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Manifold Matching using Shortest-Path Distance and Joint Neighborhood Selection

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ABSTRACT

Matching datasets of multiple modalities has become an important task in data analysis. Existing methods often rely on the embedding and transformation of each single modality without utilizing any correspondence information, which often results in sub-optimal matching performance. In this paper, we propose a nonlinear manifold matching algorithm using shortest-path distance and joint neighborhood selection. Specifically, a joint nearest-neighbor graph is built for all modalities. Then the shortest-path distance within each modality is calculated from the joint neighborhood graph, followed by embedding into and matching in a common low-dimensional Euclidean space. Compared to existing algorithms, our approach exhibits superior performance for matching disparate datasets of multiple modalities.

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1. Introduction

The abundance of data in the modern age has made it crucial to effectively deal with large amounts of high-dimensional data. For the purpose of data analysis, it is imperative to apply dimension reduction to embed data into a low-dimensional space for subsequent analysis. Traditional linear embedding techniques have solid theoretical foundations and are widely used, e.g., principal component analysis (PCA) [21, 43] and multidimensional scaling (MDS) [44, 5, 7] for datasets of a single modality, and canonical correlation analysis (CCA) [19, 1] for datasets of multiple modalities.

However, real datasets often exhibit nonlinear geometry, discovering which can be advantageous for subsequent inference. Many manifold learning algorithms have been proposed to learn the intrinsic low-dimensional structure of nonlinear datasets, including Isomap [42, 37], locally linear embedding (LLE) [30, 29], Hessian LLE [9], Laplacian eigenmaps [2, 17], local tangent space alignment (LTSA) [50, 49], among many others. Most of them start with the assumption that the data are locally linear, explore the local geometry via the nearest-neighbor

graph of the sample data, transform the data using the neighborhood graph, and eventually learn the low-dimensional manifold by optimizing some objective function. These nonlinear embedding algorithms usually serve as a preliminary feature extraction step that enables subsequent inference. They have been used successfully in object recognition and image processing.

In this paper, we consider the manifold matching task for datasets of multiple modalities, which is traditionally modeled by multiple dependent random variables. Conventional methods for identifying the relationship among multiple random variables are still very popular in theory and practice, such as canonical correlation [19, 22, 16] and Procrustes transformation [35, 36, 13, 14]. However, it has become a much more challenging task to match real datasets of multiple modalities from disparate sources due to their complex dependency structures, such as the same document in different languages, an image and its descriptions, or networks of the same actors on different social websites.

There have been many recent endeavors regarding data fusion and manifold matching [24, 48, 47, 32, 28, 39, 34]. Similar to dimension reduction for datasets of a single modality, manifold matching can serve as a feature extraction step to explore datasets of multiple modalities, and has also been shown to help subsequent inference in object recognition [23], information retrieval [38], and transfer learning [27]. Furthermore, the match-

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ing task is important on its own and has been applied to explore multiple graphs and networks [26, 46, 25]. One such application is seeded graph matching, where two large networks are collected but only a percentage of training vertices have known correspondence. Then the remaining vertices need to be properly matched to uncover potential correspondence and benefit later inference.

Due to the success of nonlinear embedding algorithms for datasets of a single modality, it is often perceived that these algorithms can be directly combined into the matching framework to improve the matching performance when one or more modalities are nonlinear. A naïve procedure is to pick one nonlinear algorithm, apply it to each modality separately, and match the embedded modalities. But such a simplistic procedure does not always guarantee a good matching performance, since many nonlinear embedding algorithms only preserve the local geometry up to some affine transformation [12]. Furthermore, using nonlinear transformations separately can even deteriorate the matching performance when compared to using simple linear transformations, as shown in our numerical simulations

To tackle the problem, we propose a manifold matching algorithm using shortest-path distance and joint neighborhood selection. By utilizing a robust distance measure that approximates the geodesic distance, and effectively combining the correspondence information into the embedding step, the proposed algorithm can significantly improve the matching quality from disparate data sources, compared to directly take linear or nonlinear embeddings for matching. All code and data are available on our website ¹.

