



# DISTANCE-BASED DIMENSIONALITY REDUCTION FOR BIG DATA

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#### Degree

Master's Degree in Data Science

#### Master's thesis

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#### Abstract

Dimensionality reduction aims to project a data set into a low-dimensional space. Many techniques have been proposed, most of them based on the inter-individual distance matrix. When the number of individuals is really large, the use of full distance matrices is prohibitive because of their quadratic memory complexity. There are algorithms that extend classical multidimensional scaling (a long-established dimensionality reduction method based on distances) to the big data setting. In this TFM, we adapt these algorithms to any generic distance-based dimensionality reduction method.

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# 1 Introduction, motivation, and objectives

#### 1.1 Introduction

Dimensionality Reduction (DR) techniques embed high-dimensional datasets into signficantly lower-dimensional spaces while preserving the structure of the original data. Their primary goal is to tackle the common challenges of working with high-dimensional data, such as sparsity (caused by the curse of dimensionality) or large computational and storage costs. That being said, DR is mainly used for visualization purposes. In this setting, complex high-dimensional data is projected into  $\mathbb{R}^2$ , making patterns and clusters more apparent.

Many DR methods have been proposed since principal component analysis (PCA) (being the standard linear method) was first introduced (Pearson, 1901), each with distinct approaches and objectives. In particular, non-linear techniques have turned out to be very useful thanks to their ability to preserve complex relationships. Some examples are classical multidimensional scaling (classical MDS) (Torgerson, 1952; Gower, 1966), local multidimensional scaling (local MDS) (Chen and Buja, 2009), Isomap (J. B. Tenenbaum, Silva, and Langford, 2000), t-distributed stochastic neighbor embedding (t-SNE) (Maaten and Hinton, 2008), uniform manifold approximation and projection (UMAP) (McInnes, Healy, and Melville, 2018) and autoencoders (Baldi and Hornik, 1989; Kramer, 1991). All these methods differ in how they define and maintain relationships between points, although most of them try to preserve global structure and local neighborhoods.

Despite their utility, DR methods face a few limitations. Many algorithms require computing and/or keeping in memory pairwise distances between all data points, resulting in quadratic time complexity and memory requirements. This becomes prohibitive for large datasets with millions of points. Parameter selection presents challenges too, since there is no general consensus on the best way to tune them nor how to measure the quality of an embedding with a validation set. The substantial time complexities of these algorithms makes k-fold cross-validation very costly as well. Finally, understanding which aspects of the data are preserved and which are distorted when reducing the dimensionality is crucial to correctly interpret the results.

#### 1.2 Motivation

The recent advent of large data has led to new challenges and technologies to face them. When the number of observations of a dataset is very big, distance-based DR methods become computationally prohibitive because of their quadratic time and memory complexities. For example, working with a dataset of 100,000 individuals would require to keep a distance-matrix of 10 billion floating-point numbers in memory. If double precision is used, then the computer in use shall have at least 80 GB of RAM.

Recently, Delicado and Pachón-García (2024) studied existing and new modifications of classical MDS to tackle big datasets, demonstrating significant improvements in computational efficiency while maintaining embedding quality. Even though most of them could not be generalized to arbitrary DR techniques, two of them followed more common approaches: divide-and-conquer and recursion. These consist in partitioning the distance matrix into submatrices small enough for the system to keep in main memory. Then, classical MDS is applied to every partition independently and the resulting embeddings are aligned and merged with Procustes transformations. That being said, the recursive approach suffers from low-quality embeddings because it might divide the distance matrix

into too small partitions. Therefore, we were interested in generalizing the divide-and-conquer proposal to DR methods beyond classical MDS.

# 1.3 Objectives

The primary goal of this thesis is to propose a divide-and-conquer framework for any generic distance-based dimensionality reduction method that effectively decreases the method's time and memory complexities. Specifically, we aim to:

- 1. Review the literature to analyze the properties and applications of the most used DR techniques.
- 2. Develop a generalized framework for distance-based DR methods that leverages the divide-and-conquer strategy and the Procrustes transformation to reduce time and memory complexities.
- 3. Implement and parallelize the proposed framework for specific DR algorithms such as non-classical MDS, Local MDS, Isomap and t-SNE.
- 4. Empirically evaluate the performance of the adapted algorithms in terms of computational efficiency, size limitations and quality of embeddings on benchmark datasets of varying sizes.
- 5. Compare the results with the traditional counterpart of each tested DR method.
- 6. Provide guidelines and best practices for selecting and tuning the proposed DR methods based on dataset characteristics and desired properties of the embedding.

The successful completion of these objectives will contribute to making dimensionality reduction techniques accessible for very large datasets, democratizing their use across scientific and industrial applications where data scale is a present challenge.

# 2 State of the art

Many nonlinear DR techniques and big data solutions discovered in recent years have lead to our divide-and-conquer proposal. Hence, we shall first review them to understand the advances they have achieved. In section 2.1, our goal is to review four dimensionality reduction techniques that we will later apply our divide-and-conquer framework to. Next, in section 2.2, we will showcase in more detail the work of Delicado and Pachón-García (2024) with respect to classical MDS in big data to motivate our choosing of the divide-and-conquer approach. Finally, sections 2.3 and 2.4 offer a discussion about three more solutions to the big data problem in DR and how they relate to our proposal: landmark Isomap (Silva and J. Tenenbaum, 2002), the out-of-core dimensionality reduction framework (Reichmann, Hägele, and Weiskopf, 2024) and novel Python t-SNE implementations.

## 2.1 A few dimensionality reduction techniques

#### 2.1.1 Non-classical MDS: the SMACOF algorithm

SMACOF (Scaling by MAjorizing a COmplicated Function) is a multidimensional scaling procedure that minimizes metric stress using a majorization technique (Joseph B. Kruskal, 1964b; Joseph B Kruskal, 1964a). Given the distance matrix  $\mathbf{D}_{\mathbf{X}} = (\delta_{ij})$  of the original high-dimensional data and its corresponding low-dimensional distance matrix  $\mathbf{D}_{\tilde{\mathbf{Y}}} = (d_{ij})$ , metric stress determines how much different  $\mathbf{D}_{\mathbf{X}}$  and  $\mathbf{D}_{\tilde{\mathbf{Y}}}$  are by the expression

$$STRESS_{M}(\mathbf{D_{X}}, \mathbf{D_{\tilde{Y}}}) = \sqrt{\frac{\sum_{i < j} (\delta_{ij} - d_{ij})^{2}}{\sum_{i < j} \delta_{ij}^{2}}}.$$

SMACOF (see Algorithm 1) is faster for this problem than general optimization methods, such as gradient descent, and it consists on iterative Guttman transformations (Guttman, 1968) applied to the embedded data.

