Visualizing Riemannian data with Rie-SNE

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Abstract

Faithful visualizations of data residing on manifolds must take the underlying geometry into account when producing a flat planar view of the data. In this paper, we extend the stochastic neighbor embedding (SNE) algorithm to data on general Riemannian manifolds. We replace standard Gaussian assumptions with Riemannian diffusion counterparts and propose an efficient approximation that only requires access to calculations of Riemannian distances and volumes. We demonstrate that the approach also allows for mapping data from one manifold to another, e.g. from a high-dimensional sphere to a low-dimensional one.

1. Introduction

Visualizations are crucial to investigators trying to make sense of high-dimensional data. The most common visualization is a two-dimensional plot (e.g. on a piece of paper or a computer screen), so we rely on dimensionality reduction when handling high-dimensional data. Most dimensionality reduction techniques assume that data resides in an Euclidean domain, which presents a problem when data is not quite that simple. Data residing on Riemannian manifolds, such as the sphere, appear in many domains where either known constraints or other modeling assumptions impose a Riemannian structure. How should we visualize such data?

There are many concerns and questions when visualizing Riemannian data. The first is generic: all dimensionality reduction tools amplify parts of the signal while reducing the remainder. This inherent limitation should always be in mind when interpreting the visualization. Since some loss of information is inevitable, should we then loosen our grip on the data or its underlying Riemannian structure? Gauss’s Theorema Egregium [GP05] informs us that if the final plot is to be presented on a flat screen or piece of paper, then a distortion of the Riemannian structure is inevitable.

In practice, even if we accept the limitations of a visualization, actual algorithms for Riemannian data are missing. In this paper, we develop an extension of Stochastic Neighbor Embedding (SNE) [HR03] to Riemannian data and thereby provide one such tool. We call this Riemannian Stochastic Neighbor Embedding, or Rie-SNE for short. Our approach can embed data observed on one Riemannian manifold on another manifold. This allows for mapping data from a Riemannian space to a two-dimensional Euclidean plane (for plotting), but also mapping to a two-dimensional sphere, or similar when the Euclidean topology is inappropriate. Rie-SNE does not claim to solve the above-mentioned limitations of visualization-through-dimensionality-reduction, but it does provide a working tool with practical merit.

Figure 1: Rie-SNE visualizes high-dimensional Riemannian data by mapping to a low-dimensional Riemannian (or Euclidean) manifold. The figure shows high-dimensional spherical data mapped to either a low-dimensional sphere or a low-dimensional plane. The method also supports other manifolds, both as input and output.

2. Background and related work

Euclidean visualization. General data visualization is a vast topic, and a complete review is beyond our scope [Van16]. We here focus on the setting where data is represented as vectors (points) in an Euclidean space of high dimension. When data is two- or three-dimensional a scatter plot can directly reveal its structure, and we focus on the more difficult setting where data dimensions vastly exceed the easily plottable. Here one may explore the data through multiple projections, such as pairwise scatter plots, which are usually manageable for data of up to around 10 dimensions. Eventually, the approach tends to become unwieldy and the greater picture is lost. Alternatives include continuously interpolating between dimensions to provide an (interactive) animation [Asi85].

The approach we here explore is to find a non-linear mapping from the high-dimensional observation space into a two- or three-
dimensional space, which is suitable for plotting. Principal component analysis (PCA) [Jol02] is arguably the most commonly used approach to dimensionality reduction. PCA is restricted to spanning a linear subspace of the observation space, which often implies that the low-dimensional view reveals little structure. Several non-linear variants exist (e.g. [Law05, TSL00, RS00, HR03]), but they tend to be brittle [FHH21]. One of the currently most popular tools is t-SNE [vdMH08], which we review in Sec. 2.1.

Riemannian data. Data is often equipped with additional knowledge such as constraints or given smooth structures that give the data a Riemannian structure. For example, if all observations have unit norm, they reside on the unit sphere, which has a well-studied Riemannian manifold. The a priori available knowledge giving rise to a Riemannian data interpretation differ between domains, and we here only name a few. The most prominent example is that of directional statistics [MMJ00, KGH13, Hau18] where data resides on the unit sphere. Other examples include shape data [Ken84, FB12, SJML05, You12, KKG*12], DTI images [LDF04, PFA06, WSCV14], image features [TPM06, FHB14], motion models [TVSC11, cV09], human poses [SCLBS07, HSP12, HSP10, HLP11], robotics [GKJH15, JBS*22], social networks [MLLM*19] and more.

