide some experimental analysis for the proposed AI-MDS algorithm. To explore the quality and the performance of the proposed AI-MDS approach discussed in this paper compared to MI-MDS [15], we have used 166-dimensional chemical dataset obtained from PubChem project database¹, which is a NIH-funded repository for over 60 million chemical molecules and provides their chemical structures and biological activities, for the purpose of chemical information mining and exploration. In this paper we have used randomly selected up to 4 million chemical subsets for our testing. The computing cluster system we have used in our experiments is demonstrated in Table 1.

6.2. Quality and Runtime Analysis

In this section, we would like to compare the mapping quality of the proposed AI-MDS to that of the MI-MDS algorithm as well as the running time of both algorithms. For the quality measurement, we use the normalized STRESS value (σ) with uniform weights ($w_{ij} = 1$) defined as in Eq. (20).

$$\sigma(X) = \frac{\sum_{i < j \leq N} (d_{ij}(X) - \delta_{ij})^2}{\sum_{i < j \leq N} \delta_{ij}^2} \quad (20)$$

The normalized STRESS will be ONE if all the points are configured at the same position. In this section, when we mention STRESS value, it means the normalized STRESS value.

¹PubChem, <http://pubchem.ncbi.nlm.nih.gov/>

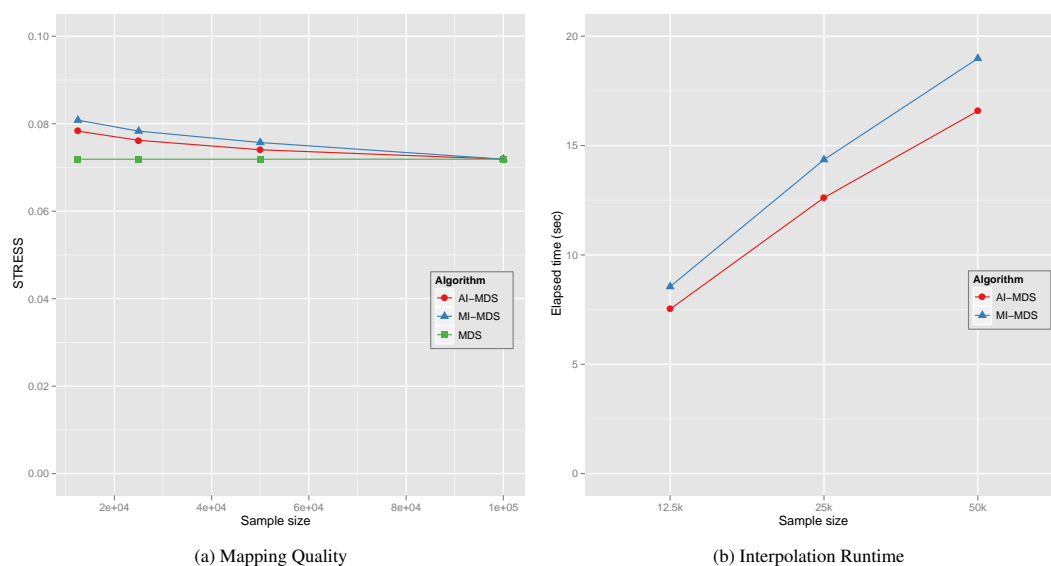


Figure 2: (a) Mapping quality comparison among full MDS, MI-MDS, and AI-MDS algorithm with 100k pubchem data set with respect to different sample sizes (n), i.e. 12.5k, 25k, and 50k, when $k = 2$, and (b) corresponding running time of MI-MDS and AI-MDS.

Fig. 2-(a) illustrates the mapping quality of the AI-MDS results and the MI-MDS results of 100,000 points (hereafter $100k$) (N) with respect to the different sample sizes (n). As shown in Fig. 2-(a), the proposed AI-MDS performs better than the MI-MDS in terms of the mapping quality. In detail, if we define the quality degradation of MI-MDS ($\sigma_{full} - \sigma_{MIMDS}$) is equal to 1.0, then the quality degradation of AI-MDS ($\sigma_{full} - \sigma_{AIMDS}$) is 71.9%, 67.2%, and 55.4% of that of MI-MDS with respect to $n = 12.5k$, 25k, and 50k, correspondingly. In other words, if we assume that the possible best quality of an MDS result via an interpolation approach will be the quality of full MDS running, the proposed AI-MDS improves the mapping quality about 28.1%, 32.8%, and 44.6% compared to the MI-MDS algorithm with the test dataset.