## 2.1.2 LMDS

Local multidimensional scaling (LMDS) is a variant of non-classical multidimensional scaling that differs in how large distances are treated (Chen and Buja, 2009). Specifically, a repulsive term between distant points is added to the stress function to further separate points in the low-dimensional configuration (see Algorithm 2). Parameters  $\tau$  (which must be in the unit interval) and k may be tuned with k'-cross validation thanks to the LC (Local Continuity) meta-criteria (Chen and Buja, 2009). (POSSIBLE ANNEX)

#### 2.1.3 Isomap

Isomap (J. B. Tenenbaum, Silva, and Langford, 2000) is a nonlinear technique that preserves geodesic distances between points in a manifold (see Algorithm 3). The key insight of Isomap is that large distances between objects are estimated from the shorter ones by the shortest path length. Then, shorter and estimated-larger distances have the same importance in a final classical MDS step.

The only tuning parameter of Isomap is the bandwidth, which can be interpreted as the limiting distance of nearest neighbors,  $\varepsilon$ , or the amount of nearest neighbors per point, k. However, there is no consensus on what is the best method to choose it.

## **Algorithm 1** SMACOF

**Require:**  $\mathbf{D}_{\mathbf{X}} = (\delta_{ij})$ , the  $n \times n$  matrix of observed distances; q, the embedding's dimensionality;  $n_i$  the maximum number of iterations; and  $\varepsilon$ , the convergence threshold. **Ensure:**  $\mathbf{Y}$ , a configuration in a q-dimensional space.

- 1: Initialize at random  $\tilde{\mathbf{Y}}^{(0)} \in \mathbb{R}^{n \times q}$
- $2: k \leftarrow 0$
- 3: repeat
- Compute the distance matrix of  $\tilde{\mathbf{Y}}^{(k)}$ ,  $\mathbf{D}_{\tilde{\mathbf{Y}}^{(k)}}$ , with entries  $d_{ij}^k = \|\tilde{y}_i^{(k)} \tilde{y}_j^{(k)}\|$ Compute the metric stress:  $STRESS_M(\mathbf{D_X}, \mathbf{D_{\tilde{\mathbf{Y}}^{(k)}}})$
- 5:
- Compute the Guttman transform:  $\tilde{\mathbf{Y}}^{(k+1)} = n^{-1}\mathbf{B}(\tilde{\mathbf{Y}}^{(k)})\tilde{\mathbf{Y}}^{(k)}$  where  $\mathbf{B}(\tilde{\mathbf{Y}}^{(k)}) = (b_{ii}^k)$ : 6:

$$b_{ij}^k = \begin{cases} -\delta_{ij}/d_{ij}^k & \text{if } i \neq j \text{ and } d_{ij}^k > 0\\ 0 & \text{if } i \neq j \text{ and } d_{ij}^k = 0\\ -\sum_{j \neq i} b_{ij}^k & \text{if } i = j \end{cases}$$

- $k \leftarrow k + 1$
- 8: until  $k \ge n$  iter or  $|STRESS_M(D_{\mathbf{X}}, \tilde{\mathbf{Y}}^{(k-1)}) STRESS_M(D_{\mathbf{X}}, \tilde{\mathbf{Y}}^{(k)})| < \varepsilon$
- 9: return  $ilde{\mathbf{Y}}^{(k)}$

#### 2.1.4 t-SNE

t-distributed stochastic neighbor embedding (t-SNE) is a nonlinear dimensionality reduction technique that preserves local neighborhoods by modeling similarities between points as conditional probabilities (Maaten and Hinton, 2008). The difference between these probability distributions in high and low-dimensional spaces is then minimized (see Algorithm 4).

t-SNE focuses on retaining the local structure of the data while ensuring that every point  $\mathbf{v}_i$  in the low-dimensional space will have the same number of neighbors, making it particularly effective for visualizing clusters. The use of the Student's t distribution with 1 degree of freedom (or Cauchy distribution) in the low-dimensional space addresses the crowding problem by allowing dissimilar points to be modeled far apart (Maaten and Hinton, 2008). The characteristic parameter of t-SNE is the perplexity, which can be interpreted as the average effective number of neighbors of the high-dimensional datapoints  $\mathbf{x}_i$ . Typical values are between 5 and 50.

#### 2.2 Multidimensional scaling for big data

Delicado and Pachón-García (2024) compared four existing versions of MDS with two newly proposed (divide-and-conquer MDS and interpolation MDS) to handle large datasets. As can be seen in Figure 1, these can be grouped into four categories:

• Interpolation-based: landmark MDS, interpolation MDS and reduced MDS apply classical multidimensional scaling to a subset of  $l \ll n$  points and then interpolate the projection of the remaining data. They differ in how the interpolation is computed: landmark MDS uses distance-based triangulation; interpolation MDS, the l-points Gower interpolation formula; and reduced MDS, the 1-point Gower interpolation formula.

## Algorithm 2 LMDS

**Require:**  $\mathbf{D}_{\mathbf{X}} = (\delta_{ij})$ , the  $n \times n$  matrix of observed distances; q, the embedding's dimensionality; k, the size of neighborhoods; and  $\tau$ , the weight of the repulsive term.

**Ensure:**  $\mathbf{Y}$ , a configuration in a q-dimensional space.

- 1: Compute the symmetrized k-NN graph of  $\mathbf{D}_{\mathbf{X}}$ ,  $\mathcal{N}_{k}$ 2: Calculate  $t = \frac{|\mathcal{N}_{k}|}{|\mathcal{N}_{k}^{C}|} \cdot \operatorname{median}_{\mathcal{N}_{k}}(\delta_{ij}) \cdot \tau$
- 3: Minimize the modified stress function:

$$\tilde{\mathbf{Y}} = \min_{\mathbf{Y} \in \mathbb{R}^{n \times q}} \sum_{(i,j) \in \mathcal{N}_k} \left( \delta_{ij} - \|\mathbf{y}_i - \mathbf{y}_j\| \right)^2 - t \sum_{(i,j) \notin \mathcal{N}_k} \|\mathbf{y}_i - \mathbf{y}_j\|$$

## 4: return $\tilde{\mathbf{Y}}$

#### Algorithm 3 Isomap

**Require:**  $\mathbf{D}_{\mathbf{X}}$ , the  $n \times n$  matrix of observed distances; q, the embedding's dimensionality; and  $\varepsilon$ , the limiting distance of nearest neighbors, or k, the amount of nearest neighbors per point.

