Looking at Non-Linear Dimension Reductions as Models in the Data Space

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Abstract

Nonlinear dimension reduction (NLDR) techniques such as tSNE, and UMAP provide a low-dimensional representation of high-dimensional (high-D) data using non-linear transformation. The methods and parameter choices can create wildly different representations, making it difficult to decide which is best, or whether any or all are accurate or misleading. NLDR often exaggerates random patterns, sometimes due to the samples observed. But NLDR views have an important role in data analysis because, if done well, they provide a concise visual (and conceptual) summary of high-D distributions. To help evaluate the NLDR we have developed an algorithm to show the 2D NLDR model in the high-D space, viewed with a tour. One can see if the model fits everywhere or better in some subspaces, or completely mismatches the data. It is used to evaluate which 2D layout is the best representation of the high-D distribution and see how different methods may have similar summaries or quirks.

Keywords: high-dimensional data, dimension reduction, hexagonal binning, low-dimensional manifold, tour, data vizualization, model in the data space

1 Introduction

Non-linear dimension reduction (NLDR) is popular for making a convenient low-dimensional (k-D) representation of high-dimensional (p-D) data. Recently developed methods include t-distributed stochastic neighbor embedding (tSNE) (van der Maaten & Hinton 2008), uniform manifold approximation and projection (UMAP) (McInnes & Healy 2018), potential of heat-diffusion for affinity-based trajectory embedding (PHATE) algorithm (Moon et al. 2019), large-scale dimensionality reduction Using triplets (TriMAP) (Amid & Warmuth 2022), and pairwise controlled manifold approximation (PaCMAP) (Wang et al. 2021). However, the representation generated can vary dramatically from method to method, and with different choices of parameters or random seeds made using the same method (Figure 1). The dilemma for the analyst is then, which representation to use. The choice might result in different procedures used in the downstream analysis, or different inferential conclusions. The research described here provides new visual tools to aid with this decision.

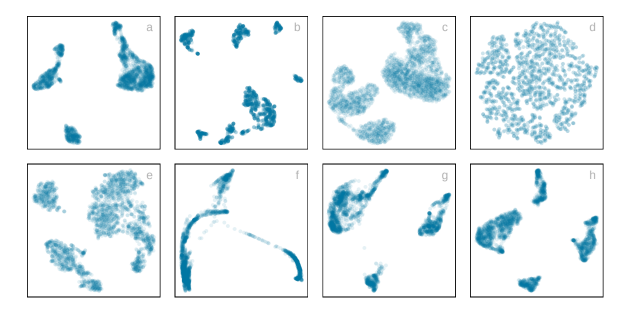


Figure 1: Six different NLDR representations of the same data. Different techniques and different parameter choices are used. Researchers may have seen any of these in their analysis of this data, depending on their choice of method, or typical parameter choice. Would they make different decisions downstream in the analysis depending on which version seen? Which is the most accurate representation of the structure in high dimensions?

The paper is organised as follows. Section 2 provides a summary of the literature on NLDR, and high-dimensional data visualization methods. Section 3 contains the details of the new methodology, including simulated data examples. Two applications illustrating the use of the new methodology for bioinformatics and image classification are in Section 4. Limitations and future directions are provided in Section 5.

2 Background

Historically, k-D representations of p-D data have been computed using multidimensional scaling (MDS) (Borg & Groenen 2005), which includes principal components analysis (PCA) (Jolliffe 2011) as a special case. The k-D representation can be considered to be a layout of points in k-D produced by an embedding procedure that maps the data from p-D. In MDS, the k-D layout is constructed by minimizing a stress function that differences distances between points in p-D with potential distances between points in k-D. Various formulations of the stress function result in non-metric scaling (Saeed et al. 2018) and isomap (Silva & Tenenbaum 2002). Challenges in working with high-dimensional data, including visualization, are outlined in Johnstone & Titterington (2009).

Many new methods for NLDR have emerged in recent years, all designed to better capture specific structures potentially existing in p-D. Here we focus on five currently popular techniques, tSNE, UMAP, PHATE, TriMAP and PaCMAP. tNSE and UMAP can be considered to produce the k-D minimizing the divergence between two distributions, where the distributions are modeling the inter-point distances. PHATE, TriMAP and PaCMAP are examples of diffusion processes (Coifman et al. 2005) spreading to capture geometric shapes, that include both global and local structure.

