A Deep Generative Model for Reordering Adjacency Matrices

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Abstract—Depending on the node ordering, an adjacency matrix can highlight distinct characteristics of a graph. Deriving a “proper” node ordering is thus a critical step in visualizing a graph as an adjacency matrix. Users often try multiple matrix reorderings using different methods until they find one that meets the analysis goal. However, this trial-and-error approach is laborious and disorganized, which is especially challenging for novices. This paper presents a technique that enables users to effortlessly find a matrix reordering they want. Specifically, we design a generative model that learns a latent space of diverse matrix reorderings of the given graph. We also construct an intuitive user interface from the learned latent space by creating a map of various matrix reorderings. We demonstrate our approach through quantitative and qualitative evaluations of the generated reorderings and learned latent spaces. The results show that our model is capable of learning a latent space of diverse matrix reorderings. Most existing research in this area generally focused on developing algorithms that can compute “better” matrix reorderings for particular circumstances. This paper introduces a fundamentally new approach to matrix visualization of a graph, where a machine learning model learns to generate diverse matrix reorderings of a graph.

Index Terms—Graph visualization, matrix visualization, machine learning, deep generative model, visualization interface.

1 INTRODUCTION

Graph-structured data are widely found in many disciplines; examples include protein-protein interaction networks in biological science, brain networks in neuroscience, and friendship networks in social science. Visualizing a graph allows people to intuitively understand and communicate the structural information in the data, which is especially helpful for non-experts in network analysis. A popular way to visualize a graph is visually encoding an adjacency matrix.

The rows and the columns of an adjacency matrix correspond to the nodes of a graph, where a matrix element $A_{i,j}$ indicates if there is an edge between node $u_i$ and node $u_j$. A view of an adjacency matrix is generally made by coloring an area mark for each element of the matrix based on its value [1]. Although node-link diagrams are more commonly used for visualizing graphs, adjacency matrix views are often preferred for dense graphs. Matrix views do not suffer from occlusion issues of node-link diagrams, such as node overlaps and edge crossings. Thus, adjacency matrix views can have higher perceptual scalability and information densities than node-link diagrams [1].

A node-link diagram of a graph can be made to display distinct structural characteristics of the graph depending on how the nodes are laid out in a two- or three-dimensional space. In an adjacency matrix view, nodes are laid out linearly in terms of their order along the rows (and the columns) of the matrix; and different node reorderings often highlight different structural aspects of the same graph (e.g., Fig. 1). Therefore, it is essential to find those “good” node reorderings that show the aspects of a graph that users wish to examine or present.

Decades of research have introduced many matrix reordering methods, which sort the rows and the columns of a matrix to reveal the structural information in the data [2], [3], [4]. However, there is no “best” method, as each method follows a different set of heuristics that can reveal certain structures in a graph. Therefore, users typically try multiple reorderings of the given adjacency matrix using different methods until they find one that best meets a specific visualization goal. Unfortunately, this trial-and-error approach is time-consuming and haphazard, which is especially challenging for novices.

This paper presents a technique that builds an intuitive interface for users to effortlessly find the desired adjacency matrix views of a graph. The fundamental approach is similar to the one for node-link diagrams presented by Kwon and Ma [5]. We first collect a set of reordering examples of the adjacency matrix of the given graph. Then, we train a generative model that learns a latent space of diverse matrix reorderings of the given graph. Finally, based on a grid of samples from the learned latent space, we build a what-you-see-is-what-you-get (WYSIWYG) interface that enables users to intuitively depict diverse reorderings of the given adjacency matrix.

However, reordering an adjacency matrix is a discrete problem, while laying out a node-link diagram is a continuous problem. Thus, designing a neural network for matrix reordering faces unique challenges. Overall, we identify three key challenges and design a novel neural network architecture addressing said challenges:

- Slight differences in node positions are negligible in a node-link diagram. In the case of an adjacency matrix,
however, a single difference may no longer represent the same graph. To ensure the generated adjacency matrices represent the same structure as the input graph, we train the model to produce permutation matrices, which represent node orderings, and then permute the given adjacency matrix, not to approximate reordered adjacency matrices naively.

- Gradient-based optimization is the standard approach to training a neural network [6]. However, it is only applicable to differentiable functions. Since reordering a matrix is not a differentiable problem, we need a differentiable approximation. We employ a relaxed sorting operator (e.g., Sinkhorn [7] and SoftSort [8]) such that standard gradient-based optimization techniques can be used for the training.

- Depending on the structure of the given graph, some distinct node orderings may produce the same matrix reordering. Training a model to learn different solutions for the same output could lead to ineffective or inefficient results. We thus design the neural network architecture and the loss function to take the resulting reordered matrices into account but not how the nodes are ordered (permutation equivariant).

Our evaluation results show that the presented architecture is capable of learning a generative model that can visualize a graph in diverse matrix reorderings that are comprehensive and distinctive. This new capability will greatly aid those who rely on matrix visualization extensively in their study of complex network data.

2 Related Work

Designing a deep generative model for matrix reordering is related to matrix reordering methods, deep learning for sorting, and deep generative models.

2.1 Matrix Reordering

Matrix reordering is a problem of finding a “good” ordering of the rows and columns of a relational matrix (e.g., an adjacency matrix and a correlation matrix) to reveal structural information from a set of objects. This problem is also known as (or equivalent to) seriation, linear arrangement, and sequencing. Since there can be $O(n!)$ different permutations of a set of $n$ objects, a brute-force enumeration approach to finding a good node ordering is infeasible even for a graph with a few dozens of nodes.