2. Manifold Matching

2.1. The Matching Framework

Suppose n objects are measured under two different sources. Then $X_l = \{x_{il}\} \in \Xi_l$ for l = 1, 2 are the actual datasets that are observed / collected, with $x_{i1} \sim x_{i2}$ for each i (\sim means the two observations are matched in the context). Thus X_1 and X_2 are the two different views / modalities of the same underlying data. This setting is extendable to datasets of more than two modalities, but for ease of presentation we focus mainly on the matching task of two modalities.

 Ξ_1 and Ξ_2 are potentially very different from each other, such as a flat manifold and its nonlinear transformation, an image and its description, or texts under different languages. A typical example is the social network, where many users have accounts on Youtube, Facebook, Twitter, etc. People sometimes post different contents and connect with different groups on each network site, such that data analysis of better quality is only possible when multiple accounts of the same person are combined. Some users already linked their accounts from different places, or unique user information are filled (like actual name, occupation), certain accounts can be automatically matched, providing a set of matched training data; but all the other accounts need to

be matched by machine (as manual match is too expensive for millions of accounts), presenting a set of testing data from each website.

We assume $x_{il} \in \Xi_l$ is endowed with a distance measure Δ_l such that $\Delta_l(i,j) = dist(x_{il},x_{jl})$. To match multiple modalities, we find two mappings $\rho_l : \Xi_l \to \mathbb{R}^d, l = 1,2$ such that the mapped data $\hat{X}_l = \{\rho_l(x_{il})\}$ are matched into a common low-dimensional Euclidean space \mathbb{R}^d . A simple example of ρ_l is MDS (e.g., classical MDS first doubly centers the distance matrices, followed by eigen-decomposition and keeping the top d eigenvalues and eigenvectors to yield the embedding) followed by CCA (find two orthogonal $d \times d$ transformation matrices for each data set to maximize their correlation), which is a linear embedding and matching procedure.

Once the mappings are learned from the training data, the learned mappings ρ_l can be applied to match any new observations $y_1 \in \Xi_1$ and $y_2 \in \Xi_2$ of unknown correspondence, i.e., compute $\hat{y}_l = \rho_l(y_l) \in \mathbb{R}^d$, and declare y_1 and y_2 as matched if and only if \hat{y}_1 is sufficiently close to \hat{y}_2 in the common space. Ideally, a good matching procedure should be able to correctly identify the correspondence of the new observations, i.e., if the testing observations are truly matched in the context, the mapped points should be very close to each other in \mathbb{R}^d . If the testing observations are not matched, the mapped points should be far away from each other.

To evaluate a given matching algorithm, a natural criterion is the matching ratio used in seeded graph matching [26]. Assume that there exist multiple testing observations in each space; and for each testing observation y_1 in Ξ_1 , there is a unique testing observation $y_2 \in \Xi_2$ such that $y_1 \sim y_2$. Then they are correctly matched if and only if \hat{y}_2 is the nearest neighbor of \hat{y}_1 among all other testing data from Ξ_2 , and vice versa. The matching ratio equals the percentage of correct matchings, and a higher matching ratio indicates a better matching algorithm.

The matching ratio based on nearest neighbor is often conservative, and can be a very small number when matching disparate real datasets. In practice, it is often more interesting to consider all neighbors within a small threshold, or rank multiple neighbors up to a limit. To that end, the testing power of the statistical hypothesis $H_0: y_1 \sim y_2$ considered in [28] is another suitable criterion, which directly takes the Euclidean distance $\|\hat{y}_1 - \hat{y}_2\|$ as the test statistic. To estimate the testing power for given data, we first split all observations into matched training data pairs, matched testing data pairs, and unmatched testing data pairs. After learning ρ_l from the matched training data and applying them to all testing data, the test statistic under the null hypothesis can be estimated from the matched testing pairs, and the test statistic under the alternative hypothesis can be estimated from the unmatched testing pairs. The testing power at any type 1 error level is directly estimated from the empirical distributions of the test statistic, and a higher testing power indicates a better manifold matching algorithm.

We used both the testing power and the matching ratio for evaluation in the numerical experiments, and in most cases they yield the same interpretation regarding which method has a better matching performance. Note that if the critical value at a given type 1 error level is used as a distance threshold, the test-

¹https://github.com/cshen6/MMSJ

ing power equals the probability that the distance between the matched pair is no larger than the distance threshold. Since the matching ratio only considers the nearest neighbor of the matched pair, the testing power is never smaller than the matching ratio.