**Ensure:**  $\dot{\mathbf{Y}}$ , a configuration in a q-dimensional space.

- 1: Find the graph  $\mathcal{N}$  given by the radius  $\varepsilon > 0$  or the k-NN
- 2: Compute the distance matrix of  $\mathcal{N}$ ,  $\mathbf{D}_{\mathcal{N}}$
- 3: Apply classical MDS to  $\mathbf{D}_{\mathcal{N}}$ :  $\mathbf{Y} = \mathrm{MDS}(\mathbf{D}_{\mathcal{N}}, q)$
- 4: return Y
  - Approximation-based: pivot MDS approximates the SVD of the full inner product matrix with the SVD of the inner product matrix between a subset of  $l \ll n$ points and all the points in the dataset.
  - Divide-and-conquer: In divide-and-conquer MDS, the dataset is randomly partitioned into subsets of up to  $l \ll n$  points into which MDS is independently applied. Then, the resulting embeddings are aligned with Procrustes transformations.
  - Recursive: fast MDS is similar in spirit to divide-and-conquer MDS, but it partitions the data recursively.

#### 2.3Landmark Isomap and the out-of-core dimensionality reduction framework

Silva and J. Tenenbaum (2002) first introduced landmark MDS as a step in landmark Isomap, a variation of Isomap that reduced the time complexity from  $\mathcal{O}(n^3)$  to  $\mathcal{O}(n^2l)$ , where  $l \ll n$  is the amount of landmark points. Other big data versions of MDS can be used with Isomap similarly. Nonetheless, interpolation-based and approximation-based algorithms cannot be trivially generalized to other nonlinear dimensionality reduction methods because they depend on the eigendecomposition computed in classical MDS.

Later, Reichmann, Hägele, and Weiskopf (2024) proposed the out-of-core dimensionality reduction framework. Similar to interpolation MDS, this algorithm applies a DR method that produces a mapping between high- and low-dimensional spaces (i.e. PCA, MDS, t-SNE, UMAP, autoencoders) to a small subset of the data and then projects the

## Algorithm 4 t-SNE

**Require:**  $\mathbf{D}_{\mathbf{X}} = (\delta_{ij})$ , the  $n \times n$  matrix of observed distances; q, the embedding's dimensionality; Perp, the perplexity.

**Ensure:**  $\mathbf{Y}$ , a configuration in a q-dimensional space.

1: For every point  $\mathbf{x}_i$ , find  $\sigma_i > 0$  so that the conditional probability ditribution

$$p_{j|i} = \frac{\exp(-\delta_{ij}^2/2\sigma_i^2)}{\sum_{k \neq i} \exp(-\delta_{ik}^2/2\sigma_i^2)} \text{ if } i \neq j, \, p_{i|i} = 0$$

has perplexity  $Perp = 2^{-\sum_{j} p_{j|i} \log_2 p_{j|i}}$ 

- 2: Symmetrize conditional distributions:  $p_{ij} = \frac{p_{j|i} + p_{i|j}}{2n}$ 3: Consider Cauchy-distributed joint probabilities for the low-dimensional data  $\mathbf{y}_i$ :

$$q_{ij} = \frac{(1 + ||y_i - y_j||^2)^{-1}}{\sum_{h \neq k} (1 + ||y_h - y_k||^2)^{-1}}$$

4: Minimize the sum of Kullback-Leibler divergences between the joint distributions over all datapoints:

$$\tilde{\mathbf{Y}} = \min_{\mathbf{Y} \in \mathbb{R}^{n \times q}} \sum_{i} \sum_{j} p_{j|i} \log \frac{p_{j|i}}{q_{j|i}}$$

# 5: return Y

remaining datapoints in blocks. In order to obtain the aforementioned mappings, Reichmann, Hägele, and Weiskopf (2024) gathered a different projection mechanism for every method. PCA and autoencoders learn a parametric mapping between the original data and the embedding, so projecting new points is straightforward. For non-classical MDS, stress is minimized for a single point while keeping others fixed, a process known as single scaling in the literature (Basalaj, 1999). A similar strategy is used for t-SNE (Zhang et al., 2021) and UMAP (McInnes, Healy, and Melville, 2018), which leverage the k-NN of the projecting point to initialize the optimizer.

#### 2.4 openTSNE, an outstanding implementation of t-SNE

Maybe because of its popularity, t-SNE has been optimized for large data environments in Python through iterative implementations. P. G. Poličar, Stražar, and Zupan (2024) considered many of them and published openTSNE, a package that includes several t-SNE extensions "to address scalability issues and the quality of the resulting visualizations". Furthermore, they compared their proposal with those of other popular packages in serial and parallel configurations. In Figure 2 it can be seen that, thanks to these advances, the challenges of big data have already been solved in the specific case of t-SNE.

(POSSIBLE ANNEX)

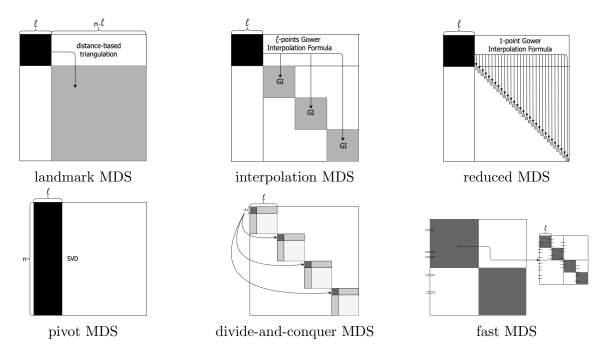


Figure 1: Schematic representation of the six MDS algorithms for big data described in Delicado and Pachón-García, 2024 (Source: original publication).

# 3 Specification and design of the solution

Our solution to the problem of applying dimensionality reduction techniques to large datasets consists in dividing the data into partitions small enough for the computer to process and then merge their embeddings. As stated earlier, we follow the same approach as Delicado and Pachón-García (2024). By means of an efficient alignment procedure named *Procustes transformation*, we manage to orient every embedding to a specific alignment, with the goal that the final embedding will be coherent. Therefore, even though partitioning the data might induce errors in the embedding's structure, our approach allows any system to tackle arbitrarily large datasets and leverage parallelization.