Decades of research across various disciplines have introduced many methods for matrix reordering, where each method uses a different set of heuristics, such as traveling salesman problem solvers [9], spectral ordering (i.e., the order of the Fiedler vector) [10], [11], and hierarchical clustering [12], [13], [14]. Several studies [2], [3], [4], [15] have reviewed various matrix reordering methods and have shown that different methods can show distinct patterns in the same matrix.

Unfortunately, it is not clear how to derive a suitable reordering for the given dataset and analysis goal [16], [17]. Users typically first produce multiple reorderings of the given matrix using various methods and then select one that fits the given circumstances. Several techniques have been introduced to support users through such a trial-and-error process by providing interactive methods to steer matrix reordering algorithms [17], [18], [19], [20]. The goal of these techniques is to narrow down the search space. Therefore, multiple trials with continuous human intervention would be required to explore other possible reorderings of the same matrix. In contrast, our approach is to build a model that can produce diverse reorderings of the given matrix, not to narrow the search space. Moreover, we train such a model in a fully unsupervised manner, thus not requiring human intervention.

2.2 Deep Generative Models

Deep generative models, such as variational autoencoders [21] and generative adversarial networks [22], have shown great success in various domains, such as image generation [23], [24], text generation [25], [26], and music generation [27], [28].

In the field of data visualization, Kwon and Ma [5] recently introduced a deep generative model for visualizing a graph in node-link diagrams. Their approach is to learn a two-dimensional latent space of diverse layouts of a graph. Then, users use the learned latent space as a map of various node-link diagrams of the given graph, such that they can effortlessly find a layout they want.

Inspired by [5], this paper aims to build such a generative model for visualizing a graph as adjacency matrices. We identify unique challenges for designing a neural network to achieve the goal and introduce a novel encode-decoder architecture that can learn a latent space of diverse matrix reorderings.

2.3 Deep Learning for Sorting

Sorting is one of the most important problems in computer science. Matrix reordering is also a sorting problem, where the goal is to find an ordering that can uncover structural information in a relational matrix. For our goal, we need to train a neural network that can produce diverse reorderings of the adjacency matrix of the given graph. However, sorting is not differentiable with respect to the input. It is thus impossible to use end-to-end gradient-based optimization to train a neural network with exact sorting functions.

Recently, a number of relaxed (differentiable) sorting operators have been introduced to enable gradient-based learning of models that include a sorting process [7], [8], [29]. The outputs of these relaxed sorting operators are “soft” permutation matrices. For example, the output of the Sinkhorn operator [7] is a doubly stochastic matrix, where the sum of any row or any column is 1. In addition, the NeuralSort [29] and SoftSort [8] operators produce a unimodal row stochastic matrix, where any row has the sum of 1 and has a unique $\text{argmax}$.

Using soft permutation matrices computed by the relaxed sorting operators, we can train a neural network model with approximated solutions. Once training is completed, these relaxed sorting operators can output “exact” permutation matrices to produce actual solutions for evaluation. For instance, since an output of the NeuralSort [29] and SoftSort [8] operators is a unimodal row stochastic matrix, we can simply apply row-wise $\text{argmax}$ to get an exact
permutation matrix. For a model with the Sinkhorn operator [7], an exact permutation can be obtained by passing the output of Sinkhorn operator to a linear assignment problem solver (e.g., Hungarian method [30]).

We use these relaxed sorting operators for learning a deep generative model that produces diverse reorderings of the adjacency matrix of a graph. In the evaluation, we compare two relaxed sorting operators for our problem, namely Sinkhorn [7] and SoftSort [8].

3 Approach
This paper introduces a technique for users to intuitively find the desired matrix view of a graph. More specifically, the goal is to learn a deep generative model that can systematically produce multiple adjacency matrix views of a graph with diverse node orderings that can reveal different structural information of the same graph. This section describes the whole process of constructing a deep generative model for matrix reordering, from gathering training samples to designing the neural network architecture.

3.1 Collecting Training Data
In order to learn a high-quality generative model that can produce various matrix reorderings, we need to collect a large and diverse set of matrix reorderings of the given graph. We collect such a dataset using various matrix reordering methods, multiple measures for computing distances between nodes, and random initial node orderings.

As we discussed in Section 2.1, decades of research have led to many matrix reordering methods. The adjacency matrix of a graph can reveal different structural patterns depending on which method is used, as shown in Fig. 1. Thus, an obvious way to collect diverse reorderings of the given graph is using multiple reordering methods.

This paper focuses on a simple, undirected, unlabeled graph. Since an adjacency matrix of an undirected graph is a symmetric matrix, we can use a single node ordering for sorting both the rows and columns of the matrix. Fig. 2 shows an overview of the matrix reordering process. To reorder the adjacency matrix $A$ of an undirected graph, a matrix reordering method typically begins with a pairwise distance (or dissimilarity) matrix $D$, where $D_{ij}$ represents the distance between nodes $u_i$ and $u_j$, and $D_{ii} = 0$. Thus, depending on the measure of distance, we can get different reordering results with the same reordering method.

