## Seminar Report

# Semi-Supervised Locally Linear Embedding: Application and Sensitivity Analysis of Critical Hyperparameters

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## **Abstract**

#### Storyline

- Goal: present SS-LLE as a local, graph-based manifold learning method incorporating prior knowledge
- Step 0: define basic mathematical concepts required to understand argumentation (plus notation)
- Step 1: introduce idea of **isometry** (most basic: MDS)
- Step 2: introduce idea of graph-based models
  - Achieve non-linearity
  - $\circ$  Common structure: build graph  $\to$  derive matrix as quadratic form over graph function  $\to$  derive embedding from eigenvalue problem
  - Most basic: ISOMAP (global, dense, convex)
- Step 3: introduce idea of locality
  - Relax global to local isometry
  - Find sparse rather than dense matrices
  - Laplacian eigenmaps as concept in which the others can be generalized
    - Define weighting scheme for neighborhood
    - Use Laplacian to derive matrix
    - o Solve sparse eigenvalue problem
- Step 4: introduce local linearity
  - o LLE
    - Obtain weights via linear reconstructions
    - Can be shown to approximate graph Laplacian (Belkin & Niyogi (2006))
  - o Hessian LLE
    - Replace Laplacian by Hessian
- Step 5: introduce **prior knowledge** 
  - o SS-LLE
  - Improve results by pre-specifying some manifold coordinates

## **Extended Abstract**

#### CHECK AGAINST INTRODUCTION

The goal of this report is to lay out the theoretical framework behind the manifold learning technique of *semi-supervised locally linear embedding (SS-LLE)*, as proposed by Yang et al. (2006), and to put it to implementation for data sampled from manifolds.

Manifold learning in general is concerned with dimensionality reduction. As data analysis employs increasingly high-dimensional data, it is frequently necessary to scale down the number of features to ensure models work as desired and remain interpretable. Dimensionality reduction is justified by the assumption that data observed in D dimensions often truly lie on a d-dimensional manifold (d-manifold), i.e., the d-dimensional generalization of a curved surface, embedded in  $\mathbb{R}^D$  (with  $d \ll D$ ). As an example for this phenomenon one might consider image data showing objects in different poses. While images are typically stored in high-dimensional pixel representations, intuitively, it is in fact a very small number of features causing the variation in the data.

A crucial property of d-manifolds embedded in  $\mathbb{R}^D$  is their local topological equivalence to  $\mathbb{R}^d$ . This locally Euclidean behavior is exemplified by a sphere embedded in  $\mathbb{R}^3$ : although the sphere as a whole is entirely non-linear, on sufficiently small patches of its surface it behaves just like a flat plane in  $\mathbb{R}^2$ . It is precisely this fact that allows manifold coordinates to be mapped to  $\mathbb{R}^d$  in a reduction of dimensionality. The goal is now to learn this mapping in an unsupervised manner. Mapping manifold coordinates to  $\mathbb{R}^d$  is in general not equivalent to simple projection onto the d-dimensional coordinate hyperplanes. Instead, models must learn the intrinsic neighborhood structure of the manifold to establish a notion of "nearness" between points. As the sphere example demonstrates, standard distance metrics do not apply (globally) since points on general manifolds are connected by curved paths rather than straight lines.

Some manifold learning techniques try to retain global isometry by mapping pairwise distances to  $\mathbb{R}^d$ . For instance, multi-dimensional scaling (MDS) does so using Euclidean distances, thereby limiting the manifolds it can learn to linear ones, while ISOMAP generalizes this approach to non-linear manifolds by applying geodesic distances. Research indicates, however, that for non-convex manifolds it is more effective to preserve local structures only. Otherwise, solutions are prone to shortcuts, i.e., placing points close in  $\mathbb{R}^D$  next to each other when they lie in fact on quite different parts of the manifold. In order to avoid such miscalculations, sparse techniques focus on merely local neighborhood structures, modeled through weighted graph representations. The information from these graphs is then condensed into a sparse matrix. Eventually, the principal eigenvectors of this matrix yield the desired low-dimensional coordinates.

One such local graph-based technique is  $Laplacian\ eigenmaps$ , a method in whose general framework other techniques may be interpreted. It employs the graph Laplacian and does well in preserving locality, yet is less adept at determining local linearity. This shortcoming is mitigated by  $locally\ linear\ embedding\ (LLE)$  and its variants. LLE is based on the idea that the embedded manifold may be approximated by locally linear neighborhoods in  $\mathbb{R}^D$ . Since weights resulting from linear reconstruction are believed to reflect the intrinsic geometry of the manifold, they are topological

properties and as such invariant to rotations, rescalings, and translations. By consequence, these same weights should also reconstruct points in d dimensions. LLE thus maps vicinity structures to the d-dimensional subspace and finds the coordinates that preserve them best. This requires solving the least-squares problem of minimizing reconstruction error and then the sparse eigenvalue problem of minimizing embedding cost. Convexity of both sub-problems ensures globality of local optima. A later proposition,  $Hessian\ LLE\ (H-LLE)$ , may be viewed as an algorithmic variant of LLE and a conceptual variant of Laplacian eigenmaps using the Hessian en lieu of the Laplacian.

These approaches have been shown to successfully retrieve manifold structures in different applications. However, their fully unsupervised functionality offers a draw-back: they may fail to find a low-dimensional embedding that has an actual reflection in the real-life setting. Such situations might require the specification of some pre-labeled instances. Also, it may simply be the case that manual analysis of a subset of the data is available at low cost.

When prior knowledge is at hand it is only natural to use it. Therefore, Yang et al. (2006) proposed SS-LLE as an extension to LLE that is able to harvest prior information. Both exact and inexact knowledge, the latter regularized with an uncertainty coefficient, are applicable. The information is incorporated in the second step of the algorithm by fixing some of the sought-for coordinates in advance. Perhaps unsurprisingly, Yang et al. (2006) find that careful selection of the prior points to be maximally scattered across the manifold surface works better than random sampling. Indeed, the presented results indicate considerable success of their technique.

It is the aim of this report to (1) reproduce these results, thereby creating an opensource implementation of SS-LLE, and (2) to apply SS-LLE to further manifold learning tasks for a more thorough assessment of its performance.

## Contents

1	Intr	oduction	1
2	2.1	Concepts in Manifold Learning	
	2.2	Formal Goal of Manifold Learning	6
3	Local Graph-Based Manifold Learning (LGML)		
	3.1	Overview	
		3.1.1 Taxonomy	
	2.0	3.1.2 Intuition	
	3.2	Conceptual Framework of LGML	
		3.2.2 Graph Approximation	
		3.2.3 Eigenanalysis	
		o.z.o Eigonamarjois I I I I I I I I I I I I I I I I I I I	
4	LGN	ML Techniques	13
	4.1	Unsupervised Techniques	
		4.1.1 Laplacian Eigenmaps (LEM)	
		4.1.2 Locally Linear Embedding (LLE)	
	4.0	4.1.3 Hessian Locally Linear Embedding (HLLE)	
	4.2	Semi-Supervised Locally Linear Embedding (SSLLE)	
		4.2.1 Employment of Prior Information	18 18
		4.2.3 SSLLE Algorithm	
	4.3	Particular Challenges	
5	Exp	eriment Results	20
_	5.1		
		5.1.1 Software Implementation	
		5.1.2 Evaluation Framework	
	5.2	Application to Synthetic Data	20
		5.2.1 Data	20
		5.2.2 Results	20
	5.3	Sensitivity Analysis	20
6	Disc	cussion	20
7	Conclusion		20
A	App	endix	V
R	Elec	tronic Appendix	VI

## List of Symbols

Number of observed dimensions  $d \in \mathbb{N}$ Number of dimensions of embedded manifold Number of dimensions of  $m \in \mathbb{N}$ low-dimensional representation  $N \in \mathbb{N}$ Number of observed data points  $oldsymbol{\mathcal{X}} \subset \mathbb{R}^D \ oldsymbol{X} = (oldsymbol{x}_1, oldsymbol{x}_2, ..., oldsymbol{x}_N) \in (\mathbb{R}^D)^N$ d-manifold embedded in  $\mathbb{R}^D$ Observed coordinates of data sampled from  $\mathcal{M}$  $oldsymbol{Y} = (oldsymbol{y}_1, oldsymbol{y}_2, ..., oldsymbol{y}_N) \in (\mathbb{R}^m)^N$ Learned coordinates of low-dimensional representation of data

# List of Figures

1	S-curve manifold	5
2	Overview on selected manifold learning models	7
3	Schematic idea of kernel PCA	9
4	S-curve neighborhood graph	12
5	Tangent hyperplane for two-dimensional unit sphere	14
6	Linear reconstruction in LLE	16

## List of Tables

## 1 Introduction

Machine learning problems increasingly employ data of high dimensionality. While a large amount of samples is beneficial to learning, high-dimensional feature spaces, such as in speech recognition or gene processing, pose serious obstacles to the performance and convergence of most algorithms (Cayton, 2005).