We have also compared the interpolation running time of the MI-MDS and the proposed AI-MDS algorithms corresponding to Fig. 2-(a), which is demonstrated in Fig. 2-(b), and it shows very interesting result. We expect the runtime of the proposed AI-MDS algorithm could be taken slightly longer than or compatible to that of the MI-MDS, since the distance adapting step is added for the interpolation procedure of each point. However, the runtime analysis in Fig. 2-(b) shows in the opposite to our expectation. The AI-MDS runs faster than the MI-MDS for all test cases. Thus, we have also looked into detail of the number of iterations for interpolation of each point by the MI-MDS and the AI-MDS with 50k sample of 100k full data set. The average of the iteration numbers by the AI-MDS is 1.731, and the average of the iteration numbers by the MI-MDS is 1.834. We could understand this phenomenon as the search space of interpolation for each point becomes more suitable via adapting the dissimilarity.

In addition to the experiments of the fixed full data set in the above analysis, we have also tested the proposed AI-MDS with larger *out-of-sample* size cases, i.e. millions of points, as shown in Fig. 3. For the large data test, we randomly selected 100k pubchem data among over 60-million compounds data as an *in-sample* set, then we have tried to interpolate 1 million, 2 millions, and 4 millions (hereafter $1M$, $2M$, and $4M$, correspondingly) chemical compounds data, which are also randomly selected from the same dataset.

As shown in Fig. 3-(a), the proposed AI-MDS outperforms the MI-MDS method in terms of the mapping quality. The full MDS is infeasible to generate a mapping of millions of points as the motivation of MDS interpolation approach [15]. However, we could still compare the mapping quality of those interpolation approach with respect to the normalized STRESS value of the sample mapping, on the basis of the assumption that the sample mapping quality could be similar to the full MDS mapping result of all points. If we define the quality degradation of the MI-MDS and the AI-MDS as same as the previous experiment, the quality degradation of the AI-MDS is only about 56.7%, 57.5%, and 57.7% compared to the quality degradation of the MI-MDS with 100k sample mapping and $1M$, $2M$, and

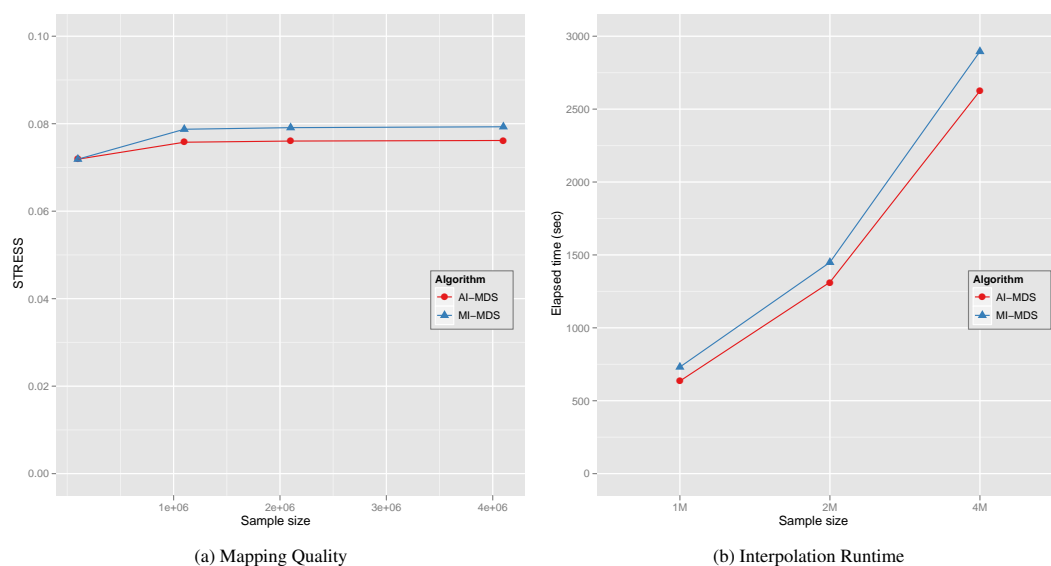


Figure 3: (a) Mapping quality comparison between MI-MDS and AI-MDS algorithm with respect to large out-of-sample size (M), i.e. 1M, 2M, and 4M, with 100k sample pubchem data when $k = 2$, and (b) corresponding running time of MI-MDS and AI-MDS.