The shortest-path distance is one of the most common measures for calculating dissimilarity between nodes in a graph. Another common approach is considering the adjacency matrix of a graph as the feature vectors of nodes, where each row/column represents the corresponding node. Then, general distance (or dissimilarity) functions (e.g., Table 2) can be used to obtain the pairwise distance matrix, and the resulting matrix reorderings can vary greatly depending on the measure, as shown in Fig. 3. We include the shortest-path distance and various distance functions (e.g., Table 2) to collect diverse reordering results of a graph.

When two nodes have the same set of adjacent nodes, their adjacency vectors (i.e., a row/column of the adjacency matrix) are the same. In this case, the distance between the two nodes can be calculated as 0, if the distance function only considers their adjacency vectors (e.g., Euclidean distance). In order to distinguish nodes with the same set of adjacent nodes, self-loops are often added (i.e., set the diagonal elements of the adjacency matrix to be 1) before computing the pairwise distance matrix, so that the distance between two different nodes is greater than 0. The same reordering method with the same distance function can produce very different results depending on whether self-loops are added or not, as shown in Fig. 4. We use both the (raw) adjacency matrix $A$ and the adjacency matrix with self-loops $\hat{A}$ of the given graph when we compute the pairwise distances.

Lastly, several reordering methods often produce different results depending on the initial node orderings. Fig. 5 shows such an example, where the same reordering method with the same distance measure produces different reordering results just because of the initial orderings of the given adjacency matrix are different. Thus, we also include a number of random initial orderings when collecting a training dataset.

Obtaining a large collection of matrix reorderings can take a significant amount of time. However, the training process can begin as soon as we have a few dozen samples with mini-batch training schemes. Therefore, the model training...
A permutation of the nodes of a graph is said to be an automorphism if the permutation preserves the adjacency of each node. In other words, if the permutation $P$ is an automorphism, the permuted adjacency matrix $A_P = P A P^T$ is equal to the original adjacency matrix $A (0 = A_P - A)$.

When a graph has a nontrivial automorphism (i.e., non-identity automorphisms), multiple different permutations of the graph can produce an equal reordered adjacency matrix. Fig. 6 shows such an example, where two different permutations $P_1$ and $P_2$ of the karate graph produce the same reordered adjacency matrix $(0 = A_{P_1} - A_{P_2})$.

Hence, we need to design a neural network architecture and a loss function considering the fact that different permutations can produce an equal reordered adjacency matrix. However, finding all the permutations that produce the same reordered adjacency matrix is practically impossible since the graph automorphism problem belongs to the class NP of computational complexity, similar to the graph isomorphism problem. Therefore, the architecture and the loss function need to be permutation equivariant: they should not depend on how the nodes are permuted but only on the resulting reordered adjacency matrices. We thus train the model to reconstruct reordered adjacency matrices, not to reconstruct permutations (i.e., node orderings). We describe our design of the architecture and the loss function in more detail in Section 3.3.

### 3.3 Architecture

This section discusses an encoder-decoder architecture to learn a generative model that can produce diverse matrix reorderings of the given graph. The architecture follows the general framework of VAEs [21]. Fig. 7 shows an overview of the architecture.

#### 3.3.1 Encoder

The encoder’s goal is to learn the low-dimensional latent representation $z$ of a reordered adjacency matrix $A_P = \text{Reorder}(A, P) = P A P^T$ (Fig. 7a), where $A$ is the original adjacency matrix and $P$ is the permutation matrix that represents a node ordering. As we discussed in Section 3.2, we use a reordered adjacency matrix $A_P$ as the input object rather than a permutation $P$ because there can be different permutations that produce the same reordered adjacency matrix.

There is another intuition of using reordered adjacency matrices as the input objects: we can treat a reordered adjacency matrix as an image since it is a visual representation of the given graph designed to help the viewers comprehend the data. With this intuition, we design the encoder inspired by neural networks for learning a generative model of images [21], [34]. For example, Kingma and Welling [21] have designed a variational autoencoder using a multilayer perceptron (a fully-connected neural network), where the input images are converted as one-dimensional vectors before they are fed to the encoder. In addition, Radford et al. [34] used a convolutional neural network, which has been widely used for various computer vision tasks [6], for designing a generative adversarial network [22].

Whether we use a multi-layer perceptron (MLP) or a convolutional neural network (Conv), the fundamental goal of the encoder is to learn a highly compressed representation $z$ that captures the essence of a reordered adjacency matrix $A_P$, where each layer of the encoder gradually reduces the dimensionality of representations. In the evaluation (Section 4), we compare two encoder designs (MLP and Conv) for learning a generative model of matrix reordering.

We set the encoder to produce a two-dimensional representation of the input $A_P$. Similar to the latent space of node-link diagrams [5], a two-dimensional latent space is straightforward to map a grid of generated samples on the
3.3.2 Decoder

The decoder’s goal is to reconstruct the input objects from their latent representations—computed by the encoder. In general, the decoder is trained to reconstruct the input objects (e.g., images of handwritten digits [21]) directly from the trainable parameters (i.e., weights and biases) of the neural network. However, for an adjacency matrix, naively training the decoder to reconstruct an input (i.e., a reordered adjacency matrix $A$) from the decoder leads to producing an invalid adjacency matrix.

For general image data (e.g., landscapes, portraits, and animals), small errors in the generated results do not have a substantial effect on the viewers’ understanding of the image content. In the case of an adjacency matrix view, however, even a small reconstruction error can lead to producing an invalid visualization that can mislead the viewers about the data, where the graph structure represented by a reconstructed adjacency matrix is different from the structure of the given graph. Therefore, we need to ensure that the decoder produces adjacency matrices representing exactly the same structure as the given graph.