2.2. Main Algorithms

Our methodology is henceforth referred to as MMSJ. Algorithm 1 serves to learn the matching transformations from the matched training data, while algorithm 2 maps any testing observation onto the learned manifolds.

Given the distance matrices Δ_l for the training data $\{X_l, l=1,2\}$, we first construct an $n\times n$ binary graph G by k-nearest-neighbor using the sum of normalized distance matrices $\sum_{l=1}^2 \frac{\Delta_l}{\|\Delta_l\|_F}$, i.e., G(i,j)=1 if and only if $\sum_l \frac{\Delta_l(x_{il},x_{jl})}{\|\Delta_l\|_F}$ is among the smallest k elements in the set $\{\sum_l \frac{\Delta_l(x_{il},x_{ql})}{\|\Delta_l\|_F}, q=1,\ldots,n\}$. Next, for each modality X_l , we calculate the shortest-

Next, for each modality X_l , we calculate the shortest-path distance matrix Δ_l^G based on the normalized Δ_l and the joint graph G, i.e., solve the shortest-path problem using the weighted graph $\frac{\Delta_l \circ G}{\|\Delta_l\|_F}$, where \circ denotes the Hadamard product. Then we apply MDS to embed Δ_l^G into \mathbb{R}^d for each l, followed by the Procrustes matching to yield the matched data \hat{X}_l , i.e., the Procrustes matching finds a $d \times d$ rotation matrix by

$$P = \arg\min_{P'P=I} ||P\tilde{X}_1 - \tilde{X}_2||_F^2,$$

and sets $\hat{X}_1 = P\tilde{X}_1$ and $\hat{X}_2 = \tilde{X}_2$, where \tilde{X}_l denotes the embedded data by MDS.

Then each testing observation $y_1 \in \Xi_1$ is mapped as follows: Given the distance between testing and training $\Delta_1(y_1, X_1)$ and the shortest-path distances for the training data Δ_1^G , we first approximate the shortest-path distances $\Delta_1^G(y_1, X_1)$ by the respective nearest neighbors of the testing data within each modality. Then the testing data y_1 are embedded by MDS out-of-sampling (OOS) technique into \mathbb{R}^d to yield \tilde{y}_1 , followed by the Procrustes matching (i.e., $\hat{y}_1 = P\tilde{y}_1$ or $\hat{y}_2 = \tilde{y}_2$). Similarly for any $y_2 \in \Xi_2$.

To better visualize the process, we summarize the main algorithm in the flowchart of Figure 1.

2.3. Implementation Details

In this subsection, we discuss some implementation details of MMSJ.

MMSJ starts with two distance matrices rather than the sample observations directly, which means it is directly applicable to multiple modalities of different feature dimensions, as long as a distance metric can be defined for each modality. Although there is no limitation on applying the algorithm once the distances are given, the actual matching performance is clearly dependent on the choice of the metric. The most common choice is the Euclidean distance, or L^p metrics in general. Other similarity or dissimilarity measures may be more appropriate in certain domains, such as the cosine distance for text data (see Section 3.2), or suitable kernels for structured data [18].

The joint neighborhood graph ensures consistent neighborhood selection when at least one of the modalities are nonlinear or noisy, and is intuitively better than two separate neighborhood graphs for matching. Alternatively, one may use a

Algorithm 1 Manifold Matching using Shortest-Path Distance and Joint Neighborhood Selection (MMSJ)

Require: The distance matrices Δ_l for the matched datasets $\{X_l, l = 1, 2\}$, the neighborhood choice k, and the dimension choice d.