In order to preserve the topology of the partition's embedding, we narrow Procustes transformations to rigid motions; that is, rotations and reflections. Moreover, we diminish the overhead computation of merging the embeddings by computing the transformation matrix with only a few datapoints. Specifically, if partitions have l (or l-1) points, we sample c < l from the first partition and stack them temporarily to every other partition. Later, we embed the first partition and extract the result of the c sampled points,  $\tilde{\mathbf{Y}}_{\mathcal{C}}$ . Afterward, for every remaining partition, we project the stacked data with the chosen DR method and separate the c points corresponding to the first partition,  $\tilde{\mathbf{Y}}_{\mathcal{C}}^{(i)}$ , from those of the current partition,  $\tilde{\mathbf{Y}}_{c}^{(i)}$  and apply it to the larger matrix  $\tilde{\mathbf{Y}}_{i}$ , all without needing to process two full partitions.

For a more detailed specification of our solution, see Algorithm 5, which depicts the divide-and-conquer dimensionality reduction algorithm, and section 3.1, which shows how the Procrustes transformation can be obtained.

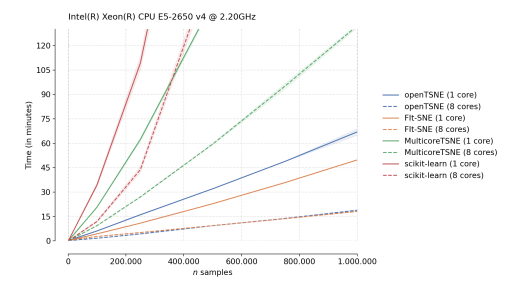


Figure 2: Benchmark of the best open-source t-SNE Python implementations by P. Poličar, 2023 (Source: original webpage).

## 3.1 Orthogonal Procrustes transformation

Our problem of aligning the partitions' embeddings is known in the literature as the *Procustes problem* (Borg and Groenen, 2005). Depending on the kind of fitting desired, many solutions can be found. For example, orthogonal transformations consist of rotations and reflections, but one may also desire dilations and shifts. In fact, the transformation could be any linear distortion.

That being said, in order to preserve the structure of every partitions' embedding, we considered best to narrow the problem down to rigid motions, or in other words, rotations and reflections. Now, let  $\mathbf{A} \in \mathbb{R}^{c \times q}$  be the target configuration ( $\tilde{\mathbf{Y}}_{\mathcal{C}}$  in Algorithm 5) and  $\mathbf{B} \in \mathbb{R}^{c \times q}$  the corresponding testee ( $\tilde{\mathbf{Y}}_{\mathcal{C}}^{(i)}$  in Algorithm 5). We wish to fit  $\mathbf{B}$  to  $\mathbf{A}$  by rigid motions. That is, we want to find the best orthogonal matrix  $\mathbf{T}$  such that  $\mathbf{A} \simeq \mathbf{B}\mathbf{T}$ .

To measure the  $\simeq$  relation, we may use the sum-of-squares criterion L. Then, the transformation  $\mathbf{T}$  should be chosen to minimize L. Expressed in matrix notation, our problem is

$$\min_{\mathbf{T} \in O(q)} L(\mathbf{T}) = \min_{\mathbf{T} \in O(q)} \operatorname{tr}(\mathbf{A} - \mathbf{B}\mathbf{T})(\mathbf{A} - \mathbf{B}\mathbf{T})',$$

where O(q) is the orthogonal group in dimension q.

 $(POSSIBLE\ ANNEX)$ 

By expanding the expression of  $L(\mathbf{T})$  and applying a lower bound inequality on traces derived by Kristof, 1970, Borg and Groenen, 2005 found a global solution to the minimization problem. Let  $\mathbf{U}\Sigma\mathbf{V}'$  be the singular value decomposition of  $\mathbf{A}'\mathbf{B}$ , where  $\mathbf{U}'\mathbf{U} = \mathbf{I}, \mathbf{V}'\mathbf{V} = \mathbf{I}$ , and  $\Sigma$  is the diagonal matrix with the singular values. Then,  $L(\mathbf{T})$  is minimal if

$$VU'$$
.

Therefore, the Procrustes procedure we used would be as follows:

#### Algorithm 5 Divide-and-conquer dimensionality reduction

**Require:**  $\mathbf{X} \in \mathbb{R}^{n \times p}$ , the high-dimensional data;  $\mathcal{M}$ , the DR method; l, the partition size; c, the amount of connecting points; q, the embedding's dimensionality; and arg,  $\mathcal{M}$ 's specific parameters.

**Ensure:**  $\tilde{\mathbf{Y}}$ , a configuration in a q-dimensional space.

- 1: if  $n \leq l$  then
- 2: **return**  $\mathcal{M}(\mathbf{X}, q, arg)$
- 3: end if
- 4: Partition data into k subsets:  $\mathcal{P} = \{\mathcal{P}_1, \mathcal{P}_2, \dots, \mathcal{P}_k\}$  where  $|\mathcal{P}_i| \leq l$  for all i
- 5: Sample c connecting points from  $\mathcal{P}_1$ :  $\mathcal{C} \subset \mathcal{P}_1$  with  $|\mathcal{C}| = c$
- 6: Apply DR method to first partition:  $\tilde{\mathbf{Y}}_1 = \mathcal{M}(\mathbf{X}_{\mathcal{P}_1}, q, arg)$
- 7: Extract embedding of C:  $\tilde{\mathbf{Y}}_{\mathcal{C}} = \tilde{\mathbf{Y}}_{1}[C,:]$
- 8: **for** i = 2 to k **do**
- 9: Stack connecting points to current partition:  $\mathbf{X}_{\text{stack}} = [\mathbf{X}_{\mathcal{C}}; \mathbf{X}_{\mathcal{P}_i}]$
- 10: Project stacked data:  $\tilde{\mathbf{Y}}_{\text{stack}} = \mathcal{M}(\mathbf{X}_{\text{stack}}, q, arg)$
- 11: Separate embedding of  $\mathbf{X}_{\mathcal{C}}$ :  $\tilde{\mathbf{Y}}_{\mathcal{C}}^{(i)} = \tilde{\mathbf{Y}}_{\text{stack}}[1:c,:]$  and  $\tilde{\mathbf{Y}}_i = \tilde{\mathbf{Y}}_{\text{stack}}[(c+1):,:]$
- 12: Align projection using Procrustes:  $\tilde{\mathbf{Y}}_i = \text{Procrustes}(\tilde{\mathbf{Y}}_{\mathcal{C}}, \tilde{\mathbf{Y}}_{\mathcal{C}}^{(i)}, \tilde{\mathbf{Y}}_i)$
- 13: end for
- 14: Combine all projections:  $\tilde{\mathbf{Y}}' = [\tilde{\mathbf{Y}}_1; \tilde{\mathbf{Y}}_2; \dots; \tilde{\mathbf{Y}}_k]$
- 15: Reorder rows to match original ordering:  $\tilde{\mathbf{Y}}' = \tilde{\mathbf{Y}}'[\text{order},:]$
- 16: Apply PCA to center and rotate for maximum variance:  $\tilde{\mathbf{Y}} = \text{PCA}(\tilde{\mathbf{Y}}', q)$
- 17: return  $\tilde{\mathbf{Y}}$