We design the decoder to generate a permutation matrix $P'$ and then reorder the given adjacency matrix $A$ with the generated permutation matrix $P'$. Specifically, the input $A_P$ is reconstructed as $A_P = \text{Reorder}(A, P') = P' A P^T$, where $P'$ is the output of the decoder and $A$ is the original adjacency matrix (Fig. 7c). In essence, the decoder learns to reconstruct the permutation matrix $P$ (or its automorphism) of the input matrix reordering $A_P = P A P^T$, not to directly reconstruct $A_P$.

For the decoder to produce a permutation matrix, we use a relaxed sorting operator that outputs permutation matrices, such as Sinkhorn [7] and SoftSort [8] (discussed in Section 2.3). A relaxed sorting operator is placed after the decoder’s layers with trainable parameters (SoftPerm Fig. 7b). We set the SoftPerm to produce “soft” permutation matrices during training to enable gradient-based learning and exact permutation matrices at the inference time. While a reconstructed adjacency matrix $A_P$ can be different from the input $A_P$, the decoder always produces a valid permutation matrix. Therefore, the decoder is guaranteed to generate adjacency matrices that represent the same structure as the given graph.

While both Sinkhorn [7] and SoftSort [8] operators produce a permutation matrix, they have several differences in terms of their computational complexity and the mathematical characteristic of produced “soft” permutation matrices. The Sinkhorn operator $S(X)$ over an $n \times n$ square matrix $X$ is defined as:

$$S^0(X) = \exp(X),$$

$$S^i(X) = \mathcal{T}_r \left( S^{i-1}(X) \right),$$

$$S(X) = \lim_{i \to \infty} S^i(X),$$

where $\mathcal{T}_r(X)$ and $\mathcal{T}_c(X)$ are the row- and column-wise normalization operators, respectively. On the other hand, the SoftSort operator takes an $n$-dimensional vector $h$ and produces a permutation matrix by:

$$\text{softmax} \left( -d\left(\text{sort}(h)1^T, 1h^T\right) / \tau \right),$$

where softmax is applied row-wise, $d$ is a differentiable semi-metric function, and $\tau$ is a temperature parameter for controlling the degree of relaxation. In addition, to get an exact permutation matrix at the inference stage, the Sinkhorn operator requires to solve a linear assignment problem. Contrastingly, the SoftSort operator only needs to apply $\arg\max$ row-wise. Therefore, the Sinkhorn operator is more computationally demanding than the SoftSort operator. We compare two decoder designs in the evaluation to investigate their differences in building a generative model of matrix reordering.

3.4 Training

The parameters of the encoder and the decoder are trained to minimize the reconstruction loss $L_X$ and the variational loss $L_Z$, as our architecture is a variational autoencoder [21]. For the variational loss $L_Z$, we use the sliced-Wasserstein distance [35], like the generative model for node-link diagrams [5].

The reconstruction loss $L_X$ quantifies the difference between the input matrix reordering $A_P = P A P^T$ and its reconstruction $A_P = P' A P^T$, where $A$ is the original adjacency matrix, $P$ is the permutation matrix that represents the input node ordering, and $P'$ is the permutation matrix produced by the decoder. We don’t consider the difference between the input node ordering $P$ and the generated node permutation $P'$ since there can be different orderings that produce the same matrix reordering of the input adjacency matrix, as we discussed in Section 3.2. We directly compare the difference between the corresponding elements of $A_P$.
The eight graphs used in the evaluation. \(|V|\): the number of nodes, \(|E|\): the number of edges, \(|A_P|\): the number of unique reorderings (Section 4.1), \(t\): the time it took to compute the training dataset in minutes.

| Name  | Type   | \(|V|\) | \(|E|\) | \(|A_P|\) | \(t\) | Source |
|-------|--------|--------|--------|---------|------|--------|
| karate| Social | 34     | 78     | 7,964   | 3.27 | [36], [37] |
| cat   | Brain  | 52     | 515    | 9,060   | 4.49 | [38]    |
| gd96c | Synthetic | 65     | 125    | 10,973  | 6.65 | [39]    |
| macaque| Brain   | 71     | 438    | 9,682   | 7.16 | [40]    |
| polbooks| Co-purchase | 105   | 441    | 10,303  | 11.15 | [36] |
| collab| Collaboration | 129   | 847    | 8,420   | 13.68 | [41] |
| ego-fb| Social | 148    | 1,692  | 10,193  | 16.15 | [42], [43] |
| jazz  | Collaboration | 198   | 2,742  | 10,212  | 27.58 | [36], [44] |

4 Evaluation

The primary goal of the evaluation is to validate whether the model can learn to produce diverse matrix reorderings of the given graphs, not just memorize the training samples. We perform quantitative and qualitative evaluations of the generative model for reconstructing unseen matrix reorderings (i.e., the test dataset).

4.1 Datasets

This evaluation uses eight real-world graphs and around ten thousand orderings per graph (Table 1). While the number of training samples could affect the model’s behaviors, this evaluation focuses on the effect of different architectures using a fixed number of training samples per graph. For each graph, we first collected 13,500 matrix reorderings using 25 distance measures between nodes (Table 2), 27 matrix reordering methods (Table 3), and 20 random initial orderings.