Ensure: The mapped datasets $\{\hat{X}_l \in \mathbb{R}^{d \times n}, l = 1, 2\}$, the shortest-path distance Δ_l^G , and the learned Procrustes transformation P.

```
1: function MMSJ(\Delta_1, \Delta_2, k, d)
              for i, j := 1, ..., n do G_{ij} \leftarrow \sum_{l} \frac{\Delta_{l}(x_{il}, x_{jl})}{\|\Delta_{l}\|_{F}} end for G = \text{Rank}(G) \Rightarrow rank distances within each row
 2:
 3:
              for i, j := 1, ..., n do G_{ij} \leftarrow I(G_{ij} \le k) end for
 4:
              for l := 1, 2 do
 5:
                     \Delta_l^G = \text{ShortestPath}(\frac{\Delta_l \circ G}{\|\Delta_l\|_F})
 6:
                     \tilde{X}_l^i = \text{MDS}(\Delta_l^G, d)
                                                                                  \triangleright embedding into \mathbb{R}^d
 7:
 8:
              [U,S,V] = \text{SVD}(\tilde{X}_2^T \tilde{X}_1)
 9:
              P \leftarrow UV^T
                                                                                ▶ Procrustes matching
10:
              \hat{X}_1 = P\tilde{X}_1
\hat{X}_2 = \tilde{X}_2
11:
12:
13: end function
```

Algorithm 2 MMSJ on Testing Data

Require: The distance vectors $\Delta_l(y_l, X_l)$ (either l = 1 or 2), the shortest-path distance matrices Δ_l and the mapped data \hat{X}_l , the learned Procrustes transformation P, and the neighborhood choice k.

```
Ensure: The mapped testing observation \hat{y}_l.
  1: function MMSJ2(\Delta_l(y_l, X_l), \Delta_l^G, \hat{X}_l, P, k)
  2:
             G_l = \text{Rank}(\Delta_l(y_l, X_l))
             \Delta_l^G(y_l, X_l) = \text{ShortestPath}([\Delta_l^G | \Delta_l(y_l, X_l) \circ G_l])
  3:
             \tilde{\mathbf{y}}_l = \text{MDS-OOS}(\hat{X}_l, \Delta_l^G(\mathbf{y}_l, X_l))
  4:
             if l = 1 then
  5:
                   \hat{y}_1 = P\tilde{y}_1
  6:
             else
  7:
  8:
                   \hat{y}_2 = \tilde{y}_2
  9:
             end if
10: end function
```

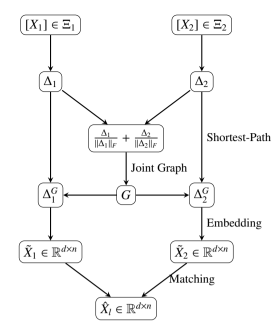


Fig. 1: Flowchart for Algorithm 1

weighted sum of distances or a rank-based method to derive the joint neighborhood graph instead. Note that joint neighborhood requires the distance matrices of the training data to be properly scaled in advance, but is clearly not applicable to the testing data with unknown correspondence.

Using the joint neighborhood graph allows the resulting shortest-path distance to utilize the dependency structure of the training data. Computationally, the shortest-path distance matrix can be effectively implemented by Floyd's algorithm or Dijkstra's algorithm [42], which can be further sped up by choosing a small set of landmark points [37, 3]. Theoretically, the shortest-path distance can recover the geodesic distance of isometric manifolds with high probability under certain sampling conditions [4, 37]. When embedding the testing data, we essentially treat the training data as landmark points and only compute the shortest-path distances from the testing data to the training data.

Embedding the shortest-path distances into the Euclidean space followed by matching is a standard procedure. Alternatively, one may match the embeddings by CCA or joint MDS, as discussed in [28, 11]. The advantages of MMSJ mostly lie in joint neighborhood and shortest-path distance; in fact, MMSJ always exhibits significant improvement, no matter which matching method to use. Thus we mainly consider the Procrustes matching for ease of presentation in the paper. For the testing data, they are embedded by out-of-sample MDS, which is a standard technique for MDS and kernel PCA [31, 3, 45] and more efficient than re-embedding all training and testing data. After all testing data are mapped onto the manifolds by the learned Procrustes matching, we may evaluate the matching performance as described in Section 2.1.

In terms of computation speed, suppose n is the sample size of training pairs, and n' is the size of all testing data. The running time complexity of MMSJ is $O(n^2 + nn')$, assuming the distance matrices are already given and the shortest-path

step uses the fast landmark approximation. The only overhead is the distance matrix construction, which takes an additional $O(n^2d + nn'd)$, where d denotes the maximal feature dimension among all modalities. Therefore MMSJ is computationally efficient for high-dimensional data or data with large amount of testing observations.