High dimensionality. Three aspects strike as particularly problematic: computational operations, interpretation of results, and geometric idiosyncrasies. Computational cost must be considered but is becoming less of an issue with technological evolution (Leist et al., 2009). By contrast, the demand for explainable results (for reasons of, say, safety or ethics) is rather intensified by the advance of complex technology. Alas, interpretation in more than a few dimensions is virtually inaccessible to humans (Doshi-Velez and Kim, 2017). The geometric aspect is often addressed as curse of dimensionality, a term subsuming various phenomena of high-dimensional spaces. It is generally not straightforward to infer properties of objects in these spaces as geometric intuition developed in lower dimensions can be misleading. Crucially, the exponential increase of spatial volume induces sparsity. Consequences of this behavior are, among others, a sharp incline in the number of points required to sample the feature space and a loss in meaningfulness of distances. Many learners, however, rely on these concepts<sup>1</sup> and see their functionality deteriorate (Verleysen and Francois, 2005).

**Manifold learning.** These challenges make the case for dimensionality reduction, that is, the endeavor of compressing problem dimensionality to a manageable size. Far from undue simplification, dimensionality reduction is justified by the belief that the latent data-generating process is indeed of much lower dimension than is observed. Consider, for example, image data showing objects in different poses. Such data are typically stored in high-dimensional pixel representations, yet it is reasonable to suppose that variation in these images is in fact caused by a small number of latent features. More formally, the data are assumed to lie on a d-dimensional manifold embedded in the D-dimensional observation space, with  $d \ll D$ . This belief is referred to as manifold assumption (Cayton, 2005). A crucial property of d-manifolds, i.e., the d-dimensional generalization of a curved surface, embedded in  $\mathbb{R}^D$ , is their local topological equivalence to  $\mathbb{R}^d$  (Ma and Fu, 2011). It is precisely this fact that allows manifold coordinates to be mapped to  $\mathbb{R}^d$  in a reduction of dimensionality<sup>2</sup>. The goal is now to learn this mapping in an unsupervised manner (Cayton, 2005). Mapping manifold coordinates to  $\mathbb{R}^d$  is in general not equivalent to simple projection onto the d-dimensional coordinate hyperplanes. Instead, models must learn the intrinsic neighborhood structure of the manifold to establish a notion of "nearness" between points. Standard distance metrics do not apply (globally) as points on general manifolds are connected by curved paths rather than straight lines (Ma and Fu, 2011).

 $<sup>^{1}</sup>$ For instance, consider support vector machines and k-nearest neighbors, both of which rely on distances, or tuning, which often requires extensive sampling of the hyperparameter space.

<sup>&</sup>lt;sup>2</sup>The most intuitive example of this is probably the representation of the Earth, which is a two-dimensional manifold enclosed in three-dimensional space, on two-dimensional maps.

#### Adapt to structure of chapter 3

Local graph-based techniques. Various approaches have been proposed to retrieve points' manifold. A taxonomy may, for example, be found in van der Maaten et al. (2009). Many rely on spectral techniques, trying to find a matrix representation of the data whose principal eigenvectors are used to span a d-dimensional subspace. One group of spectral methods attempts to retain global isometry by mapping pairwise distances to  $\mathbb{R}^d$ . Among them, some are based on Euclidean distances and thus confined to learning linear embeddings (such as principal component analysis (PCA) or multi-dimensional scaling (MDS)). Since linearity is a strong assumption that will not hold for general manifolds, non-linear techniques are more widely applicable (van der Maaten et al., 2009). For example, Isomap achieves non-linearity by applying geodesic distances in the MDS setup (Tenenbaum et al., 2000). Research indicates, however, that for non-convex manifolds it is more effective to preserve local structures only. Otherwise, solutions are prone to shortcuts, i.e., placing points close in  $\mathbb{R}^D$  next to each other when they lie in fact on quite different parts of the manifold (Belkin and Niyogi, 2001). In order to avoid such miscalculations, sparse techniques focus on merely local neighborhood structures, modeled through weighted graph representations. The information from these graphs is then condensed into a sparse matrix. Eventually, the principal eigenvectors of this matrix yield the desired low-dimensional coordinates (van der Maaten et al., 2009).

**Locally linear embedding.** One such local graph-based technique is *locally linear* embedding (LLE), the unsupervised algorithm SSLLE builds upon (Roweis and Saul, 2000). LLE is based on the idea that the embedded manifold may be approximated by locally linear neighborhoods in  $\mathbb{R}^D$ . Weights for the resulting graph are obtained by linear reconstruction of points from their neighbors. As these weights are believed to reflect the intrinsic geometry of the manifold, they are topological properties and should as such also reconstruct points in d dimensions. LLE thus maps vicinity structures to the d-dimensional subspace and finds the Euclidean coordinates that preserve them best by means of spectral decomposition (Roweis and Saul, 2000). Much of the theoretical foundation for LLE has been discussed only in later work. In particular, Belkin and Niyogi (2001) proposed Laplacian eigenmaps (LEM), a method which employs the graph Laplacian, and provided evidence for the fact that, under certain assumptions, LLE may be generalized to the same framework (Belkin and Niyogi, 2003). A later proposition by Donoho and Grimes (2003), Hessian LLE (HLLE), may be viewed as an algorithmic variant of LLE and a conceptual variant of LEM (using the Hessian en lieu of the Laplacian). The theoretical link between LLE and LEM, centered around the Laplace-Beltrami operator, has recently been found to hold less generally than previously assumed (Wu and Wu, 2018). It still appears beneficial to interpret all methods in this common framework also found by Bengio et al. (2003); a more thorough study of convergence guarantees is left to future research.

Semi-supervised extension. The above approaches have been shown to successfully retrieve manifold structures in different applications (Wu and Wu, 2018). However, their fully unsupervised functionality offers a drawback: they may fail to find a low-dimensional embedding that has an actual reflection in the real-life setting. Such situations might require the specification of some pre-labeled instances. Also, it may

simply be the case that some observations already come with labels, or that annotation of a subset of the data is available at low cost (Yang et al., 2006). When prior knowledge is at hand it is only natural to use it. Therefore, Yang et al. (2006) proposed semi-supervised locally linear embedding (SSLLE), an extension to LLE that is able to harvest prior information. Note that this intuition is rather different from general semi-supervised learning where unlabeled data is used to aid an inherently supervised task with few available labels: SSLLE tackles an inherently unsupervised task by providing some labeled data points as reference<sup>3</sup>.

Outline. Indeed, the presented results indicate considerable success of their technique. It is the aim of this report to (1) reproduce these results, thereby creating an open-source implementation of SSLLE, and (2) to apply SSLLE to further manifold learning tasks for a more thorough assessment of its performance. The rest of the report is organized as follows: chapter 2 provides a mathematical framework where fundamental concepts are briefly introduced; chapter 3 explains the idea of local graph-based manifold learning; chapter 4.2 presents SSLLE in detail; chapter 5 discusses the results of the conducted experiments; and chapter 7 draws final conclusions.

Remark. This report addresses a particular sub-field of dimensionality reduction rooted, more or less, in machine learning. A lot of research has been attracted by deep learning in the recent past. The manifold learning task naturally lends itself to neural networks where the derivation of intermediate data representations is an integral component (perhaps most notably so in autoencoder and generative adversarial systems; see for example Khayatkhoei and Elgammal (2018), Huang et al. (2020)). So, while the focus here lies on a different family of methods, it should not go unmentioned that deep dimensionality reduction is certainly a very promising area.

## 2 Basic Manifold Theory

## 2.1 Concepts in Manifold Learning

This chapter introduces the main geometric concepts considered necessary to provide a solid understanding of SSLLE<sup>4</sup>. It must be noted that everything discussed here is presented through the lens of machine learning, deliberately forsaking the generality inherent to topology. Therefore, assuming features can be represented by coordinates in D-dimensional Euclidean space, all concepts are examined with regard to their meaning in  $\mathbb{R}^D$ . Dimensionality reduction techniques take the data observed in  $\mathbb{R}^D$  to actually lie on a d-dimensional topological space that is not necessarily Euclidean but exhibits some specific properties.