The 25 distance measures consist of the shortest-path distances between nodes, 12 distance measures (Table 2) based on the adjacency matrix (a row of adjacency matrix represents a node), and the 12 distance measures based on the adjacency matrix with self-loops (i.e., the diagonal elements of the matrix is 1). The 27 matrix reordering methods (Table 3) are selected from the R package seriation [45], which provides publicly available, robust implementations of multiple matrix reordering methods. Although other methods are available in the package, these methods could not compute reorderings of all the eight graphs in a reasonable amount of time (24 hours). The 20 random initial node orderings are used as some matrix reordering methods produce different results depending on the initial node ordering of the given graph (Fig. 5).

A reordering can be the same as the reverse of another. Therefore, a reordering is reversed if its reverse has a higher similarity with a reference reordering—a Spectral reordering is used in this evaluation.

To properly evaluate the generalization capability of a model, the reorderings in the test set should not be used for training the model. Thus, we extract unique reordered adjacency matrices from the 13,500 reorderings of a graph since there can be equal reorderings, as we discussed in Section 3.2. The numbers of unique matrix reorderings and the time it took to collect the training dataset of each graph are shown in Table 1. For the reconstruction evaluation, we perform 5-fold cross-validations, where 80% of unique reorderings of a graph are used for training, and the other 20% reorderings are used for testing. Extracting unique adjacency matrices is required only for the evaluation purpose; it is optional for using our technique in practice.

4.2 Architectures and Configurations

We compare four different architectures, which are the combinations of two encoder designs (MLP and Conv) and two decoder designs (Sinkhorn and SoftSort): MLP-Sinkhorn, MLP-SoftSort, Conv-Sinkhorn, and Conv-SoftSort. Fig. 8 shows the encoders and decoders in detail. The encoder architectures generally follow the ones for images. The MLP encoders and decoders use four fully-connected layers, where it takes the input reordering \(A_P\)—an \(n \times n\) matrix—as an input, and the SoftSort operator is more computationally efficient [8].
The 27 matrix reordering methods used for producing training data in the experiment. The method names follow the documentation of the R package seration [45].

ARSA: Minimize the linear seriation using simulated annealing [32].

TSP: Minimize the Hamiltonian path length using a traveling salesperson problem solver [9], [31].

RZE: Rank-two ellipse seriation method [33]

MDS-[Metric, Nonmetric, Angle]: Minimize stress of the linear order using multidimensional scaling (MDS) [46]. Three variants are used.

HC-[Average, Single, Complete, Ward]: Uses a linear order of the leaf nodes in a dendrogram obtained by hierarchical clustering [12]. Each variant uses a different clustering method.

GW-[Average, Single, Complete, Ward]: Hierarchical clustering methods with the leaf-node ordering method developed by Gravaeus and Wainer [14].

OLO-[Average, Single, Complete, Ward]: Hierarchical clustering methods with the optimal leaf ordering method by Bar-Joseph et al. [13].

VAT: Uses the order of node addition in Prim’s algorithm, which finds a minimum spanning tree [47].

Spectral-[σ, Norm]: Uses the order of the Fiedler vector [10], [11]. “Spectral” uses the (unnormalized) Laplacian matrix, and “Spectral-Norm” uses the normalized Laplacian matrix.

SPIN-[NH, STS]: Sorting Points Into Neighborhoods [48]. Two variants are used: “SPIN-NH” uses the neighborhood algorithm, and “SPIN-STS” implements the side-to-side algorithm [48].

QAP-[LS, 2Sum, BAR, Inertia]: Formulates reordering as a quadratic assignment problem. Each variant uses different heuristics, namely the linear seriation problem formulation (QAP-LS) [49], the 2-sum problem formulation (QAP-2Sum) [10], the banded anti-Robinson form (QAP-BAR) [50], and the inertia criterion (QAP-Inertia) [19].

$\mathbf{z}$ is the number of nodes of the given graph. Each layer learns a more compressed representation of the input $\mathbf{A}_P$ by gradually reduces the dimensionality of representations. From the first to third layers, the $i$-th fully-connected layer (FC in Fig. 8) takes an $[n/2^{(i-1)}]^{2}$-dimensional vector and outputs an $[n/2^{i}]^{2}$-dimensional vector representation of the input $\mathbf{A}_P$. For example, the first layer takes the $n^2$-dimensional representation of the input $\mathbf{A}_P$ and learns an $[n/2]^{2}$-dimensional representation. The last layer produces the two-dimensional latent representation $z$ of the input $\mathbf{A}_P$ from the $[n/8]^{2}$-dimensional representation of it.

Conv encoders use three convolution layers and one fully-connected layer, where the input reordering $\mathbf{A}_P$ is considered as an $n \times n \times 1$ tensor. From the first to third layers, the $i$-th convolution layer (Conv in Fig. 8) has the kernel size of $(7-2(i-1)) \times (7-2(i-1))$ and the output feature size of $16 \cdot 4^{(i-1)}$. For example, the second layer has the kernel size of 5 and the output feature size of 64. Each convolution layer is followed by layer normalization [51], ELU [52], and the adaptive max pooling. The $i$-th adaptive max pooling (Pool in Fig. 8) has the target size of $[n/2^{i}] \times [n/2^{i}]$. Thus, the first convolution layer takes the input $\mathbf{A}_P$ as an $n \times n \times 1$ tensor and outputs an $[n/2]^{2} \times [n/2]^{2} \times 16$ tensor. The last layer computes the two-dimensional latent representation $z$ from the output of the third convolution layer by reshaping the $[n/8] \times [n/8] \times 256$ tensor as a $256 \times [n/8]^{2}$-dimensional vector.