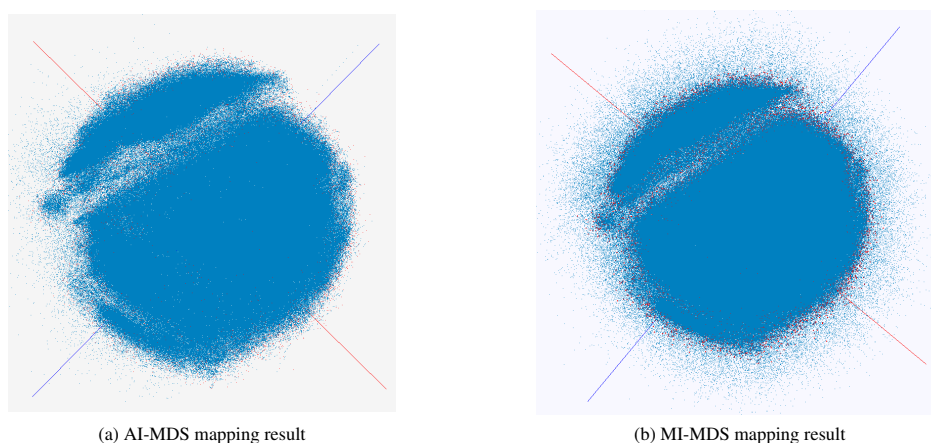


Figure 4: Interpolation Mapping results of 2M compounds by (a) AIMDS and (b) MIMDS with 100k sample data, when $k = 2$.

4M out-of-sample data, respectively, which means the AI-MDS improves the quality of interpolation more than 40% compared to the MI-MDS.

Relating to the runtime of those algorithms, the AI-MDS is about 10% faster than the MI-MDS as shown in Fig. 3-(b), which is consistent to Fig. 2-(b). The runtime results in Fig. 2-(b) and Fig. 3-(b) are the running times of both the AI-MDS and the MI-MDS algorithms in hybrid parallel method by using 384 cores of the system in Table 1.

Fig. 4 illustrates the actual interpolation mapping result of 2M out-of-sample points (shown in blue color) based on 100k prior mapping (which is represented in red color) via (a) the AI-MDS and (b) the MI-MDS methods. As shown in Fig. 4-(b), the MI-MDS result has points which are configured in the outside of the boundary of the prior mapping much more than the AI-MDS result in Fig. 4-(a). We could interpret those outside mappings in Fig. 4-(b) are affected by the distance discrepancy between the original dissimilarity of each interpolated point and its k -NN and the established mapping space in the target dimension by the sample mapping.

7. Conclusion

Majorizing interpolation method for multidimensional scaling (MI-MDS) was proposed for the purpose of configuring millions of points via commodity cluster systems based on the prior mapping of sample data [15]. As in [15], the MI-MDS method produces mappings of millions of points, which is infeasible via normal MDS methods, in reduced computational complexity by the cost of mapping quality degradation. Although the quality of MI-MDS is acceptable, we have investigated how to improve the mapping quality of MI-MDS algorithm in [15].

In this paper, we propose an adaptive interpolation method of MDS (AI-MDS) which aims to improve the mapping quality of MI-MDS, based on the distance ratio between the original dissimilarities and the corresponding mapping distances in the target space. The proposed AI-MDS shows significant improvement of the mapping quality for the tested cases. For instance, the AI-MDS algorithm configures mappings of millions of out-of-sample data cases which are improved more than 40%, with regard to the quality degradation from the full MDS mapping quality, compared to corresponding mappings of the MI-MDS. Furthermore, the proposed AI-MDS generates better configuration of the tested data during faster running time than the MI-MDS. The average of iterations for all interpolated points via the proposed AI-MDS is less than via the previous MI-MDS method, interestingly.

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