To compare with MMSJ, we use the standard procedure that embed each modality separately by MDS / Isomap / LLE / LTSA, followed by Procrustes matching. Note that MDS / Isomap / LLE can all operate directly on a distance matrix, but some nonlinear algorithms like LTSA have to start with the Euclidean data rather than a distance measure. Thus, if only the distance matrices are available, the distance matrices are preembedded into a Euclidean space $\mathbb{R}^{d'}$ with $d' \geq d$ by MDS, followed by embedding into \mathbb{R}^d via LTSA and matching by Procrustes.

3. Numerical Experiments

In this section, we demonstrate the numerical advantages of MMSJ. Overall, we observed that our method is significantly better than all the benchmarks (MDS, Isomap, LLE, and LTSA) in matching ratio and testing power under various simulated settings and real experiments.

3.1. Swiss Roll Simulation

The Swiss roll data from [42] is a 3D dataset representing a nonlinear manifold, which is intrinsically generated by points on a 2D linear manifold. Figure 2 shows the 3D Swiss roll data with 5000 points in colors, along with its 2D embeddings by MDS, Isomap, and LLE. Clearly, MDS fails to recognize the nonlinear geometry while both Isomap and LLE succeed. However, the LLE embedding has a distorted geometry, while the Isomap embedding is more faithful to the underlying 2D linear manifold.

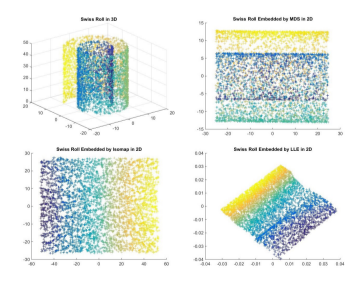


Fig. 2: The 3D Swiss roll dataset (top left), its 2D embedded data by MDS (top right), 2D embedding by Isomap at neighborhood size k=10 (bottom left), and 2D embedding by LLE at k=10 (bottom right).

For the first simulation, we match the 3D Swiss roll with its underlying 2D linear manifold at varying training data size. A total of n points from the 3D Swiss roll are randomly generated to construct the first modality X_1 , and the corresponding points on the underlying 2D linear manifold are taken as the second modality X_2 . Thus X_1 and X_2 are matched training data with distinct geometries. Once the training data are matched, we embed and apply the learned mappings to new testing observations y_1 and y_2 in each space.

We set the neighborhood size as k = 10, the dimension choice as d = 2, and generate n' = 100 testing pairs to compute the matching ratio. We repeat the above process for 100 Monte-Carlo replicates, and show the average matching ratio in Figure 3(A) with respect to increasing training data at $n = 50, 100, \ldots, 1000$. The MMSJ algorithm exhibits a significant advantage over all other algorithms: it enjoys a better matching ratio from small sample size onwards, and achieves almost perfect matching as sample size grows.

Next, we check the robustness of the manifold matching algorithms against noise, by adding white noise to the linear modality X_2 and fixing the training data size to n=1000. The noise is independently and identically distributed as $Normal(0, \epsilon I_{2\times 2})$, and the same testing procedure is applied to compute the matching ratio at increasing noise levels. The results are plotted in Figure 3(B) with respect to $\epsilon=0,1,2,\ldots,10$. The MMSJ algorithm is clearly better than all the benchmarks as the noise level increases.

For the third simulation, we consider an outlier scenario that randomly permutes a portion of the training data. For $\epsilon = 0, 0.01, \ldots, 0.1$, we randomly pick ϵn training data from X_2 and permute their indices, such that those training data are no longer matched with the corresponding observations from X_1 , i.e., there exists ϵn outliers in the training data. Fixing n = 1000, we apply the same testing procedure and plot the matching ratio in Figure 3(C) with respect to the outlier percentage ϵ . The MMSJ algorithm again has better matching ratio throughout increasing ϵ ; and all methods exhibit insignificant matching ratio when the outlier percentage ϵ reaches beyond 0.1, implying that the matching task may benefit significantly from excluding outliers prior to matching.

3.2. Wikipedia Articles Experiments

In this section, we apply the manifold matching algorithm to match disparate features of Wikipedia articles. The raw data contains 1382 pairs of articles from Wikipedia English and the corresponding French translations, within the 2-neighborhood of the English article "Algebraic Geometry". On Wikipedia, the same articles of different languages are almost never the exact translations of each other, because they are very likely written by different people and their contents may differ in many ways.