**Topological spaces.** A topological space is constituted by a set T equipped with a

<sup>&</sup>lt;sup>3</sup>On this note, alternative proposals for a semi-supervised LLE have been made, for instance by Zhang and Chau (2009), that build upon a fully supervised LLE model such as the one de Ridder and Duin (2002) had designed for classification tasks.

<sup>&</sup>lt;sup>4</sup>Obviously, the list of concepts discussed is by no means extensive. Theory is presented much more in detail (and mathematical rigor) in, for example, good book.

topology  $\mathcal{T}$ . A topology is a general way of describing relations between elements in T. Consider a function  $\mathcal{T}: T \to 2^T, t \mapsto \mathcal{T}(t)$ , which assigns to  $t \in T$  a set of subsets of T called *neighborhoods*. For  $\mathcal{T}$  to be a topology<sup>5</sup> on T, the following properties must hold (Brown, 2006):

- (T1) If  $\mathcal{T}$  is a neighborhood of t, then  $t \in \mathcal{T}$ .
- (T2) If  $\mathcal{T}$  is a subset of T containing a neighborhood of t, then  $\mathcal{T}$  is a neighborhood of t.
- (T3) The intersection of two neighborhoods of t is again a neighborhood of t.
- (T4) Any neighborhood  $\mathcal{T}$  of t contains a neighborhood  $\mathcal{T}'$  of t such that  $\mathcal{T}$  is a neighborhood of each element in  $\mathcal{T}'$ .

Note that, in this general definition, neighborhoods are based on an abstract notion of "nearness". Learning the structure of a topological space effectively boils down to learning neighborhood relations. In Euclidean topological space these are directly based on distance: neighborhoods around t are constructed by  $\epsilon$ -balls containing all elements within a Euclidean distance of  $\epsilon$  from t. The resulting topology is also called the *metric topology* (McCleary, 2006).

Topological spaces in general are not accessible via distances (or angles, for that matter) known from Euclidean spaces. The ultimate goal therefore is the interpretation of the data in a space that is again Euclidean, albeit of lower dimensionality, where such concepts are meaningful. The next step is then to study how (potentially highly non-linear) topological spaces relate to  $\mathbb{R}^d$ .

**Homeomorphisms.** Consider two topological spaces  $(S, \mathcal{T}_S)$ ,  $(T, \mathcal{T}_T)$  (denoted by the respective shorthands S, T from here) and a mapping function  $f: S \to T$ . If f is bijective and continuous and  $f^{-1}: T \to S$  is also continuous, f is called a homeomorphism (Brown, 2006). Topological spaces for which such a mapping exists are homeomorphic to each other. Any properties of S that T shares when it is homeomorphic to S are referred to as topological properties. Two homeomorphic spaces are thus topologically equivalent (McCleary, 2006).

If there exists a non-negative integer d such that for every s in a topological space S a local neighborhood  $U \ni s$ ,  $U \subset S$ , is homeomorphic to an open subset of  $\mathbb{R}^d$  (sometimes called parameter space), S is locally Euclidean<sup>6</sup> (Ma and Fu, 2011). In other words, there is a homeomorphism  $f: U \to \mathbb{R}^d$  for every element in S. The neighborhoods U are also referred to as coordinate patches and the associated maps f are called coordinate charts (Cayton, 2005). In local neighborhoods S then behaves like  $\mathbb{R}^d$  (Ma and Fu, 2011).

**Manifolds.** Manifolds are now precisely such locally Euclidean topological spaces, with some additional properties. For a topological space  $\mathcal{M}$  to be a d-dimensional manifold<sup>7</sup> (also: d-manifold) it must meet the following conditions (Waldmann, 2014):

- (M1)  $\mathcal{M}$  is Hausdorff.
- (M2)  $\mathcal{M}$  is second-countable.

 $<sup>\</sup>overline{{}^{5}\text{Alternative definitions employ open subsets of } T$ , see for example Waldmann (2014).

<sup>&</sup>lt;sup>6</sup>For locally Euclidean topological spaces it is thus meaningful to speak of elements as points.

 $<sup>^{7}\</sup>mathcal{M}$  is again a shorthand, omitting the explicit notation of the corresponding topology.

#### (M3) $\mathcal{M}$ is locally homeomorphic to $\mathbb{R}^d$ .

The Hausdorff condition is a separation property and ensures that for any two distinct points from  $\mathcal{M}$  disjoint neighborhoods can be found (Brown, 2006). Second-countability restricts the manifold's size via the number of open sets it may possess (Waldmann, 2014).

**Embeddings.** Recall that the data are observed in  $\mathbb{R}^D$  but taken to lie on  $\mathcal{M}$ , locally homeomorphic to  $\mathbb{R}^d$ . This implies the assumption  $\mathcal{M} \subset \mathbb{R}^D$  and  $\mathcal{M}$  is said to be embedded in the ambient D-dimensional Euclidean space (Cayton, 2005). The associated *embedding* is but a map  $f: \mathcal{M} \to$  $\mathbb{R}^D$  whose restriction to  $\mathcal{M}$  is a homeomorphism (Brown, 2006), or, more specifically, the canonical inclusion map identifying points on the manifold as particular points of  $\mathbb{R}^D$  (Waldmann, 2014). It can be shown that K = 2d + 1 is sufficient to create an embedding (Ma and Fu, 2011). Figure 1 shows the so-called S-curve manifold embedded in  $\mathbb{R}^3$ . Clearly, the S-curve as a whole is far from linear, but locally homeomorphic to  $\mathbb{R}^2$  and thus intrinsically two-dimensional.



Figure 1: 10,000 points sampled from the S-curve manifold. *Source:* own representation, inspired by implementation in Python's scikit-learn library (Pedregosa et al., 2011).

Geodesics. One last aspect remains open and shall be briefly touched here, namely how to handle distances on manifolds where Euclidean metrics are not meaningful. Rather than measuring "shortcuts" between points across  $\mathbb{R}^D$  (where, for instance, points in the red upper area of figure 1 would be considered deceptively close to points in the cyan mid area), it makes intuitive sense to constrain distances to the manifold surface. In order to enable the construction of such a metric, manifolds must fulfill two additional properties:  $smoothness^8$  and  $connectedness^9$  (Ma and Fu, 2011). For smooth, connected manifolds,  $geodesic\ distance$  is the length of the shortest curve (geodesic) on  $\mathcal{M}$  between two points on  $\mathcal{M}$ . A curve c in  $\mathcal{M}$  is a smooth mapping from an open interval  $\Lambda \subset \mathbb{R}$  into  $\mathcal{M}$ . c is parametrized by a point  $\lambda \in \Lambda$ , such that

$$c(\lambda) = (c_1(\lambda), ..., c_d(\lambda))^T$$

is a curve in  $\mathbb{R}^d$  (all  $c_j, j = 1, ..., d$  having a sufficient number of continuous derivatives). Component-wise differentiation with respect to  $\lambda$  yields *velocity* in  $\lambda$ :

<sup>&</sup>lt;sup>8</sup>The smoothness property is based on differentiability of coordinate charts and ensures that concepts of curvature, length and angle remain meaningful (Ma and Fu, 2011). A detailed derivation may be found, for example, in Mukherjee (2015).

<sup>&</sup>lt;sup>9</sup>Connectedness means that no separation  $\{U, V\}$  of a manifold  $\mathcal{M}$  exists with open, non-empty and disjoint  $U, V \subset \mathcal{M}$ ,  $\mathcal{M} = U \cup V$ . This may be loosely put as paths linking arbitrary pairs of manifold points (McCleary, 2006).

$$c'(\lambda) = (c'_1(\lambda), ..., c'_d(\lambda))^T.$$

The *speed* of c is given by  $\|c'(\lambda)\|_2^2$ , where  $\|\cdot\|^2$  denotes the square norm. Distance along this curve is measured by the arc-length

$$L(c) = \int_{\mathbf{p}}^{\mathbf{q}} \|c'(\lambda)\|^2 d\lambda.$$

Eventually, geodesic distance can be derived as the length of the shortest such curve, out of the set C(p, q) of differentiable curves in  $\mathcal{M}$  that connect p and q:

$$d^{\mathcal{M}}(\boldsymbol{p}, \boldsymbol{q}) = \inf_{c \in \mathcal{C}(\boldsymbol{p}, \boldsymbol{q})} L(c). \tag{1}$$

Intuitively, geodesic distance can be identified with Euclidean distance in Euclidean spaces where shortest curves are just straight lines (Ma and Fu, 2011).