## Algorithm 6 Procrustes procedure

**Require:**  $\mathbf{A} \in \mathbb{R}^{c \times q}$ , the target matrix;  $\mathbf{B} \in \mathbb{R}^{c \times q}$ , the testee matrix; and  $\mathbf{C} \in \mathbb{R}^{m \times q}$ , the matrix to transform.

Ensure: C', the matrix C after alignment.

- 1: Multiply  $\mathbf{M} = \mathbf{A}'\mathbf{B}$
- 2: Compute singular value decomposition:  $\mathbf{U}, \mathbf{\Sigma}, \mathbf{V}' = \text{SVD}(\mathbf{M})$
- 3: Construct orthogonal matrix: T = VU'
- 4: Align C: C' = CT
- 5: return C'

# 4 Development of the proposal

# 4.1 Python implementation of divide-and-conquer DR

Given that R and Python are the standard programming languages in the data science field, we chose them as appropriate to implement the divide-and-conquer DR algorithm. Initially, we aimed to develop and publish an R library because the thesis directors already had experience with the language. However, after reviewing the literature on DR for big data, we realized that many solutions were implemented in Python instead (Reichmann, Hägele, and Weiskopf, 2024). So, in order to leverage the existing coding ecosystem, we switched to Python. From that moment onward, we documented the code development on an open-source GitHub repository (https://github.com/airdac/TFM\_Adria). This system allowed us to easily update and share our implementations and experiments.

With time, the Pyhton functions and classes we have written in different modules have become structured in a directory tree, hence taking the form of a package. Even though our project has not been published in any Python package index yet, it effectively works as a library. The main function is divide\_conquer, which implements Algorithm 5 in parallel through the concurrent.futures module. divide\_conquer also depends on private methods and requires a DRMethod object as one of its arguments. This class, inherited from enum.Enum, lists the supported DR methods in our package, which can be called through the get\_method\_function method.

We have implemented and tested four DR algorithms: SMACOF, LMDS, Isomap and t-SNE. All but LMDS are wrappers to methods coded in other efficient and parallelized Python libraries. Specifically, we used the sklearn.manifold module (Pedregosa et al., 2011) for Isomap and SMACOF and openTSNE (P. Poličar, 2023) for t-SNE. LMDS, on the other hand, is a less popular method and, up to our knowledge, it has no public Python implementation at the moment. However, the R library smacofx (Leeuw and Mair, 2009) does, so we translated it to Python with the help of the scipy.spatial.distance and sklearn.neighbors modules. This also allowed us to optimize the LMDS runtime with the numba just-in-time compiler (Lam, Pitrou, and Seibert, 2015; Aycock, 2003).

As shown in section 5, our implementation does not add a significant computation overhead when compared to standard DR techniques. In fact, divide-and-conquer DR leverages parallelization and provides a significant speed up to standard DR methods. Most DR techniques have quadratic time complexity (Joseph B. Kruskal, 1964b; Joseph B Kruskal, 1964a; Chen and Buja, 2009; Maaten and Hinton, 2008), except for Isomap, whose computation time is  $\mathcal{O}(n^2\log(n))$  (J. B. Tenenbaum, Silva, and Langford, 2000). Meanwhile, divide-and-conquer DR applies a DR method on n/l partitions of l individuals, resulting in linear time complexity:  $\mathcal{O}(nl)$  usually and  $\mathcal{O}(nl\log(l))$  for Isomap. The only DR method we have not been able to accelerate has been t-SNE, given that it already was thouroughly optimized in the openTSNE library (P. G. Poličar, Stražar, and Zupan, 2024). Finally, since we only keep in memory the inter-individual distance matrix of the partitions being embedded at each moment, our DR framework reduces space complexity from  $\mathcal{O}(n^2)$  to  $\mathcal{O}(l^2)$ .

# 4.2 Experiment's methodology and parameter tuning

Once our framework was developed and we had implemented a few DR methods to test it, we run a series of experiments on the MNIST (Cohen et al., 2017) and the swiss roll (J. B. Tenenbaum, Silva, and Langford, 2000) datasets. We downloaded the MNIST dataset

from the NIST webpage (Standards and Technology, 2024) and generated points on the swiss roll manifold with the sklearn.datasets.make\_swiss\_roll function (Pedregosa et al., 2011). All these experiments were executed, logged, plotted and dicussed on Jupyter notebooks (Kluyver et al., 2016) with the help of the Python packages numpy (Harris et al., 2020), pandas (McKinney, 2010), matplotlib.pyplot (Hunter, 2007), os, time, sys, shutil, logging and warnings. Furthemore, results were serialized with the pickle module in final tests.

The general experimental methodology we used consisted in a series of steps. Given that working with big datasets is costly, we randomly sampled a subset of 1000 points and embedded them into  $\mathbb{R}^2$  with a traditional DR technique. To distinguish it from its divide-and-conquer variation, we called it a *bare* technique. Then, before increasing the size of the dataset, we tuned the parameters of the method by applying it to the same dataset with different value combinations. In the tuning process, we took into account the runtime and embedding quality for each value combination. Our goal was to find parameter values that provided a good trade-off between speed and preserving the structure of the data. In section 5, we enter in more detail on the qualitative assessment of each dataset's embeddings.

Once parameters were tuned with a small enough subset of the data for our system to handle, we applied divide-and-conquer DR to larger datasets with partitions of l = 1000 points. In comparison, when we applied the bare techniques to big datasets, execution would crash because of a lack of main memory. Hence, as shown in section 5, we managed to extend the functionality of DR techniques to arbitrarily large datasets.