For the English articles and their French translations, a text feature and a graph feature are collected separately under each language. For the texts of each article, we use latent semantic indexing (LSI) (i.e., first construct a term-document matrix to describe the occurrences of terms in documents, then apply the low-rank approximation to the term matrix to 100 dimensions by singular value decomposition, see [8] for details) followed

by cosine dissimilarity to construct two dissimilarity matrices TE and TF (representing the English texts and French texts). For the networks, two shortest-path distance matrices GE and GF (representing the English graph and French graph) are calculated based on the Internet hyperlinks of the articles under each language setting, with any path distance larger than 4 imputed to be 6 to avoid infinite distances and scaling issues.

Therefore, there exist four different modalities for pairs of Wikipedia articles on the same topic, making TE, TF, GE, and GF matched in the context. Furthermore, as the text matrices are derived by cosine similarity while the graph matrices are based on the shortest-path distance with imputation, the former probably have nonlinear geometries while the latter are linear from the view of our matching algorithm.

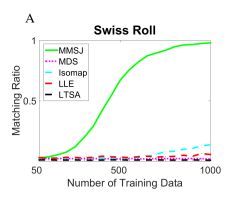
For each Monte-Carlo replicate, we randomly pick n = 500 pairs of training observations, 100 pairs of testing matched observations, and 100 pairs of testing unmatched observations for evaluation. The parameters are set as k = 20, d = 10, d' = 50 (for LTSA only), and the manifold matching algorithms are applied for every possible combination of matching two modalities. We perform a total of 100 Monte-Carlo replicates. The mean matching ratio is reported in Table 1, and the estimated testing power is presented in Table 2 at type 1 error level 0.05.

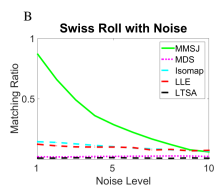
Clearly, MMSJ achieves the best performance throughout all combinations. From the tables and figures, we further observe that without using shortest-path distance or joint neighborhood, separate nonlinear embeddings from LLE or LTSA are worse than the linear MDS embeddings in matching. Isomap does fairly well in the testing power, as it also uses shortest-path distance, but it can still be occasionally similar or slightly inferior to MDS in the matching ratio. Our proposed MMSJ algorithm is consistently the best manifold matching algorithm in both the testing power and the matching ratio.

For the next experiment, we demonstrate that MMSJ algorithm is also robust against misspecification of parameters. The first two panels of Figure 4 plot the MMSJ and Isomap testing powers (the best two algorithms in our matching experiments) for matching (TE, GE), against different choices of $d \in [2, 30]$ and $k \in [10, 30]$. Comparing the two panels not only shows that MMSJ is always better than Isomap in matching, but it also attains comparable testing power in a wide range of parameter choices. The same robustness holds for MMSJ under all other matching combinations.

Table 1: Wikipedia Documents Matching Ratio

Modalities	MMSJ	MDS	Isomap	LLE	LTSA
(TE, TF)	0.2942	0.2546	0.2003	0.1265	0.0491
(TE, GE)	0.1209	0.0675	0.0866	0.0143	0.0260
(TF,GF)	0.0624	0.0419	0.0522	0.0134	0.0144
(GE,GF)	0.1347	0.1280	0.1081	0.0157	0.0236
(TE,GF)	0.0677	0.0429	0.0560	0.0132	0.0138
(TF, GE)	0.0946	0.0545	0.0698	0.0132	0.0238





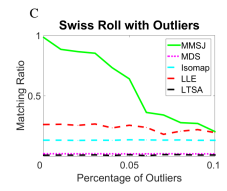


Fig. 3: Matching Ratio of 3D Swiss Roll versus its 2D Underlying Linear Manifold. (A) Matching Ratio with respect to Increasing Size of Training Data. (B) Matching Ratio with respect to Increasing Noise at n = 1000. (C) Matching Ratio with respect to Growing Number of Outliers at n = 1000.