### 2.2 Formal Goal of Manifold Learning

Building on the above concepts, the data situation in manifold learning might be summarized as follows: data are observed in  $\mathbb{R}^D$  but assumed to be really samples from a d-manifold  $\mathcal{M}$  embedded in  $\mathbb{R}^D$ , meaning they can be analyzed in  $\mathbb{R}^d$  if a faithful translation between respective coordinates in  $\mathbb{R}^D$  and  $\mathbb{R}^d$  is found<sup>10</sup>. The challenge is thus to unravel the manifold in a way that preserves its intrinsic structure to maximum extent (Saul et al., 2006). This goal shall be formalized in a way that will be referenced throughout the remainder of this report and that is inspired by the works of Cayton (2005) and Saul et al. (2006).

**Given.** Data  $\mathcal{X} = (\boldsymbol{x}_1, \boldsymbol{x}_2, ..., \boldsymbol{x}_N)$ , with  $\boldsymbol{x}_i \in \mathbb{R}^D \ \forall i \in \{1, 2, ..., N\}$  and  $N, D \in \mathbb{N}$ .  $\mathcal{X}$  thus consists of N real-valued data vectors observed in D dimensions. The true data-generating process is taken to have dimensionality  $d \in \mathbb{N}$ , such that  $\mathcal{X}$  is in fact a sample from a smooth, connected d-manifold with  $\mathcal{X} \sim \mathcal{M} \subset \mathbb{R}^D$ .  $\mathcal{M}$  may be described by a single coordinate chart<sup>11</sup>  $\psi : \mathcal{M} \to \mathbb{R}^d$ . For manifold learning methods to yield satisfying results,  $\mathcal{M}$  is always assumed to be sampled well by  $\mathcal{X}$ .

**Goal.** Find the *d*-dimensional representation of the data, i.e., compute  $\mathcal{Y} = (\boldsymbol{y}_1, \boldsymbol{y}_2, ..., \boldsymbol{y}_N)$ , where  $\boldsymbol{y}_i = \psi(\boldsymbol{x}_i) \in \mathbb{R}^d \ \forall i \in \{1, 2, ..., N\}$ . The map  $\psi$  itself is not always explicitly retrieved.

Note that, while D is given a priori, the intrinsic dimensionality d is often unknown in real-life applications. Due to the lack of (finite-sample) convergence guarantees for many methods dimensionality estimations may not always be equal to d.  $\mathcal{Y}$  as found by manifold learning techniques must therefore be expected to differ from the true coordinates and, in particular, to even have incorrect dimension (Saul et al., 2006). Notwithstanding this potential gap, solutions of the subsequently presented methods will be denoted by  $\mathcal{Y} \in (\mathbb{R}^d)^N$  to avoid overly complicated notation.

 $<sup>\</sup>overline{^{10}}$ It is actually a simplification to assume all data to lie *exactly* on  $\mathcal{M}$ , but the more general case of data lying *near*  $\mathcal{M}$  is rarely considered explicitly.

<sup>&</sup>lt;sup>11</sup>This is possible for any smooth, compact manifold (Cayton, 2005).

## 3 Local Graph-Based Manifold Learning (LGML)

#### 3.1 Overview

#### 3.1.1 Taxonomy

After the goal of manifold learning has been formalized, it shall now be laid out how the problem is approached by LLE as the conceptual parent of SSLLE (the incorporation of prior information is a rather different matter that will be addressed in chapter 4.2; apart from this, the basic functionalities of SSLLE and LLE are identical and will therefore be presented as one). Much of the theoretical foundation for LLE has been discussed only in later work. In order to provide a more integrated background, explanations will therefore be given in a broader context of local graph-based manifold learning (LGML), which also comprises LEM and HLLE. The particular relationship of the three methods shall be made clear along the way. LGML arises from a variety of geometric intuitions and computational implementations. Nonetheless, methods share common structures that allow for interpretation in a more abstract framework (Bengio et al. (2003), Bengio et al. (2004)). It should be noted that such a framework might be established from several angles; after all, the different approaches attempt to solve the same problem and can thus be translated into one another in various ways.

Figure 2 depicts a schematic overview on the models studied here, representing the specific perspective taken within this report. All of these belong to the realm of *spectral* models. The non-spectral group includes, among others, techniques based on neural networks and is not discussed here (van der Maaten et al., 2009).

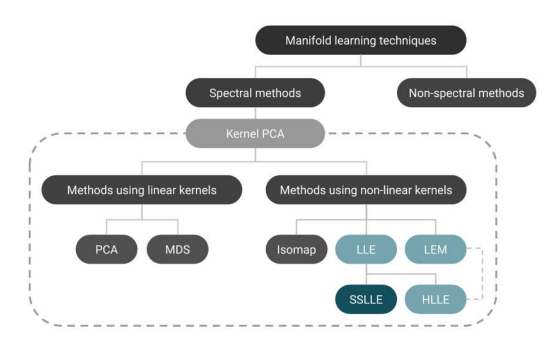


Figure 2: A schematic overview on selected methods of manifold learning (the list is by no means extensive and could arguably be ordered in several alternative ways). *Source:* own representation, inspired by a similar example given in van der Maaten et al. (2009) and re-interpreted with the findings in Bengio et al. (2004).

#### 3.1.2 Intuition

As indicated in figure 2, this report will sketch the idea behind LGML in the light of kernel principal component analysis (kernel PCA). Kernel PCA was actually proposed first and later shown to link the other concepts by a unified idea (Ham et al. (2003). It makes for an appealing framework that provides a useful general intuition to manifold learning and subsumes the other methods in a way beneficial to the important task of out-of-sample extension (Bengio et al., 2004).

**Kernel PCA.** Kernel PCA builds upon two fundamental concepts in machine learning: it performs *principal component analysis (PCA)* on data transformed by the *kernel trick*. It undertakes two subsequent steps. First, features of interest are extracted from the data by kernelization. These are taken to capture the intrinsic data structure and may therefore be understood as an approximation to the latent manifold properties. In the end, they constitute a matrix representation. Second, PCA finds the principal axes along which these intrinsic properties vary, yielding the desired reduction in dimensionality by preserving the most relevant latent dimensions (Schölkopf et al., 1998).

**Kernelization.** By kernelization, mapping the data to a space  $\mathcal{F}$  of arbitrarily high dimension, features may be obtained that relate to the input in a non-trivial way<sup>12</sup>. Crucially, the feature map  $\phi: \mathbb{R}^D \to \mathcal{F}$  need not be computed explicitly, which might prove prohibitively expensive. Kernelization instead solely relies on inner products  $\langle \phi(\boldsymbol{x}_i), \phi(\boldsymbol{x}_j) \rangle$  of the transformed inputs. Employing Mercer's theorem of functional analysis, these inner products may be interpreted as performed by a continuous kernel  $\kappa(\boldsymbol{x}_i, \boldsymbol{x}_j)$  in some space with Hilbert property. Appropriate choice of  $\kappa$  then allows for the data to be represented by a matrix  $\boldsymbol{K} \in \mathbb{R}^{N \times N}, \boldsymbol{K}_{ij} = \kappa(\boldsymbol{x}_i, \boldsymbol{x}_j)$ . This matrix is the numerical data representation derived with respect to their latent properties. (Schölkopf et al., 1998). Precisely how it is computed depends on the choice of the kernel function  $\kappa$  and gives rise to different techniques (Ham et al., 2003).

**PCA.** PCA is a quite powerful technique by itself. It finds the directions of maximum variance through eigenanalysis of the empirical covariance matrix, yielding the most important axes of inter-feature relations that coincide with the principal eigenvectors of the covariance matrix. The data are projected into the linear subspace spanned by these d eigenvectors, thereby mapping the observations to a coordinate system given by the linear feature combinations that represent the strongest (co)variability. Note that for this transformation to actually rotate the coordinate system about the origin, the data must be mean-centered beforehand. PCA thus performs an orthogonal input transformation that allows for dimensionality reduction at minimal information loss (Cayton, 2005). In kernel PCA, this eigenanalysis is implicitly performed in the feature space  $\mathcal{F}$ . Algorithmically, it boils down to diagonalizing the kernel matrix  $\mathbf{K}$  (Schölkopf et al., 1998).