Sinkhorn decoders use four fully-connected layers and the Sinkhorn operator [7] for computing a permutation matrix. The fully-connected layers of this decoder design is similar to the reverse of the MLP encoder. The first fully-connected layer takes the two-dimensional latent representation $z$ and outputs an $[n/8]^{2}$-dimensional vector. From the second to fourth layers, the $i$-th fully-connected layer takes an $[n/2^{(5-i)}]^{2}$-dimensional vector and outputs an $[n/2^{(4-i)}]^{2}$-dimensional vector. For example, the fourth layer takes an $[n/2]^{2}$-dimensional vector and outputs an $n^2$-dimensional vector. The output of the fourth layer is reshaped as an $n \times n$ matrix and fed to the Sinkhorn operator, which produces a “soft” permutation matrix. For computing an exact permutation matrix after training, we apply the Hungarian method [30] on the output of the Sinkhorn operator.

SoftSort decoders use four fully-connected layers and the SoftSort operator [8]. While the input of the Sinkhorn operator is an $n \times n$ matrix, the input of the SoftSort operator is an $n$-dimensional vector. Thus, the SoftSort decoders can produce a permutation matrix with a significantly fewer number of parameters. In this evaluation, we designed the SoftSort decoders to have $8n$-dimensional hidden representations (see Fig. 8 for details). The fourth layer takes an $8n$-dimensional vector and outputs an $n$-dimensional vector, which is passed to the SoftSort operator for computing a permutation matrix. After training, an exact permutation matrix is obtained by applying the row-wise argmax operation to the output of the SoftSort operator.

All the architectures use layer normalization [51] and the exponential linear unit (ELU) [52] on every hidden layer. Batch normalization [53] is commonly used in the neural network architectures for computer vision tasks, especially with convolutional neural networks. However, our pilot study showed that the models with batch normalization often result in numerical instability. The elements of an adjacency matrix are binary, whereas the pixels of a natural image (e.g., landscape, portrait, or an animal) are often 24-bit RGB colors or 8-bit grayscale. Therefore, in our case, batch normalization can lead to numerical instability due to the lack of diverse values of a feature across the instances of a batch.

We use the Adamax optimizer [54] with a learning rate of 0.001 for all the models. In our pilot study, Adamax—a variant of Adam based on infinity norm—showed more stable training than the standard Adam with the $L^2$ norm. For the reconstruction loss $L_X$, we use the binary cross-entropy loss averaged over the elements of the adjacency matrix. We set $\tau$ to 1.0 for both the Sinkhorn and SoftSort operators. We train each model for 500 epochs.

### 4.3 Implementation

We used the R package seration [45] to compute reorderings for the datasets (Section 4.1). The models are implemented with PyTorch [55]. To generate the datasets and run the evaluation, we used a machine with an Intel i7-5960X (8 cores at 3.0 GHz) CPU and an NVIDIA Titan RTX GPU.
### 4.4 Quantitative Evaluation

To quantitatively evaluate the generalization capabilities, we compare four different architectures (Section 4.2) for each graph in terms of the average reconstruction error rate of the test dataset—around two thousand reorderings that are not used for training the models. Specifically, a trained model takes an input matrix reordering \( A_P \) that is not used for training the model, encodes it to the latent space, and then reconstructs a matrix reordering \( A'_P \) from the latent representation. The difference between an input matrix reordering \( A_P \) and its reconstruction \( A'_P \) of the test dataset quantifies the generalization capability of a generative model as it estimates how close the model can produce unseen matrix reorderings. For a single reconstruction, the error is measured by the ratio of the different elements of the input matrix \( A_P \) and its reconstruction \( A'_P \): \( \sum_{i,j} 1_{(A_P)_{i,j} \neq (A'_P)_{i,j}} / n^2 \). We repeat 5-fold cross-validations ten times and report the mean error rates to reduce the effects of fold assignments. We also compare the four architectures’ computational costs based on the training time and the peak GPU memory usage, which measures the GPU memory capacity required for training.

#### 4.4.1 Results

Fig. 9a shows the mean reconstruction error rates of the 10 trials of the 5-fold cross-validations. Overall, MLP-Sinkhorn shows the lowest error rates for all eight graphs. The results show two different rankings of the other three architectures, depending on the graph. For polbooks, ego-fb, and jazz, the models with MLP encoders show lower error rates than the ones using Conv encoders, where the ranking is as follows (from lowest to highest): MLP-Sinkhorn, Conv-Sinkhorn, MLP-SoftSort, and Conv-SoftSort. For the other graphs, the models using Sinkhorn decoders show lower error rates than the ones with SoftSort decoders: MLP-Sinkhorn, Conv-Sinkhorn, MLP-SoftSort, and Conv-SoftSort.