The MNIST dataset consisted of 345035 images represented in  $\mathbb{R}^{784}$ . On the other hand, the swiss roll dataset was sampled from a bidimensional object in  $\mathbb{R}^3$  shaped like a sheet of paper curved into a spiral along its longest axis. In order to measure the time complexity of divide-and-conquer DR, we randomly generated datasets with sizes ranging between  $10^3$  and  $10^8$  points. A good embedding of the swiss roll into  $\mathbb{R}^2$  should intuitively be a rectangle, as if the swiss roll were unfolded. In MNIST, we would like to obtain 10 clearly distinct clusters corresponding to each digit present in the dataset.

# 5 Experimentation and evaluation of the proposal

The goal of this section is to present the experiments we have conducted on the divide-and-conquer DR algorithm to assess its quality and performance. Following the methodology described in section 4.2, we were able to corroborate the space and time complexity of the procedure and understand better it's embedding mechanism. We chose the swiss roll and MNIST datasets for the tests because of their popularity, which makes it easier to compare our method with others in the literature, and because of how well Isomap and t-SNE embedded them. That allowed us to go even further and successfully unfold a 10<sup>8</sup> points swiss roll with divide-and-conquer Isomap. On the other hand, the classification task in the MNIST dataset proved more complicated to our algorithm, which was slower and separated digits worse than openTSNE's implementation of t-SNE.

The experimentation and evaluation process also provided us some insights on the standard SMACOF, LMDS, Isomap and t-SNE techniques. For example, we realized that LMDS, a nonlinear method intended to overcome the limitations of the SMACOF algorithm (Chen and Buja, 2009), was not able to unfold the swiss roll dataset. Even after having tuned the k and  $\tau$  parameters (see Algorithm 2), its embedding was porous and irregular instead of uniform and rectangular (see Figure 11). Further research on this problem might lead to the development of a variation of LMDS that reduces the dimensionality of the swiss roll better.

Finally, we will describe the computer system we performed the tests on. Even though our main development computer was a Macbook Pro (14-inc, Nov 2023) with 16 GB of RAM and the Apple M3 chip, we noticed that the concurrent.futures module, which parallelized the execution of Algorithm 5, did not work in Mac computers with ARM chipsets. Therefore, we ended up using a Windows system. Specifically, our PC was an Asus ROG G513QM-HF026 laptop with the AMD Ryzen 7 5800H CPU, 16 GB of DDR4-3200MHz RAM, an SSD M.2 NVMe PCIe 3.0 and an NVIDIA RTX 3060 GPU.

### 5.1 Initial runtime benchmarks

Isomap was the first DR method we implemented into our divide-and-conquer framework, so it also was the first method we benchmarked. We tested divide-and-conquer Isomap on the swiss roll dataset with different parameter combinations and with parallel and serial computation. However, in all tests the number of connecting points for the Procrustes transformation was 100. We chose this number because, based on the thesis directors' experience on big data MDS (Delicado and Pachón-García, 2024), it guaranteed good links between partitions' embeddings and efficient computations when  $1,000 \le l \le 10,000$ . Figure 3 shows the average runtimes of 20 experiments with different sets of parameters and dataset sizes. Parameters were previously tuned to ensure embeddings would preserve the structure of the data. As described in section 4.2, we applied bare Isomap to swiss rolls of 1,000, 3,162 and 10,000 points, following a logarithmic sequence. Notice that we increased the number of neighbors k for larger values of l because partitions were denser.

After some experimentation, we realized that if l = 3162 and k = 10, the embedding was nearly perfect no matter the amount of individuals (see Figure 4). Meanwhile, when l = 10,000 and k = 15, quality was similar and time was significantly larger, so we used l = 3162 and k = 10 for the largest datasets. This way, we managed to embed  $10^8$  three-dimensional points into the Euclidean plane in about 3 hours. Hence, we showed that divide-and-conquer Isomap is capable of handling arbitrarily large datasets on a standard

computer while maintaining the quality of the embedding.

Regarding parallelization, we can observe in Figure 3 that it effectively reduces the time complexity of divide-and-conquer DR, although its overhead slows down the algorithm when  $n \leq 10^4$ . Overall, results show that divide-and-conquer Isomap is linear in time with respect to n.

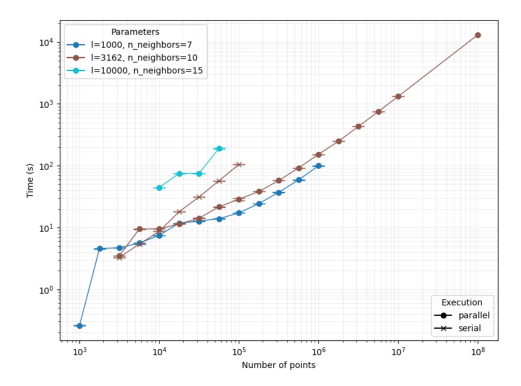


Figure 3: Runtime (s) of divide-and-conquer Isomap averaged over 20 experiments. Tests were performed on datasets generated on the swiss roll manifold (Spiwokv, 2007) with sizes ranging from  $10^3$  to  $10^8$ . Data was embedded into  $\mathbb{R}^2$  with different parameter combinations and c = 100. Runtime in parallel and serial execution is also compared.

Afterwards, we tested divide-and-conquer t-SNE on the same datasets (see Figure 5). However, t-SNE performed notably slower than Isomap on the swiss roll, so we only run its divide-and-conquer variation with one parameter combination, c = 100, l = 1,000, Perp = 30,  $n\_iter = 250$ .  $n\_iter$  is the number of iterations carried out to minimize the Kullback-Leibler divergence (Kullback and Leibler, 1951).

Even though divide-and-conquer t-SNE is about two orders of magnitud slower than divide-and-conquer Isomap, time complexity is linear as well, proving the expected results. The quality of the embedding, on the other hand, is very low. See Figure 6 to observe that the structure of the data is broken into separate parts and the spiral shape is not unfolded.

# 5.2 Analysis of possible overheads in divide-and-conquer DR

One reasonable question to formulate about the divide-and-conquer approach is whether splitting the data and merging the resulting embeddings constitue a significant computational cost in comparison to reducing each partition's dimensionality. Theoretically, fractionating the data should be rapid. As for the merging of partitions' embeddings, in section 3 we depicted how we align and overlap them with Procustes transformations

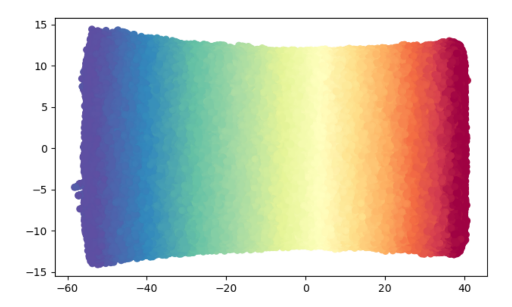


Figure 4: Bidimensional embedding of a  $10^8$  points swiss roll dataset (Spiwokv, 2007) computed by divide-and-conquer Isomap with l = 3, 162, c = 100 and k = 10. Color represents the angle of rotation along the swiss roll spiral.

between each embedding and one in specific (the first one). We intentionally find a rigid transformation only with a random subset of c < l points to accelerate its computation, and then multiply the full partition's embedding with the  $q \times q$  matrix of the transformation. Therefore, the overhead of merging embeddings should be insignificant.