<sup>&</sup>lt;sup>12</sup>Support vector machines use the kernel trick to achieve linear separability. An intuitive example may be given by data observed in two classes that form concentric circles in  $\mathbb{R}^2$ . While such data are not linearly separable in two dimensions, they are in three: mapping the classes to different heights enables separation by a horizontal hyperplane. This example also hints at the idea of (spectral) clustering to which kernel PCA is indeed intimately related (Bengio et al., 2004).

Figure 3 visualizes the idea of kernel PCA. The original data (left) are observed in two dimensions but clearly intrinsically one-dimensional, where the non-linear manifold feature is expressed by coloring. The kernel trick creates a feature map, visualized here as a projection of the intrinsic feature to a third coordinate axis (middle). Coercing the data to this dimension as the sole axis of variation yields the desired one-dimensional representation (right).



Figure 3: Schematic idea of kernel PCA: from data observed in two dimensions, but clearly of intrinsic dimensionality one (*left*), create a mapping to a higher-dimensional feature space (*middle*), reduction of which to its principal axes yields the desired one-dimensional representation (*right*). Source: own representation, using a subset of mlbench's noise-free spirals data. Note that this is but a schematic depiction where the mid and right representation have not been created by an actual implementation of kernel PCA.

## 3.2 Conceptual Framework of LGML

#### 3.2.1 Motivation

If kernel PCA sounds like a powerful concept, the crux of course lies in finding an appropriate kernel function. The nature of the feature map applied to the input data determines the kind of mapping that may be learned and serves to distinguish the various techniques. As foreshadowed in figure 2, spectral methods decompose into groups using linear and non-linear kernels, respectively. This distinction directly translates to the feature map  $\phi$ . Linear methods suffer from the confinement to finding linear subspaces (van der Maaten et al., 2009). PCA in its standard form can be interpreted as kernel PCA by identifying the kernel function with the covariance function. It thus returns the subspace of greatest variability in the original input features (Ham et al., 2003). The closely related multi-dimensional scaling (MDS) yields the same result<sup>13</sup>, albeit from a different intuition (Saul et al., 2006).

As extensively discussed above,  $\mathcal{X}$  must often be assumed to lie on a non-linear manifold  $\mathcal{M} \subset \mathbb{R}^D$ , which is precisely why kernelization is usually performed such that the resulting feature space is related to the input space in a non-linear way (Schölkopf et al., 1998). Conceivably, there is no obvious way to arrive at such a mapping. Graph-based models therefore approach the problem from an alternative angle. In fact, they do not even perform kernelization explicitly: they transform the data in a way that can be shown to correspond to applying a (data-dependent)

<sup>&</sup>lt;sup>13</sup>At least, in its metric form; there are alternative formulations of MDS that are not equivalent to PCA (see, for example, Williams (2002)).

kernel function<sup>14</sup>, but the fundamental intuition is a different one.

Non-linearity. The key idea in graph-based learning is to approximate the manifold by a discretized graph representation. Such a graph may be intuitively imagined as a skeletal model of  $\mathcal{M}$  (an example is given in 4). This way, distances may be measured along the approximated manifold surface rather than in the ambient Euclidean space, effectively enabling non-linearity. Functionals that vary across methods are used to describe properties of the graph – essentially a proxy of the latent manifold properties (Saul et al., 2006).

**Locality.** A second desideratum in general manifold learning is the ability to treat highly non-linear manifolds with sufficiently local focus. Non-convexity means  $\mathcal{M}$  is isometric to a non-convex subset of Euclidean space (Donoho and Grimes, 2003). Intuitively, such behavior requires careful tracing of the manifold surface. LGML methods therefore focus on solely local manifold properties (Cayton, 2005). They are frequently contrasted to *Isomap*, one of the earliest and most prominent examples of global manifold learning. Isomap retains pairwise distances between points on the manifold surface as measured along graph edges via geodesic curves<sup>15</sup> (Tenenbaum et al., 2000). Its central assumptions are global isometry and convexity of the parameter space (Tenenbaum et al., 2000). While it yields good results in many applications, Isomap does not sufficiently account for the curvature of strongly nonconvex manifolds. In order to avoid this drawback, local methods limit isometry to only hold between neighboring points and relax the parameter space condition to open, connected subspaces (Donoho and Grimes, 2003).

**Algorithmic procedure.** Summing up the above, LGML methods use functionals defined on graph approximations of the manifold to capture the intrinsic data structure. This information is stored in a matrix representation M that is directly linked to the kernel matrix K. Eigenanalysis of M then leads to the sought-for low-dimensional subspace coordinates.

The resulting algorithmic pattern may be stated as follows (Bengio et al., 2003):

- 1. Construct a neighborhood graph  $\mathcal{G}$  from the observed data.
- 2. Analyze the graph properties with an appropriate functional and derive a matrix representation M thereof.
- 3. Find the eigenvalues and associated eigenvectors of M.
- 4. From the principal (top or bottom) eigenvectors, as determined by the ordered eigenvalues, retrieve the low-dimensional coordinates.

#### 3.2.2 Graph Approximation

All LGML methods fundamentally build on graph approximations of the manifold surface. Note that these graphs are constructed from neighborhood relations in the high-dimensional observation space and do not require any prior information about

<sup>&</sup>lt;sup>14</sup>The report does not discuss the actual kernel function as their illustrative ability is considered rather limited. For an explicit formulation of kernels in LLE, LEM and HLLE see for example Bengio et al. (2004) and Weinberger et al. (2004).

<sup>&</sup>lt;sup>15</sup>It is thus a non-linear variant of MDS, which uses standard Euclidean distances (Tenenbaum et al., 2000).

the latent manifold coordinates.

**Neighborhoods.** A neighborhood of  $x \in \mathcal{X}$  is a subset of  $\mathcal{X}$  containing another, open subset of  $\mathcal{X}$  of which x is an element. Members of the neighborhood are called neighbors of x. In metric spaces neighborhoods are defined via distances and therefore translate to open balls around each point (Waldmann, 2014). This distance-based construction locally applies to manifolds as a direct consequence of their local isometry to the Euclidean observation space (Ma and Fu, 2011). There are two principal ways to build a neighborhood around  $x \in \mathcal{X}$ , both of which usually employ the squared Euclidean norm<sup>16</sup>  $\|\cdot\|^2$ . Let  $\mathcal{N}: \mathcal{X} \to \mathcal{X}^{\ell}, x \mapsto \mathcal{N}(x)$  be a constructor that assigns a set of neighbors to x. The first possibility is to restrict the size of the neighborhood to the k points<sup>17</sup> with the smallest distance to x, such that  $\ell = k$  and

$$\mathcal{N}_k(\boldsymbol{x}) = \{ \boldsymbol{x}_i \in \mathcal{X} : \|\boldsymbol{x} - \boldsymbol{x}_i\|^2 \le d_{(k)} \}, \tag{2}$$

with  $d_{(k)} \in \mathbb{R}$  being the k-th instance of ordered pairwise distances between  $\boldsymbol{x}$  and all other points. Alternatively, the neighborhood may be constructed by collecting all points that have a maximum distance of  $\epsilon \in \mathbb{R}$  to  $\boldsymbol{x}$ , yielding

$$\mathcal{N}_{\epsilon}(\boldsymbol{x}) = \{\boldsymbol{x}_j \in \mathcal{X} : \|\boldsymbol{x} - \boldsymbol{x}_j\|^2 \le \epsilon\}$$
(3)

and  $\ell = |\mathcal{N}_{\epsilon}(\boldsymbol{x})|$  (He et al., 2005).

Both k and  $\epsilon$  are hyperparameters that must be specified up-front. Their choice reflects beliefs about the topological structure of  $\mathcal{M}$  – smaller neighborhoods corresponding to a higher degree of non-linearity – and may affect performance rather strongly (Sudderth, 2002). Chapter 4.3 will discuss this trade-off, which is also addressed in the practical implementation (section 5), in more detail. In this, the remainder of the report will focus on the k-neighborhood notion as it is typically more easily specified due to its inherent scale invariance and has attracted rather more attention in general research<sup>18</sup>.