Even if Riemannian data is becoming increasingly common, tools for visualization have not followed. The most common approach for visualizing Riemannian data is to locate a point on the manifold, e.g. the intrinsic mean [Pen06], and map the data to its tangent space. That gives an Euclidean view of the data, which can be visualized with one of the many methods for this domain. Applying PCA tangentially is the gold standard [FLP04]. Unless data concentrates around the point of tangency this approach gives a highly distorted view of the data, and in practice, most knowledge reflected in the geometry is lost by the linearization [SLHNN10]. Several Riemannian PCA generalizations exist, e.g. [HHM10, PPY14, JDM12]. These generalize the classic linear model, and less work has been done on extending nonlinear methods. Two notable extensions are a ‘wrapped’ extension of the ear model, and less work has been done on extending nonlinearmost knowledge reflected in the geometry is lost by the linearization. Unless data concentrates around the point of tangency this approach gives a highly distorted view of the data, and in practice, with $p_{ij} = 0$. We can renormalize this to form a distribution over all $n$ observations

$$p_{ij} = \frac{p_{ji} + p_{i|j}}{2n}.$$  (2)

To learn a low-dimensional representation, the key idea is to repeat the above over the low-dimensional space to form

$$q_{ij} = \frac{s_{low}(y_i, y_j)}{\sum_{k} s_{low}(y_i, y_k)}, \quad q_{ij} = \frac{q_{ji} + q_{i|j}}{2n}. $$  (3)

We can now compare the similarity of our data and the representation with the Kullback-Leibler divergence between $p_{ij}$ and $q_{ij}$.

$$\text{KL}(P||Q) = \sum_{i=1}^{n} \sum_{j=1}^{n} p_{ij} \log \frac{p_{ij}}{q_{ij}}. $$  (4)

This can then be minimized using gradient descent with respect to the low-dimensional representation $(y_1, \ldots, y_n)$. In its classic form, SNE picks the measures of similarity as Gaussian functions

$$s_{high}(x_i, x_j) = \frac{1}{2\pi \sigma^2} \exp \left( -\frac{||x_i - x_j||^2}{2\sigma^2} \right),$$  (5)

$$s_{low}(y_i, y_j) = (2\pi)^{-d/2} \exp \left( -\frac{||y_i - y_j||^2}{2} \right). $$  (6)

With this choice, the normalization constants of Eq. 5 cancel out when computing $p_{ij}$ and $q_{ij}$. Note that this approach gives a per-observation variance $\sigma^2$, such that different points effectively can have different sizes of neighborhoods. To determine $\sigma^2$, the user specifies a perplexity parameter, which can be thought of as a measure of the effective number of neighbors [vdMH08],

$$\text{perplexity} = 2^{H(P_i)},$$  (7)

where $H(P_i)$ is the Shannon entropy [Sha48] of $P_i$. For a specific user-provided perplexity, we can perform a binary search over $\sigma^2$ such that Eq. 6 holds. In practice, the user experiments with different choices of perplexities to see which reveals a pattern.

### 2.1. Stochastic neighbor embedding

Stochastic neighbor embedding (SNE) [HR03] preserves similarity between neighboring points when mapped to a low-dimensional representation. Assume that we have access to functions $s_{high}$ and $s_{low}$, which measure the similarity between observation pairs in the high-dimensional observation space and the low-dimensional representation space, respectively. Now define the conditional probability, $p_{ji}$ that $x_i$ would pick $x_j$ as its neighbor [vdMH08]

$$p_{ji} = \frac{s_{high}(x_i | x_j)}{\sum_{k} s_{high}(x_i | x_k)},$$  (1)