Table 2: Wikipedia Documents Testing Power at Type 1 Error Level 0.05

Modalities	MMSJ	MDS	Isomap	LLE	LTSA
(TE, TF)	0.8124	0.4974	0.7476	0.3594	0.1930
(TE,GE)	0.5184	0.2563	0.4255	0.0948	0.1116
(TF,GF)	0.2782	0.1128	0.1877	0.0903	0.1028
(GE,GF)	0.3108	0.2141	0.2485	0.0961	0.1063
(TE,GF)	0.3199	0.1130	0.2141	0.0923	0.1021
(TF,GE)	0.4464	0.2114	0.3595	0.0943	0.1064

3.3. Brain Structural Networks

In this section, we assess the matching performance via brain structural networks. There are a total of n = 109 subjects, each with diffusion weighted magnetic resonance imaging (MRI) data. For the raw brain imaging data, we derived two different modalities. For each scan, (i) process diffusion and structural MRI data via MIGRAIN, a pipeline for estimating brain networks from diffusion data [15], (ii) compute the distance between brain networks using the semi-parametric graph test statistic [40, 41], then embed each graph into two dimensions and align the embeddings via a Procrustes analysis. The Euclidean distance is used on both modalities.

Therefore the first modality seems like a more faithful representation of the brain imaging, while the second modality is inherently a graph representation of the brain structure. Although these two modalities are merely different transformations of the same raw data, they are distinct in many aspects such that there is no guarantee that one can recover their underlying geometry or succeed the matching task via machine learning algorithms.

For each Monte-Carlo replicate, we randomly pick n pairs of matched observations for training, with all remaining sample observations for testing. The parameters are set as k=7, d=2, d'=2, with a total of 100 Monte-Carlo replicates. The mean matching ratio is shown in Figure 4(C) with respect to increasing size of training data. It is clear that MMSJ is the best matching method among all algorithms, and all matching ratios improve significantly as the training data size increases relative to the testing data size.

4. Concluding Remarks

In summary, we propose a nonlinear manifold matching algorithm using shortest-path distance and joint neighborhood selection. The algorithm is straightforward to implement, efficient to run, and achieves superior matching performance. It is able to significantly improve the testing power and matching ratio for multiple modalities of distinct geometries, and is robust against noise, outliers, and model misspecification. Our experiments indicate that the shortest-path distance and joint neighborhood selection are two key catalysts behind the improvement of the matching performance.

There are a number of potential extensions of this work. First, pursuing theoretical aspects of the manifold matching task is a very challenging but rewarding task: so far there is a very limited number of literatures even for manifold learning of a single modality, and no nonlinear transformation can always recover the linear manifold under a wide range of geometries. On the other hand, the task of matching multiple modalities is unique on its own. As a first step towards better theoretical understanding, we successfully proved in [33] that testing dependence via local correlations (which makes use of joint neighborhood graph and local distance in a similar manner) can successfully detect almost all relationships as sample size grows large, which shall shed more lights into the consistency of the matching task and may further advance the MMSJ algorithm.

Second, MMSJ requires a pair of metrics (or distances) for each modality. In this work we assume such metrics are predefined by domain knowledge, or use the Euclidean distance otherwise. If an optimal metric can be reasonably selected for each modality, it is likely to further boost the performance of MMSJ. From another point of view, if we are given high-dimensional or structured data (say graphs or images) to match, the algorithm may further benefit from first carrying out an appropriate feature selection down to certain landmark features [20, 6, 10], then use MMSJ. This quest is a valuable future direction to work on.

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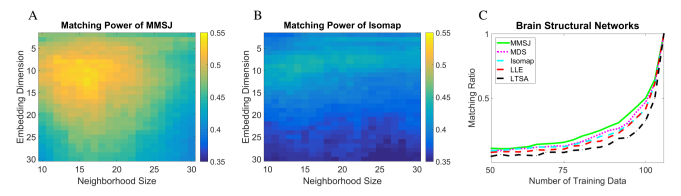


Fig. 4: (A) The Testing Power of MMSJ for Matching Wikipedia English Text and English Graph with respect to Different Dimension Choices and Neighborhood Sizes at Type 1 Error Level 0.05. (B) Same as (A) but for The Testing Power of Isomap. (C) Matching Ratio of Brain Structural Networks with respect to Increasing Size of Training Data.

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