**Neighborhood graphs.**  $\mathcal{M}$  can now be characterized by a *neighborhood graph*  $\mathcal{G} = (\mathcal{V}, \mathcal{E})$ , still assuming it is sampled well by  $\mathcal{X}$ . Inputs  $\mathbf{x} \in \mathcal{X}$  form vertices  $\mathcal{V}$  and edges  $\mathcal{E}$  indicate neighborhood relations (Belkin and Niyogi, 2001). Each vertex is connected to its k nearest neighbors or all points within  $\epsilon$ -radius, depending on the neighborhood definition.

<sup>&</sup>lt;sup>16</sup>In principle, alternative metrics are applicable, for instance such that measure angles (Belkin and Niyogi, 2004).

<sup>&</sup>lt;sup>17</sup>In presence of ties in pairwise distances k may vary across the data, but with zero probability in continuous feature spaces.

<sup>&</sup>lt;sup>18</sup>Yet, Tenenbaum et al. (2000) note that, when local dimensionality is not constant across the observed data,  $\epsilon$ -neighborhoods might provide more reliable results.

It is easy to see that k-neighborhoods are an asymmetric notion; for one point to be among another's k nearest neighbors the reverse need not be true. k-neighborhoods therefore lead to directed graphs. Conversely, the  $\epsilon$ -distance boundary holds in both directions and produces undirected graphs (He et al., 2005). Figure 4 shows how a neighborhood graph may be used as an approximation for the example of the S-curve manifold. It was built using k-neighborhoods with k=3. For a densely sampled set of points, the graph representation should yield a fairly good approximation of the manifold surface.



Figure 4: k-neighborhood graph for 300 points sampled from the S-curve with k = 3. Source: own representation.

#### 3.2.3 Eigenanalysis

Eventually, spectral manifold learning boils down to an eigenanalysis of the matrix M believed to hold information about the intrinsic manifold structure. As explained in chapter 3.1, PCA finds the principal eigenvectors of empirical covariance, thereby defining a low-dimensional subspace containing most of the data-inherent variability. The very same idea applies when diagonalizing the more general matrix corresponding to the non-linear feature map: the top (or bottom<sup>19</sup>) d eigenvectors of M span a subspace into which the data may be projected under minimal loss of information. More precisely, the representation of  $\mathcal{X}$  by the d selected eigenvectors of M is loss-optimal with respect to the least-squares error (Schölkopf et al., 1998).

**Eigenvectors and eigenvalues.** Formally, eigenanalysis is the decomposition of a square matrix into pairs of eigenvectors and eigenvalues. Let  $\mathbf{A} \in \mathbb{R}^{N \times N}$  be a square matrix and  $\lambda \in \mathbb{R}$  a scalar value.  $\lambda$  is said to be an eigenvalue to  $\mathbf{A}$  if there exists  $\mathbf{v} \in \mathbb{R}^N \setminus \{0\}$  such that  $\mathbf{A}\mathbf{v} = \lambda \mathbf{v}$ . Then,  $\mathbf{v}$  is the eigenvector corresponding to the eigenvalue  $\lambda$ , and their tuple is also called an eigenpair.

**Null spaces.** A closely related notion is that of the *null space*, consisting of the vectors that map  $\boldsymbol{A}$  to 0 upon multiplication from the right:  $\{\boldsymbol{v} \in \mathbb{R}^N : \boldsymbol{A}\boldsymbol{v} = 0\}$ . It can be easily seen that the null space consists of those eigenvectors of  $\boldsymbol{A}$  that are associated with an eigenvalue of zero, and the zero vector itself. For a specific eigenvalue  $\lambda$  of  $\boldsymbol{A}$ , the null space of  $\lambda \boldsymbol{I} - \boldsymbol{A}$  (with  $\boldsymbol{I}$  the N-dimensional identity matrix) constitutes the *eigenspace* of  $\boldsymbol{A}$  (Börm and Mehl, 2012).

Generalized eigenvalue problems. Eigendecomposition of a matrix A can be framed as the solution of a generalized eigenvalue problem. Generalized eigenvalue problems are posed subject to a constraint on a second, also symmetric matrix  $B \in \mathbb{R}^{N \times N}$ . As

<sup>&</sup>lt;sup>19</sup>This differs across methods and shall be made clear later.

the standard eigenvalue problem results immediately from B = I, the generalized form subsumes both cases. It is given by

$$AV = BV\Lambda, \tag{4}$$

where  $V = [v_1, v_2, ..., v_N] \in \mathbb{R}^{N \times N}$  is the matrix of eigenvectors of A and  $\Lambda = diag[\lambda_1, \lambda_2, ..., \lambda_N]^T \in \mathbb{R}^{N \times N}$  is the diagonal matrix of the associated eigenvalues (ordered from smallest to largest). The generalized eigenvalue problem may be stated equivalently as

$$\min_{\mathbf{V}} trace(\mathbf{V}^T \mathbf{A} \mathbf{V}), \quad \text{s.t.} \quad \mathbf{V}^T \mathbf{B} \mathbf{V} = \mathbf{I},$$
 (5)

and translated to the first form by means of a Lagrangian multiplier (Ghojogh et al., 2019). It must be noted that solving eigenvalue problems becomes computationally challenging rather quickly, an issue on which chapter 4.3 also briefly comments with regard to the methods discussed here (Börm and Mehl, 2012).

Building upon the concepts of neighborhood graph approximation and subsequent eigenanalysis, the following chapter will now present in detail how LEM, LLE and HLLE approach the manifold learning task, and, eventually, how SSLLE seeks to improve the low-dimensional embedding through anchoring with prior points.

## 4 LGML Techniques

### 4.1 Unsupervised Techniques

#### 4.1.1 Laplacian Eigenmaps (LEM)

The reason for LEM to appear in this report alongside the LLE family is its underlying theory both providing a foundation for LLE (Belkin and Niyogi, 2003), which was originally proposed lacking such, and closely relating to the theoretical concepts in HLLE (Donoho and Grimes, 2003).

LEM are centered around the preservation of locality, i.e., mapping nearby inputs to nearby outputs. Locality is enforced via the *Laplace-Beltrami operator* defined on smooth, compact manifolds, and operationalized by means of the *graph Laplacian* acting as a discrete approximator (Belkin and Niyogi, 2003). This idea is best understood recalling that the similarity of outputs for similar inputs is essentially a notion of smoothness and can thus be controlled by a size constraint on the gradient of the mapping function.

**Laplace-Beltrami operator.** Consider the twice differentiable function  $f: \mathcal{M} \to \mathbb{R}$  mapping two points  $\mathbf{p}, \mathbf{q} \in \mathcal{M}$  to  $f(\mathbf{p})$  and  $f(\mathbf{q})$ , respectively. On  $\mathcal{M}$  these points are connected by a length-parametrized curve c(t). Denote the geodesic distance between  $\mathbf{p}$  and  $\mathbf{q}$  by  $\ell$ , such that  $\mathbf{p} = c(0)$  and  $\mathbf{q} = c(\ell)$ .

Gradients of f with respect to p are defined in the local tangent space  $T_p(\mathcal{M})$  spanned by vectors tangent to  $\mathcal{M}$  at p. As  $\mathcal{M}$  is embedded in  $\mathbb{R}^D$ , its tangent spaces are Euclidean and of dimension d, i.e., d-dimensional hyperplanes (Sudderth, 2002) as depicted in figure 5. If p is identified with the origin of  $T_p(\mathcal{M})$ , the tangent space inherits an orthonormal coordinate system obtained from endowing  $T_p(\mathcal{M})$  with the inner product from  $\mathbb{R}^d$  (Donoho and Grimes, 2003). With this, the distance |f(p) - f(q)| of mappings can be expressed as the length of the integral  $\int_0^\ell \langle \nabla f(c(t)), c'(t) \rangle dt$ . In other words, the geodesic curve connecting p and q is projected onto  $T_p(\mathcal{M})$ , and the length of this projection depends on the gradient of f and the curve velocity.

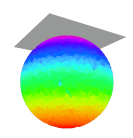


Figure 5: Tangent hyperplane for an exemplary point on the two-dimensional unit sphere manifold, embedded in  $\mathbb{R}^3$ . Source: own representation.