The t-distributed stochastic neighbor embedding (t-SNE) is the most popular variant of SNE [vdMH08]. This is motivated by the ‘crowding problem’, where the low-dimensional representations significantly overlap without revealing much underlying structure. The idea is to use a similarity in representation space with heavy tails. Specifically, $s_{high}$ is a t-distribution with one degree of freedom centered around one representation,

$$s_{high}(x_i | x_j) = \pi^{-1} \left( 1 + ||y_j - y_i||^2 \right)^{-1}.$$  (7)
3. Method

Brownian motion on a Riemannian manifold. The key building block for generalizing SNE to Riemannian manifolds is a suitable generalization of the Gaussian distribution. Here we consider a density derived from a Brownian motion on a Riemannian manifold for high-dimensional probability computations.

The end-point \( x \) of Brownian motion in Euclidean space follows a Gaussian distribution. Infinitesimal increments of the Brownian motion can be projected onto the tangent space of corresponding points on a Riemannian manifold without error. Then, a Brownian motion starting at point \( \lambda \) running for some time \( t \) can be projected onto a \( D \)-dimensional Riemannian manifold. The resulting random variable can be interpreted as the probability that a Brownian motion starting at \( \lambda \) will end in point \( x \) on the manifold. This approximately has density [HS02, KEAH20]:

\[
B_{\lambda}(x \lambda, t) \approx (2\pi)^{-D/2} H_0 \exp \left(-\frac{\text{dist}^2(x, \lambda)}{2t}\right),
\]

where \( t > 0 \) is the duration of the Brownian motion (variance in Euclidean space), \( \lambda \) is the starting point of the Brownian motion, and \( H_0 \) is the ratio of Riemannian volume measures evaluated at points \( x \) and \( \lambda \) respectively. Superficially, Eq. 8 looks like the density of the normal distribution, with the Euclidean distance being replaced by its Riemannian counterpart. The normalization factor \( H_0 \), however, differs from the Euclidean distribution.

3.1. Rie-SNE

Rie-SNE works similarly as SNE and t-SNE, i.e. it will also produce two probability distributions \( P \) and \( Q \) from the data and aim to make them as similar as possible and in the process capture some underlying structure in the produced low dimensional embedding. However, computing the high-dimensional probability distribution \( P \) comes with an added cost. To preserve the Riemannian nature of the data, a different density is used when computing high-dimensional probabilities belonging to \( P \), namely the approximate density induced by the heat kernel of a Brownian motion on a Riemannian manifold given in Eq. 8, i.e. we pick

\[
s_{\text{high}}(x_j | x_i) = B_{\lambda}(x_j, x_i, t_i).
\]

The added computational cost is that the evaluation of \( B_{\lambda}(\cdot, \cdot) \) is more demanding than the conventional Gaussian similarity (5). Specifically, the normalization \( H_0 \) and the geodesic distance may be demanding, depending on the manifold on which the data resides.

With the Brownian motion model, we get

\[
p_{ji} \approx \frac{H_0[i, j] \cdot \exp \left(-\frac{\text{dist}^2(i, j)}{2t}\right)}{\sum_k H_0[i, k] \cdot \exp \left(-\frac{\text{dist}^2(i, k)}{2t}\right)},
\]

where we use the notations \( \text{dist}^2(i, j) = \text{dist}^2(x_i, x_j) \) and \( H_0[i, j] = \sqrt{\frac{\det G_s}{\det G_0}} \) to emphasize that these quantities can be pre-computed. As with SNE, we can optimize \( t_i \) to match a pre-specified perplexity using a binary search.

Choice of representation. As mentioned in Sec. 1, Gauss’s Theorema Egregium [GP05] informs us that we cannot isometrically embed data from a curved space into a space of different curvature without introducing distortion. Specifically, if we embed data from a non-linear manifold onto a flat two-dimensional representation (for plotting) then the curvature mismatch between spaces induces a distortion. This is a fundamental limitation that any visualization of Riemannian data will face, but we may nonetheless try to limit its impact. One approach is to embed the data onto a manifold of similar curvature as that of the manifold on which the data resides. For example, if the data resides on a high-dimensional sphere, it is perhaps more prudent to embed onto a two-dimensional sphere for plotting, rather than a Euclidean space.