Fig. 9b shows the mean training time per epoch. For all graphs, training the models with SoftSort decoders took a significantly shorter time than those with Sinkhorn decoders. Given the same encoder, training the SoftSort decoders took about half the time it took to train the Sinkhorn decoders, on average, 50.2% shorter. Except for the largest graph (jazz), the MLP encoders took, on average, 13.3% shorter to train than the Conv encoders with the same decoder. Thus, for these graphs, the ranking in terms of the training time is as follows (from shortest to longest): MLP-SoftSort, Conv-SoftSort, MLP-Sinkhorn, and Conv-Sinkhorn. For jazz, training the MLP encoders took, on average, 26.7% longer than the Conv encoders given the same decoder: Conv-SoftSort (28.4 s), MLP-SoftSort (38.8 s), Conv-Sinkhorn (61.6 s), and MLP-Sinkhorn (71.8 s).

Fig. 9c shows the peak GPU memory usage for training each model, which measures the required GPU memory capacity. Given the same encoder, the models using SoftSort decoders use, on average, 36.9% less GPU memory than those using Sinkhorn decoders. For the graphs with less than 100 nodes (karate, cat, gd96c, and macaque), the MLP encoders use less GPU memory than the Conv encoders with the same decoder. The difference decreases as the number of nodes increases; it becomes the opposite for the other graphs with more than 100 nodes (polbooks, collab, ego-fb, and jazz), where the Conv encoders use less GPU memory than the MLP encoders given the same decoder.

#### 4.4.2 Discussion

For many computer vision tasks, including image generation [34], [56], convolutional neural networks (CNNs) generally outperform fully-connected neural networks (i.e., multilayer perceptrons), even though a CNN often has a significantly fewer number of trainable parameters. However, in this evaluation, the MLP encoders consistently show lower error rates than the Conv encoders when using the same decoder. A further study using explainable artificial intelligence techniques [57] is required to better understand
the main cause of the accuracy differences. However, this difference suggests that matrix reordering requires capturing global patterns of the images (i.e., reordered adjacency matrices), which is what CNNs are not good at as they process a local neighborhood at a time [58], [59].

While the MLP encoders are able to capture global patterns, they are not scalable in terms of the number of nodes, as shown in the GPU memory usage (Fig. 9c). Thus, a memory-efficient architecture is necessary to support larger graphs. For images, several non-local modules have been introduced to understand global image structures by capturing long-range dependencies between pixels [58], [59], [60]. Although general non-local modules also consume a significant amount of memory, there have been some attempts to simplify non-local modules for images [61], [62] to reduce memory usage. Moreover, we believe taking the sparsity of real-world networks (i.e., an adjacency matrix is often a sparse matrix) [63] into account could drastically decrease memory consumption.

The Sinkhorn decoders show lower error rates than the SoftSort decoders given the same encoder. This could be simply because the Sinkhorn decoders have more trainable parameters than the SoftSort decoders. However, the experiments of Grover et al. [29] have shown that the NeuralSort operator [29]—which shares many characteristics with the SoftSort operator [8]—outperforms the Sinkhorn operator for sorting tasks, although the models using the NeuralSort operator have a significantly smaller number of parameters than those with the Sinkhorn operator. While both the Sinkhorn and SoftSort operators produce permutation matrices, the Sinkhorn operator is designed for a general matching problem (e.g., align, canonicalize, and sort), whereas the SoftSort (and NeuralSort) operator is designed specifically for sorting the input values (i.e., argsort). Our encoder-decoder architecture and loss function are designed more like a matching problem (e.g., jigsaw puzzles [7]), where the goal is to produce a permutation matrix that matches the input matrix reordering. This design difference can affect the flow of gradients and thus causes the performance differences.

4.5 Qualitative Results

Overall, the MLP-Sinkhorn models show the lowest error rates for all the graphs (Section 4.4). Fig. 10 shows several test reconstructions of five different graphs (karate, cat, gd96c, macaque, and jazz) using the MLP-Sinkhorn models. The first three rows of reconstructions of Fig. 10 demonstrate that the MLP-Sinkhorn models can produce diverse styles of matrix reorderings of the given graph. Especially, the reconstructions in the first row are exact reconstructions: the models produce unseen matrix reorderings without any difference. Although exact test reconstructions do not guarantee that the model generates high-quality samples of a generative model, it does demonstrate the generalization capability of our models.

Further investigation of exact reconstructions also revealed that our architectures and loss function are indeed
Fig. 10. Test reconstruction results of the five graphs using the MLP-Sinkhorn models. For each pair of matrix reorderings, the left is test input, and the right is the reconstructed matrix reordering. The label of a pair describes how the input matrix reordering is created: matrix reordering method (Table 3), the type of the adjacency matrix (\(A\): the original adjacency matrix, \(\hat{A}\): the adjacency matrix with self-loop), and the distance metric (Table 2). The results are discussed in detail in Section 4.5. The color of the matrices and node-link diagrams represents the community structure of the graph [64], so the readers can easily compare different matrix reorderings.

permutation equivariant (Section 3.2 and Section 3.3). For the karate, collab, ego-fb, and jazz graphs, 99.5% of exact reconstruction cases have the generated permutation matrix \(P'\) that is not the same as the permutation matrix of the input matrix reordering \(P\). However, this ratio depends on the number of automorphisms of a graph. For other graphs, all the generated permutation matrix \(P'\) is the same as the permutation matrix of the input matrix reordering \(P\).

The bottom two rows of Fig. 10 show a number of reconstructions with higher losses (i.e., higher error rates). However, the reconstructions in the fourth row are perceptually similar, although they have relatively higher losses than the first three rows. These results show a limitation of our loss function: the per-cell loss can disagree with the human’s perceptual similarity, which is a known issue in other computer vision tasks [65]. We believe a perceptual loss function [65] can be used for improving the quality of the generated samples.