Exploiting the Schwartz equality and relations formally proved by Belkin and Niyogi (2008), it can be shown that  $|f(\mathbf{p}) - f(\mathbf{q})| \leq \|\nabla f(\mathbf{p})\| \cdot \|\mathbf{p} - \mathbf{q}\| + o$ , where o marks a term of vanishing size. As the distance between  $\mathbf{p}$  and  $\mathbf{q}$  is a datum,  $\|\nabla f\|$  controls how far apart points are mapped on the real line. Consequently, the goal is to find a mapping that, on average, preserves locality by posing a second-order penalty on  $\|\nabla f\|$  and minimizing  $\int_{\mathcal{M}} \|\nabla f\|^2$ . This is just equal to minimizing  $\int_{\mathcal{M}} \mathcal{L}(f)f$  with the Laplace-Beltrami operator  $\mathcal{L}$  (Belkin and Niyogi, 2003). Applying the operator  $\mathcal{L}$  to f results in a function from the same function space as f, and for  $\mathcal{L}f = \lambda f$ , f is an eigenfunction of  $\mathcal{L}$  with  $\lambda \in \mathbb{R}$  as its associated eigenvalue. Crucially, these eigenfunctions are orthogonal for  $\mathcal{L}$  and their eigenvalues are real, meaning they are natural candidates for forming a functional basis (Levy, 2006). The optimal embedding map is then given by the d principal eigenfunctions of  $\mathcal{L}$  after removing the bottom one which would map  $\mathcal{M}$  to a single point (Belkin and Niyogi, 2003).

**Graph Laplacian.** Now the same reasoning can be applied to the neighborhood graph approximation of  $\mathcal{M}$ . Recall the desideratum of mapping nearby inputs to nearby outputs. LEM achieves this by assigning edge weights<sup>20</sup>

$$w_{ij} = \begin{cases} \exp(\frac{\|x_i - x_j\|^2}{t}), t \in \mathbb{R}, & \text{if } x_i, x_j \text{ are connected,} \\ 0 & \text{otherwise,} \end{cases}$$

forming a weight matrix  $\mathbf{W} = (w)_{ij} \in \mathbb{R}^{N \times N}$ . Clearly, edges between closer points receive larger weights. A second matrix  $\mathbf{D} = (d)_{ij} \in \mathbb{R}^{N \times N}$  takes the row sums of  $\mathbf{W}$  on its diagonals. Penalizing output disparities more severely for pairs of nearby points, i.e., pairs with a large weight coefficient, the smoothness requirement may be stated as follows:

<sup>&</sup>lt;sup>20</sup>These weights stem from the heat kernel intimately related to the Laplace-Beltrami operator and ensure positive semi-definiteness of the resulting graph Laplacian. As an alternative, Belkin and Niyogi (2003) propose a simpler kernel that is equal to 1 for connected nodes and 0 otherwise.

$$\min_{\mathcal{Y}} \sum_{i,j} \|\boldsymbol{y}_i - \boldsymbol{y}_j\|^2 w_{ij} = \min_{\mathcal{Y}} \sum_{i,j} \boldsymbol{y}_i^T \boldsymbol{y}_i w_{ij} + \boldsymbol{y}_j^T \boldsymbol{y}_j w_{ij} - 2 \boldsymbol{y}_i^T \boldsymbol{y}_j w_{ij} 
= \min_{\mathcal{Y}} \sum_{i} \boldsymbol{y}_i^T \boldsymbol{y}_i d_{ii} + \sum_{j} \boldsymbol{y}_j^T \boldsymbol{y}_j d_{jj} - 2 \sum_{i,j} \boldsymbol{y}_i^T \boldsymbol{y}_j w_{ij}.$$

Now, define the graph Laplacian as  $\boldsymbol{L} = \boldsymbol{D} - \boldsymbol{W} \in \mathbb{R}^{N \times N}$ , thereby coercing all information about the graph structure into a single matrix representation<sup>21</sup>. With  $\boldsymbol{L}$  the above can be rewritten as

$$\min_{\mathcal{V}} trace(\mathcal{Y}^T L \mathcal{Y}), \quad \text{s.t. } \mathcal{Y}^T D \mathcal{Y} = I,$$
(6)

which has precisely the form of the generalized eigenvalue problem in equation 4 and is therefore solved by eigendecomposition of  $\boldsymbol{L}$  (Belkin and Niyogi, 2003). As in the continuous case, the bottom eigenvector with zero eigenvalue is constant and must be discarded. The subsequent d eigenvectors yield the desired low-dimensional embedding (Levy, 2006).

#### 4.1.2 Locally Linear Embedding (LLE)

In proposing LEM, Belkin and Niyogi (2003) also demonstrated how the somewhat earlier LLE algorithm may be reinterpreted within the LEM framework: it can be shown to approximate the graph Laplacian under certain conditions and thus asymptotically approach the Laplace-Beltrami operator. More recent research, however, suggests that these conditions might be more restrictive than previously assumed. In particular, convergence appears to depend on the choice of a regularization parameter required in the case of D < k (Wu and Wu, 2018).

Idea. The initial proposal by Roweis and Saul (2000), ignorant to these findings, was made with a different, and rather heuristically motivated, intuition. LLE relies on a simple yet powerful idea. Each point  $x_i$  in the D-dimensional input space is expressed as a convex combination of its neighbors, such that the weighting coefficients of this reconstruction essentially represent the edge weights of the neighborhood graph around  $x_i$ . These (generalized) barycentric coordinates now bear a crucial property: they are invariant to rotation, rescaling and translation of the neighborhood, and thus topological properties that equally hold in the low-dimensional embedding space. In other words, the same weights that serve to reconstruct  $x_i$  in  $\mathbb{R}^D$  should do so in  $\mathbb{R}^d$  (Roweis and Saul, 2000). Obviously, this belief is only justified if  $\mathcal{M}$  is indeed locally linear and the graph edges run along the manifold surface rather than short-circuiting it, hinting at the important role of neighborhood size which will be discussed in chapter 4.3.

<sup>&</sup>lt;sup>21</sup>This corresponds to the general matrix M introduced in chapter 3.2.1; the deviating notation shall merely emphasize the special role of M as the Laplacian.

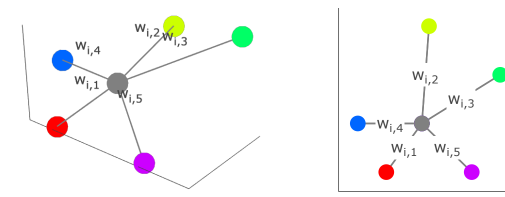


Figure 6: Reconstruction in three (left) and two (right) dimensions, employing the same reconstruction weights. *Source:* own representation.

Algorithmically, LLE performs two subsequent steps (Roweis and Saul, 2000):

- 1. Compute the reconstruction weights in  $\mathbb{R}^D$  minimizing reconstruction loss.
- 2. Compute the embedding coordinates in  $\mathbb{R}^d$  minimizing embedding loss.

**Reconstruction loss minimization.** Reconstruction errors are measured by a quadratic loss function. Optimization of the objective is subject to a sum-one constraint for the weights of each point. A second constraint, zero weights for non-neighboring points, is implicitly enforced during construction of the neighborhood graph, where edges are only drawn to vertices belonging to  $x_i$ 's neighborhood (Ghojogh et al., 2020). The resulting optimization problem is convex and has a unique closed-form solution<sup>22</sup> (Roweis and Saul, 2000):

$$\min_{\boldsymbol{W}} \varepsilon(\boldsymbol{W}) = \min_{\boldsymbol{W}} \sum_{i} \left\| \boldsymbol{x}_{i} - \sum_{j} w_{ij} \boldsymbol{x}_{j} \right\|^{2}$$

$$= \min_{\boldsymbol{W}} \sum_{i} \left\| \boldsymbol{x}_{i} - \boldsymbol{N}_{i} \boldsymbol{w}_{i} \right\|^{2},$$
s.t. 
$$\mathbf{1}^{T} \boldsymbol{w}_{i} = 1 \quad \forall i \in \{1, 2, ..., N\}.$$
(7)

Here,  $N_i \in \mathbb{R}^{D \times k}$  denotes the matrix of feature vectors of  $\boldsymbol{x}_i$ 's neighbors and  $\boldsymbol{w}_i = \sum_j w_{ij} \in \mathbb{R}^k$ .