With this in mind, we choose different distributions over the low-dimensional representation, depending on user preference.

- **Euclidean.** If the user prefers a Euclidean low-dimensional representation, we use a student-t in as regular t-SNE.
- **Spherical.** If the data manifold has positive curvature it may be beneficial to embed it on a sphere, in which case we opt to use a von Mises-Fisher distribution [MJM00]. Spherical embeddings have been explored elsewhere, e.g. [LCY19].
- **Other.** When the user wants to embed on another low-dimensional manifold, we suggest using the Riemannian Brownian over the low-dimensional representation.

Once we have defined both \( s_{\text{high}} \) and \( s_{\text{low}} \), we can estimate the representations using gradient descent just as regular SNE. Having performed \( T \) iterations (with a sufficiently large value for \( T \)) of the gradient descent, the two probability distributions \( P \) and \( Q \) will have a minimal KL-divergence resulting in near-optimal positions of the points in the low-dimensional embedding.

Implementation details. Our implementation of Rie-SNE relies on two approximation techniques that are traditionally also used when performing regular t-SNE. First, we compute the high-dimensional probabilities in \( P \) using a sparse nearest neighbor-based approximation technique [vdM14]. Second, we use the

![Figure 2: Two-dimensional Euclidean embeddings of spherical MNIST. Rie-SNE (left) discovers structure, which is lost on tangent space PCA (right).](image)
Barnes-Hut approximation \cite{BH86}, whenever we use a student’s t-distribution over the low-dimensional representation. This reduces the complexity to $O(n \log n)$.

### 4. Results

The performance of Rie-SNE is shown by comparing it to the gold standard of visualizing non-Euclidean data. This amounts to first computing the intrinsic mean, mapping all data to the tangent space at this point, and performing PCA over the tangential data.

**Spherical MNIST.** We start with the classic MNIST dataset consisting of $24 \times 24$ dimensional gray-scale images. We consider digits 0–5 to reduce clutter. To induce a non-Euclidean data geometry, we project the data onto the unit sphere of $\mathbb{R}^{24 \times 24}$ and denote the resulting data spherical MNIST. We visualize the resulting data using both Rie-SNE and tangent space PCA. First, we embed the data onto the plane, $\mathbb{R}^2$, and show the resulting plots in Fig. 2. Here it can be seen that Rie-SNE captures well the underlying relationship between the data points (the same digits are grouped), while tangent space PCA produces a cluttered view that does not reveal the underlying structure. Since the data has a spherical geometry, it may be beneficial to embed onto a low-dimensional sphere to preserve topology and curvature better. Figure 3 shows the Rie-SNE embedding onto $S^2$, where the clustering is again evident. As a baseline, the figure also shows a tangent space PCA embedding on $\mathbb{R}^3$. Although some structure can be captured with tangent space PCA here, Rie-SNE still gives better separation of the digit classes.

**Crypto-tensors.** Following Mallasto et al. \cite{MHF18} we consider the price of 10 popular crypto-currencies over the period 2.12.2014 — 15.5.2018. As is common in economy \cite{WG10}, the relationship between prices is captured by a $10 \times 10$ covariance matrix constructed from the past 20 days. This gives rise to a time series of covariance matrices, each of which resides on the cone of symmetric positive definite matrices. We provide visualizations of the data in Fig. 4. Rie-SNE is used to produce visualizations on both the plane $\mathbb{R}^2$ and the sphere $S^2$, showing a one-dimensional structure capturing the time evolution behind the data. In contrast, tangent space PCA produces $\mathbb{R}^2$ and $\mathbb{R}^3$ visualizations showing little to no structure in the embeddings.

### 5. Conclusions

We presented a new type of visualization technique, Rie-SNE, that is aimed at data residing on Riemannian manifolds, such as spheres. It is an SNE-based technique that can additionally produce different kinds of low-dimensional embeddings depending on user preference and the curvature of the original data manifold. We achieve favorable results using Rie-SNE compared to the gold standard tangential PCA. We hope that the present work also paves the way for other visualization tools for Riemannian data in order to support investigators relying on geometric models.

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