5 Usage Scenario

We discuss how a learned latent space can support users in designing effective matrix visualizations of the given graph. Browsing multiple matrix reorderings and selecting the desired one from an unsorted list often lead to tedious trials and errors. Since we train a model to construct a 2D latent space, a grid of generated samples can be created to show several representative matrix reorderings of the given graph that the model learned. Fig. 11 and Fig. 12 show the latent spaces of matrix reorderings for the karate, cat, and macaque graphs learned by MLP-Sinkhorn models. The sample grids provide an organized overview of diverse matrix reorderings of the given graph, where similar matrix reorderings are placed close to each other in the latent space.

Using a sample grid as a map of the latent space of diverse matrix reorderings, we build an intuitive interface, where users can directly set the latent variable \(z\) to produce different matrix reorderings and select one they want by simply pointing to a location in the latent space. For example, Fig. 11 shows a grid of \(8 \times 8\) samples throughout
Fig. 11. The latent space of matrix reorderings for the karate graph learned by a MLP-Sinkhorn model. The grid of samples (center) is generated by decoding $8 \times 8$ latent variables $z$ in $[-1, 1]^2$. Users can intuitively explore diverse matrix visualizations and select one they want by simply pointing to a location in the latent space (e–g). The heatmaps (a–d) show four quality metrics of $256 \times 256$ matrix reorderings: (a) $A$ (the original adjacency matrix) + Shortest-path distance (distance measure) + the number of anti-Robinson events (AR), (b) $A$ + Hamming + AR, (c) $A$ + Euclidean + Correlation, and (d) $\hat{A}$ (the adjacency matrix with self-loops) + Euclidean + Correlation. The brighter colors represent “better” values in terms of the corresponding quality metric [45]. See Section 4.5 for a detailed discussion.

When the user points the location (e) in the latent space, the latent variable $z$ is set to $(-0.746, 0.873)$. Then, the model produces the corresponding reordering shown in Fig. 11e on the right. Users can effortlessly produce different matrix reorderings by pointing to different locations. Pointing the location (f) generates a matrix reordering that highlights the highly-connected nodes of the graph: the blue and red cells that form lines in Fig. 11f on the right. From the location (g) in the latent space, users can produce a matrix reordering that can show a pattern that is not obvious in the other reorderings (e and f): the olive cells are the edges that connect the blue and red clusters.

Since a learned latent space is $\mathbb{R}^2$, we can create sample grids with different levels of granularity. For instance, a grid of a smaller number of samples can be used for showing more details of the generated adjacency matrices of a larger graph in a limited display area, as shown in Fig. 12. In addition, we can visualize a latent space as a heatmap with a large number of samples to support experts in matrix reordering (e.g., algorithm designer).

Multiple quality metrics have been introduced to measure the “goodness” of a matrix reordering [45]. Moreover, we can define different distance measures between the nodes of a graph (Section 3.1). Thus, finding the quality metric and distance measure that can quantify the desired quality of a matrix reordering for the given graph and analysis goal is not a trivial task.

Fig. 11a–d and Fig. 12a–h show several heatmaps that represent quality metrics of $256 \times 256$ matrix reorderings of the corresponding graph, where the samples from the brighter areas have “better” values in terms of the definition of each metric (e.g., lower loss values or higher merit values [45]). For example, Fig. 11a is a map of the number of anti-Robinson events [33, 66] using the shortest-path distances between nodes, where the areas with brighter colors have a smaller number of anti-Robinson events. This metric could find a matrix reordering like Fig. 11g, which can better reveal the olive edges between the blue and red clusters.

By visualizing the latent space with quality metrics, experts in matrix reordering can understand what patterns the given metric can capture or not with diverse and concrete samples. For example, we can clearly see the differences between (1) different quality metrics (Fig. 12a and b, and Fig. 12c and d), (2) different distance measures (Fig. 11a and b, and Fig. 11c and d), and (3) different combinations of quality metrics and distance measures (Fig. 12e–h). Thus,
these heatmaps can support the process of designing a new quality metric or distance measure.

In summary, the learned latent spaces support users in designing effective matrix visualizations of a graph by providing a map of diverse matrix reorderings. A further evaluation similar to the user study of [67] could quantify the quality of the latent space by comparing the proximity in the latent space and the perceptual similarity of generated matrix reorderings.

6 Conclusion

Representing a graph as an adjacency matrix is a popular approach to graph visualization in addition to using a node-link diagram. However, different node orderings can reveal varying structural characteristics of the same graph. Therefore, finding a “good” node ordering is an important step for visualizing a graph as adjacency matrix views. Users generally rely on a trial-and-error effort to find a good node ordering.

In this paper, we have introduced a deep generative model that can learn a latent space of matrix reorderings of the given graph. The learned latent space is used as a WYSIWYG interface, where users can effortlessly explore diverse matrix reorderings and select the desired one for the purpose of visualization. To achieve this, we have designed a novel neural network architecture to address unique challenges in learning a deep generative model for matrix reordering.

A graph can be visualized in many different ways, where each visualization can highlight different characteristics of the same graph, leading to new insights and discoveries. We hope this paper encourages others to join a new area of graph visualization study, where the goal is to design a method that produces diverse visualizations of the same graph, not just a single visualization that follows only a certain set of heuristics.

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