Table 1 shows the results of an experiment where each part of the divide-and-conquer DR algorithm was independently timed. We uniformly sampled 5,000 points from the MNIST dataset and embedded them into the Euclidean plane with divide-and-conquer SMACOF. The arguments used were l=1000, c=100,  $n\_iter=300$ ,  $\varepsilon=0.001$ . As it was expected, neither partitioning the dataset nor aligning the partial embeddings entail a noteworthy overhead in divide-and-conquer DR. Indeed, the prior and the latter were about 5,506 times and 50,710 times swifter than embedding all partitions with SMACOF, respectively.

| Operation    | Divide                | Embed | Merge                 |
|--------------|-----------------------|-------|-----------------------|
| Duration (s) | $6.88 \times 10^{-3}$ | 37.89 | $7.47 \times 10^{-4}$ |

Table 1: Runtime (s) of each step of divide-and-conquer SMACOF on a 5000-point random subset of MNIST. The arguments used were l = 1000, c = 100,  $n\_iter = 300$ ,  $\varepsilon = 0.001$ .

# 5.3 Divide-and-conquer SMACOF

#### 5.3.1 Swiss roll

In this experiment, we applied bare and divide-and-conquer SMACOF to swiss rolls of different sizes. The largest dataset the standard method could handle ended up having 7,500 individuals and taking about 16 minutes to calculate. When we applied bare SMA-

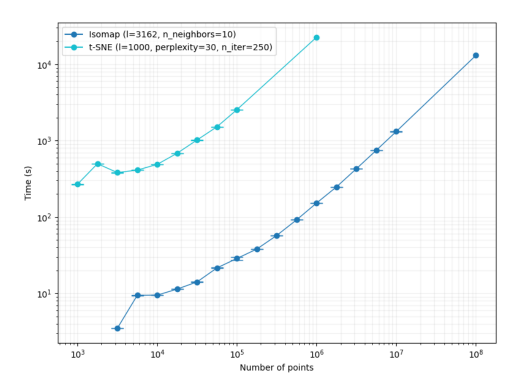


Figure 5: Runtime (s) of divide-and-conquer Isomap and divide-and-conquer t-SNE averaged over 20 experiments. Tests were performed on datasets generated on the swiss roll manifold (Spiwokv, 2007) with sizes ranging from  $10^3$  to  $10^8$ . Data was embedded into  $\mathbb{R}^2$  with different parameter combinations and c = 100.

COF to a 10,000 points swiss roll, the system crashed because it lacked main memory. Therefore, we show in Figure 7 the results of the test with 7,500 points.

The spiraled shape of both embeddings suggest that SMACOF cannot identify the intrinsic bidimensionality of the swiss roll. Moreover, SMACOF condenses more the outer part of the manifold than the inner one and presents rugged edges. Regarding the qualitative differences between bare and divide-and-conquer versions, there is not much to be said. Divide-and-conquer SMACOF makes a good job aligning the partial configurations and obtains a similar embedding than bare SMACOF up to a 180° rotation.

To conclude, even though the SMACOF algorithm is not capable of properly embedding the swiss roll, we can see a clear advantage in using it with the divide-and-conquer framework on other datasets. Indeed, divide-and-conquer SMACOF is about 22 times faster than the bare method while returning a similar low-dimensional configuration.

#### 5.3.2 MNIST

Next, we try with a subpartition of 5000 images, since that worked properly with the swiss roll. We tried a few parameter combinations and obtained the best one for ... Figure 8 shows it is not bad, although 4, 7 and 9 are expectedly mixed. 5 could also be confused by 2 and 8 and viceversa. All normal here, their shapes are indeed similar.

Maybe I should talk about dimension correlation between SMACOF and divide-and-conquer SMACOF. The point of this is the framework, not the DR method, you know?

Total correlation of dim1: -0.8996107183598692 Total correlation of dim2: 0.8785425518794896

Merda, i falta el temps.

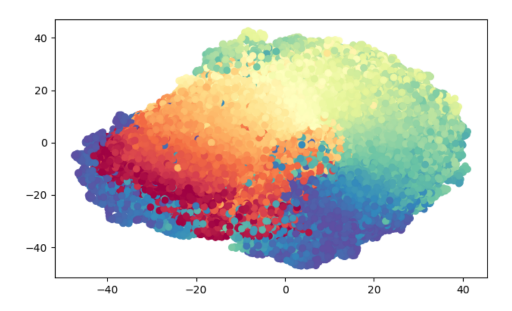


Figure 6: Bidimensional embedding of a  $10^6$  points swiss roll dataset (Spiwokv, 2007) computed by divide-and-conquer t-SNE with  $l=1,000,\,c=100,\,Perp=30$  and  $n\_iter=250$ . Color represents the angle of rotation along the swiss roll spiral.

# 5.4 Divide-and-conquer LMDS

#### 5.4.1 Swiss roll

Consider a swiss roll of 1,000 points. Well, LMDS makes trash out of it. Figure 11.

POSSIBLE ANNEX EXPLICANT EL QUE HEM PROVAT PER MILLORAR LMDS (LMDS\_examples.ipynb)

#### 5.4.2 MNIST

Same 5000 points MNIST subset, mate. Oh, but this is very bad... Like, worse than SMACOF HAHAHAHAHA.

Parla del temps i la correlació dimensionaaaaaal.

Total correlation of dim1: 0.9489962757561992 Total correlation of dim2: 0.8206029210441554

## 5.5 Divide-and-conquer Isomap

#### 5.5.1 Swiss roll

Aaaaah... I don't have the same here. Gotta get those damn plots. Well we've seen this before, in the initial benchmark. Just go see Figure 4.