Equation 7 can be re-arranged by use of the sum-one constraint and simplified by introduction of the gram, or local covariance, matrix  $G_i$  (Saul and Roweis, 2001):

<sup>&</sup>lt;sup>22</sup>Note that the weight matrix W is different from the one computed in LEM, but the same symbol is used as not to overload notation.

$$\min_{\boldsymbol{W}} \varepsilon(\boldsymbol{W}) = \min_{\boldsymbol{W}} \sum_{i} \|\boldsymbol{x}_{i} \mathbf{1}^{T} \boldsymbol{w}_{i} - \boldsymbol{N}_{i} \boldsymbol{w}_{i}\|^{2}$$

$$= \min_{\boldsymbol{W}} \sum_{i} \boldsymbol{w}_{i}^{T} (\boldsymbol{x}_{i} \mathbf{1}^{T} - \boldsymbol{N}_{i})^{T} (\boldsymbol{x}_{i} \mathbf{1}^{T} - \boldsymbol{N}_{i}) \boldsymbol{w}_{i}$$

$$= \min_{\boldsymbol{W}} \sum_{i} \boldsymbol{w}_{i}^{T} \boldsymbol{G}_{i} \boldsymbol{w}_{i},$$
s.t. 
$$\mathbf{1}^{T} \boldsymbol{w}_{i} = 1 \quad \forall i \in \{1, 2, ..., N\}.$$
(8)

By standard use of a Lagrange multiplier, the solution for the above constrained optimization problem collapses to

$$w_i = \frac{G_i^{-1} \mathbf{1}}{\mathbf{1}^T G_i^{-1} \mathbf{1}}.$$
 (9)

Solving the reconstruction problem therefore requires N matrix inversions, which may prove problematic if the gram matrices do not achieve full rank. In the case of D < k,  $G_i$  is indeed singular and must be robustified by adding a small numerical constant to its diagonal (Ghojogh et al., 2020). Chapter 4.3 will discuss how this regularization is applied.

**Embedding loss minimization.** The second optimization problem minimizes the embedding cost arising from mapping neighborhood geometries, derived in the form of reconstruction weights, into the d-dimensional subspace. Keeping the weight coefficients fixed, the aim is to find such embedding coordinates that best preserve the vicinity structures and adhere to the constraints of summing to zero (i.e., being centered around the origin) and having unit covariance (Roweis and Saul, 2000):

$$\min_{\mathcal{Y}} \Phi(\mathcal{Y}) = \min_{\mathcal{Y}} \sum_{i} \left\| \mathbf{y}_{i} - \sum_{j} w_{ij} \mathbf{y}_{j} \right\|^{2},$$
s.t. 
$$\frac{1}{N} \sum_{i} \mathbf{y}_{i} \mathbf{y}_{i}^{T} = \mathbf{I} \quad \text{and} \quad \sum_{i} \mathbf{y}_{i} = \mathbf{0} \quad \forall i \in \{1, 2, ..., N\}.$$
(10)

The objective can be equivalently stated as an eigenvalue problem. For this purpose, define  $\mathbf{E} = (\mathbf{I} - \mathbf{W})^T (\mathbf{I} - \mathbf{W})$  and set  $\tilde{\mathcal{Y}} = \mathcal{Y}^T$  (Cayton, 2005), such that:

$$\min_{\tilde{\mathcal{Y}}} trace(\tilde{\mathcal{Y}}^T \boldsymbol{E} \tilde{\mathcal{Y}}), \quad \text{s.t. } \frac{1}{N} \tilde{\mathcal{Y}}^T \tilde{\mathcal{Y}} = \boldsymbol{I} \quad \text{and} \quad \tilde{\mathcal{Y}}^T \mathbf{1} = \mathbf{0}.$$
 (11)

Again, the solution is found by eigenanalysis of the matrix encoding the intrinsic manifold structures. Note that the first constraint carries a factor  $\frac{1}{N}$  as originally proposed. In fact, any such quadratic form, provided its right hand side is of full rank, would suffice to ensure the embedding vectors actually span a d-dimensional space (Burges, 2010). The additional sum-zero condition is implicitly met by discarding the constant eigenvector associated with the bottom, zero eigenvalue (Ghojogh et al., 2020).

As mentioned before, the close resemblance to the optimization problem in LEM (equation 6) is not coincidental. Recall that LEM builds upon eigenfunctions of the Laplace-Beltrami operator  $\mathcal{L}$ . Belkin and Niyogi (2003) show that LLE approximates the eigenfunctions of the iterated form  $\frac{1}{2}\mathcal{L}^2$ , which are identical to those of  $\mathcal{L}$ . Therefore, with a somewhat different intuition, LLE essentially arrives at the same conclusion.

#### 4.1.3 Hessian Locally Linear Embedding (HLLE)

- Hessian instead of Laplacian (eigenmaps)
- Hessian instead of LS fit (LLE)

### 4.2 Semi-Supervised Locally Linear Embedding (SSLLE)

#### 4.2.1 Employment of Prior Information

- Why use labels in the first place?
- How will that help?
- Exact vs inexact knowledge

#### 4.2.2 Finding Prior Points

Prior points: take minmax approach from sparse MDS (Sparse multidimensional scaling using landmark points Vin de Silva and Joshua B. Tenenbaum 2004). Easy and deterministic after choosing seed value. Instead Euclidean distances, though, take geodesics as estimated in isomap.

can we view the prior info as some kind of active learning? like we choose some points to label in a hopefully cleverish way and then hand them to you (e.g., to look at some pictures instead of all droelf thousand)

#### 4.2.3 SSLLE Algorithm

• What is different wrt standard LLE?

## 4.3 Particular Challenges

A number of computational and design-related challenges arise from this procedure that must be faced in implementation.

Choice of intrinsic dimensionality. Until now, it has been assumed, rather implicitly, that the intrinsic dimension d of the data is a known parameter. This is obviously not always the case in practical applications. Some methods offer the advantage of estimating d in a built-in fashion. PCA, MDS and Isomap, for instance, typically show an indicative gap in their eigenvalue spectrum, distinctly pointing out the dimensions with the largest share of variability (Saul et al., 2006). For LLE, LEM and HLLE, no such tell-tale gap exists. While Sha and Saul (2005) have indeed drawn a mathematical relation between the respective eigenspectra in LLE and LEM and intrinsic data dimensionality, they immediately discarded this finding for practical

applications due to large computational overhead and lack of reliability in finite-sample situations. There have been various other proposals to tackle the problem of dimensionality estimation (for an extensive discussion, see for example Wissel (2017)). However, as the focus of this report lies on a semi-supervised method of manifold learning, it is mainly concerned with situations where prior knowledge of coordinates, and of d in particular, is actually available.

Choice of neighborhood size. Choosing the size of neighborhoods for graph approximation does pose a challenge. It is a standard hyperparameter optimization problem in which a trade-off between locality and overall approximation must be balanced. If neighborhoods are too small, the model will not be able to learn the global manifold structure; with overly large neighborhoods, it will forgo the advantages of locality and non-linearity and essentially behave like PCA (de Ridder and Duin, 2002). Describe applied approach

Robustness of eigendecomposition. Mainly problem in LLE (?)

#### Computational cost. Text

- Number of neighbors (diss grilli (referenced in lle manual) discusses regression model, ghojogh propose different things)
- Intrinsic dimensionality
- Singularity of gram matrix (LLE-specific?!)
- Large data (landmarks)

Comment on difficulty of finding neighbors in high dimensions What about using RF proximities for neighbor search? Unsupervised RF works with simulating new data from the estimated dist of the present ones, see e.g. https://horvath.genetics.ucla.edu/html/RFclustering/RFclustering/RandomForestHorvath.pdf, https://arxiv.org/pdf/2004.02121.pdf

## 5 Experiment Results

- 5.1 Experimental Design
- 5.1.1 Software Implementation
- 5.1.2 Evaluation Framework
- 5.2 Application to Synthetic Data
- 5.2.1 Data
- 5.2.2 Results
- 5.3 Sensitivity Analysis

#### 6 Discussion

Pros and Cons

Various extensions

See (van der Maaten et al., 2009) for extensive discussion of manifold learning Theoretical convergence? (e.g., ISOMAP has this)

Determination of d: actually requires to know d, right? Must be automatically known if prior points are known

Potential shortcoming: what if manifold is not well-sampled? Not a problem with synthetic data, but IRL. But probably problematic with all manifold approaches This is directly related to the COD – local methods require dense sampling (van der Maaten et al., 2009)

Also: generalization to new points (w/o recomputing everything) neighborhood-preserving propositions -> fundamental problem: except for prior points, it is deterministic (as opposed to generative approaches, such as autoencoders)

## 7 Conclusion

Lorem ipsum

# A Appendix

# **B** Electronic Appendix

Data, code and figures are provided in electronic form.

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