The array of layouts in Figure 1 illustrate what can emerge from the choices of method and parameters, and the random seed that initiates the computation. Key structures interpreted from these views suggest: (1) highly **separated clusters** (a, b, e, g, h) with the number ranging from 3-6; (2) **stringy branches** (f), and (3) **barely separated clusters** (c, d) which would **contradict** the other representations.

It happens because these methods and parameter choices provide different lenses on the interpoint distances in the data.

The alternative approach to visualizing the high-dimensional data is to use linear projections. PCA is the classical approach, resulting in a set of new variables which are linear combinations of the original variables. Tours, defined by Lee et al. (2021), broaden the scope by providing movies of linear projections, that provide views the data from all directions. Lee et al. (2021) provides an review of the main developments in tours. There are many tour algorithms implemented, with many available in the R package tourr (Wickham et al. 2011), and versions enabling better interactivity in langevitour (Harrison 2023) and detourr (Hart & Wang 2022). Linear projections are a safe way to view high-dimensional data, because they do not warp the space, so they are more faithful representations of the structure. However, linear projections can be cluttered, and global patterns can obscure local structure. The simple activity of projecting data from p-D suffers from piling (Laa et al. 2022), where data concentrates in the center of projections. NLDR is designed to escape these issues, to exaggerate structure so that it can be observed. But as a result NLDR can hallucinate wildly, to suggest patterns that are not actually present in the data.

The solution is to use the tour to examine how the NLDR is warping the space. This approach follows what Wickham et al. (2015) describes as model-in-the-data-space. The fitted model should be overlaid on the data, to examine the fit relative the spread of the observations. While this is straightforward, and commonly done when data is 2D, it is also possible in p-D, for many models, when a tour is used.

Wickham et al. (2015) provides several examples of models overlaid on the data in p-D. In hierarchical clustering, a representation of the dendrogrom using points and lines can be constructed by augmenting the data with points marking merging of clusters. Showing the movie of linear projections reveals shows how the algorithm sequentially fitted the cluster model to the data. For linear discriminant analysis or model-based clustering the model can be indicated by (p-1)-D ellipses. It is possible to see whether the elliptical shapes appropriately matches the variance of the relevant clusters, and to compare and contrast different fits. For PCA, one can display the k-D plane of the reduced dimension using wireframes of transformed cubes. Using a wireframe is the approach we take here, to represent the NLDR model in p-D.

3 Method

3.1 What is the NLDR model?

At first glance, thinking of NLDR as a model fitted to the data might seem strange. It is a model in the sense that it is a "a simplified representation or abstraction of a system, process, or phenomenon in the real world". The p-D observations are the realization of the phenomenon, and the k-D NLDR layout is the simplified representation. From a statistical perspective we can consider the distances between points in the k-D layout to be variance that the model explains, and the (relative) difference with their distances in p-D is the error, or unexplained variance. Abstractly, we can also imagine that the positioning of points in 2D represents fitted values, that will have some prescribed position in p-D that can be compared with the observed value.

Table 1 summarises the notation used to explain the new methodology. The observed data is denoted as $x_{n \times p}$ where x_{ij} would indicate the i^{th} observation on the j^{th} variable sampled from a population X. To refer to variable j, we would use X_i .

To illustrate the method, we use 7-D simulated data, which we call the "S-curve". It is constructed by setting $X_1 = \sin(a), X_2 = U(0,2), X_3 = \mathrm{sign}(a) \times (\cos(a) - 1), \forall a \in [-3\pi/2, 3\pi/2]$. The remaining variables X_4, X_5, X_6, X_7 are all uniform error, with small variance. We would consider $T = (X_1, X_2, X_3)$ to be the true model.

We define the NLDR model as a function that maps p-D to k-D, and takes some (hyper)parameters $\vec{\theta}$. That is,

$$g: \mathbb{R}^p \to \mathbb{R}^k$$
.

The parameter vector, $\vec{\theta}$, depends on the choice of g, and can be considered part of 'model fitting' in the traditional sense. Common choices for g include tSNE, uMAP, PACMAP, or PHATE, although in theory any function that maps the entire space \mathbb{R}^p to a subset of \mathbb{R}^k is suitable. Any input requirements for the data (such as normalisation, or preprocessing through the use of PCA or similar) is considered part of the function g.

Next, we create a model of the k-D representation of the data using hexagonal binning. The aim of this process is to reduce the noise in the original data that may be preserved

| Notation | Description |
|---|--|
| n, p, k | number of observations, variables, embedding dimension, respectively |
| $oldsymbol{X},oldsymbol{x}$ | p-dimensional data (population, sample) |
| y P T | k-dimensional layout orthonormal basis, generating a d -dimensional linear projection of p -dimensional data true model |
| $\begin{array}{c} y_{\max} \\ r_1, r_2 \\ h_b, w_b \\ a_r \\ h_r \end{array}$ | maximum value of the scaled second embedding component range of the first and second embedding component height and width of the hexagon aspect ratio of the 2D layout before scaling hexagon ratio |
| $egin{aligned} s \ (x_s,\ y_s) \ q_x,\ q_y \ h_s,\ v_s \end{aligned}$ | hexagonal size (radius of the outer circle surrounding the hexagon) starting coordinates of the hexagonal grid buffer amount along the x and y axes horizontal and vertical distance between adjacent hexagon bin centroids horizontal spacing and adjusted vertical spacing for hexagonal coordinates |
| $\begin{array}{c} b_{x},b_{y}\\ b,b'\\ n_{k}\\ (h_{x_{i}}^{k},h_{y_{i}}^{k})\\ C^{(j)} \end{array}$ | number of hexagon bins along the x and y axes total and non-empty hexagon bins in the grid number of observations within the k^{th} hexagon hexagonal grid coordinates of the k^{th} hexagon j -dimensional bin centers |

Table 1: Summary of notation for describing new methodology.

by g. Define a hexagonal grid across the k-D space, defined by the height and width of the hexagon (h_b and w_b respectively), the aspect ratio of the hexagon (h_r), and the starting coordinates of the hexagonal grid (x_s, y_s). We deliberately separate out the creation of the hexagonal grid from the mapping of points on the grid.

With the grid defined, the k-D points will belong to a hexagonal bin. That is, for each $y \in \mathbf{Y}$, we can (uniquely) identify the hexagon that the point belongs to. Let,

$$h: \mathbb{R}^k \to \mathbb{R}^k$$
,

be the function that maps a point in k-D space to it's hexagonal centroid. It follows that $h \circ g : \mathbb{R}^p \to \mathbb{R}^k$ maps each point in X to a k-D centroid.

The final step of the algorithm is lifting the k-D model back to p-D, completing the 'model-in-data-space' picture. For this, define the set H^k to be the set of points that belong to centroid k. That is, $H^k = \{y \mid g(y) = (h_x^k, h_y^k)\}$. We use the p-D Euclidean mean of the points in H^k to map the centroid (h_x^k, h_y^k) to a point in p-D. Let the i-th component of the p-D mean be

$$f_i = \frac{1}{|H^k|} \sum_{y \in H^k} y_i,$$

with corresponding p-D vector \vec{f}_i . Then, $f(h \circ g)(x)$ maps a p-D point to the p-D model estimate, completing the model cycle.

3.2 Displaying the model in p-D

3.3 Measuring the fit

3.4 What can be learned

- Overview: Generate a form that maps the model, that is the interpoint distances. What is the model?
- Notation
- Create a representation of the model
 - using hex-binning in 2D,
 - parameters,
 - tuning,
 - pre-processing
- How does this map to the representation in high-d
 - Centroids,
 - Edges

- Measuring fit
 - Fitted values
 - Error calculation
- What is learned about simulated examples
 - Interesting organisation of points in UMAP

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4 Applications

4.1 pbmc

- NLDR view used to illustrate clusters
- Use our method to assess is it a reasonable representation
- Demonstrate that it is not
- Illustrate how to use out method to get a better representation

4.2 digits: 1

- NLDR is used to illustrate different ways 1's are drawn
- Use our method to assess is it a reasonable representation
- Demonstrate that it is, except for the anomalies

5 Discussion

- Summarise contributions
- Explain where it is expected or not expected to work, eg higher dimensional relationships
- Human behaviour, the desire to have more certainty, and a tendency to prefer the well-separated views
- Diagnostic app to explore differences in distances
- What might be useful enhancements

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