#### 5.5.2 MNIST

There you go, Figure 14. It's just so bad HAHAHAHAH I can't handle this anymore. Total correlation of dim1: 0.9195527127381181 Total correlation of dim2: -0.8623571714329816

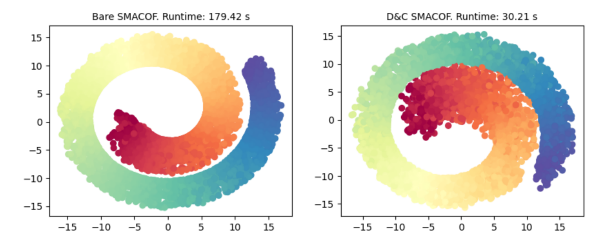


Figure 7: Comparison of the bidimensional embeddings of a 7,500 points swiss roll dataset (Spiwokv, 2007) by bare (left) and divide-and-conquer (right) SMACOF. The arguments used were  $n\_iter = 300$ ,  $\varepsilon = 0.001$  and in divide-and-conquer there also were l = 1000 and c = 100. Color represents the angle of rotation along the swiss roll spiral.

## 5.6 Divide-and-conquer t-SNE

#### 5.6.1 Swiss roll

t-SNE is specially useful for visualization, so we will focus on MNIST. Nonetheless, here's the swiss roll. Figure 6.

#### 5.6.2 MNIST

EXPLICAR EXEMPLE DUES PARTICIONS: LA QUALITAT POT SER DOLENTA PERQUÈ ELS CLUSTERS TENEN UNA TOPOLOGIA DIFERENT

Total correlation of dim1: 0.8170763484715383 Total correlation of dim2: 0.32817864544487557

AAAAAAA bé, l'original no només és millor; també és més ràpid! Va, mira què passa quan busquem una combinació de paràmetres pel t-SNE que funcioni bé amb 1000 dibuixets de números. Figure 18.

Conclusió: no hem guanyat aquesta batalla perquè la competència és un F1 i el nostre és un turisme. Però oi que no t'emportaries un F1 a la muntanya? Aaaaah-mic!

I ja estaria, xd. Anem bé de pàgines, béeeeeeeee.

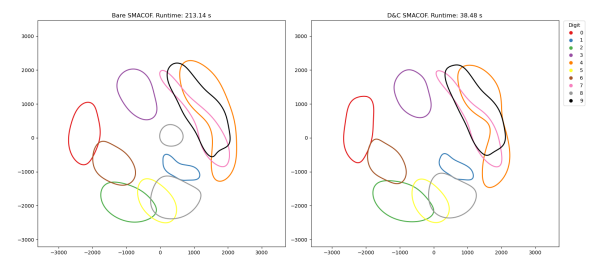


Figure 8: Kernel density estimation of the bidimensional embeddings of a 5,000 points subset of MNIST (Cohen et al., 2017) by bare (left) and divide-and-conquer (right) SMA-COF. The arguments we used were  $n\_iter = 300$ ,  $\varepsilon = 0.001$  and in divide-and-conquer there also were l = 1000 and c = 100. We applied a Procrustes transformation to align both configurations. Countour lines are at 70% of the maximum estimated density for each digit and embedding.

# 6 Analysis of sustainability and ethical implications

#### DESCRIPTION OF THIS SECTION FROM THE REGULATION:

It must include an analysis of the impact of the following gender-related technical aspects:

- issues related to data management and analysis
- issues related to equity, where possible biases are identified and assessed both in the data and in the processes carried out in relation to data management and analysis
- actions carried out to eliminate or mitigate such biases

#### ACTUAL CONTENT:

- D&C can be used in normal computers, while traditional DR methods require supercomputers to work on big datasets. Hence, we will reduce computing emissions.
- Maybe (it needs testing) DR methods could emphasize biases in the data. This happens because when projecting only a few coordinates, small clusters could be left behind in the remaining not projected coordinates and do not show in the final embedding. Hence, in theory, small communities could become invisible.

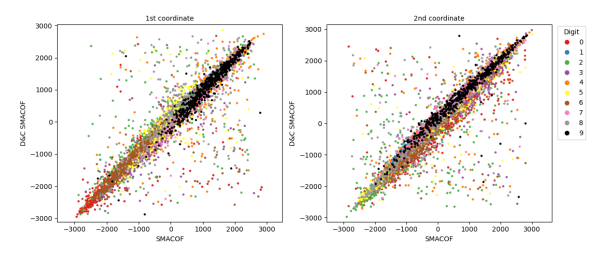


Figure 9: Scatter plots of first (left) and second (right) coordinates in the embeddings represented in Figure 8. SMACOF is compared against divide-and-conquer SMACOF.

# 7 Conclusions

Està molt bé.

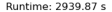
# 7.1 Future work

Crear un paquet de Python.

Comparar amb Reichmann, Hägele, and Weiskopf (2024).

Investigar perquè LMDS falla a l'Isomap.

Fer més experiments per veure com afecta c al rendiment i la qualitat. Aplicar a més datasets.



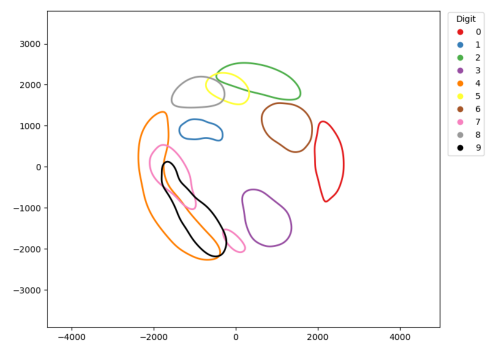


Figure 10: Kernel density estimation of the bidimensional embeddings of the MNIST dataset (Cohen et al., 2017) by divide-and-conquer SMACOF. The arguments used were  $n\_iter = 300$ ,  $\varepsilon = 0.001$ , l = 1000 and c = 100. Countour lines are at 70% of the maximum estimated density for each digit and embedding.

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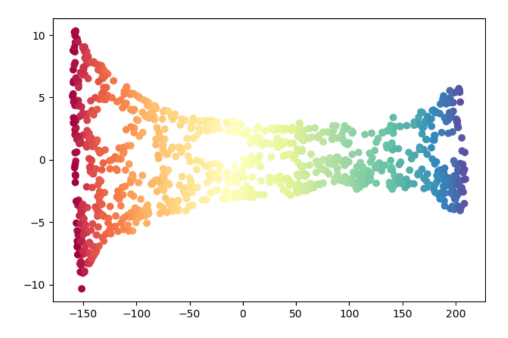


Figure 11: Bidimensional embedding of a 1,000 points swiss roll dataset (Spiwokv, 2007) computed by LMDS with k = 10 and  $\tau = 0.1$ . Color represents the angle of rotation along the swiss roll spiral.

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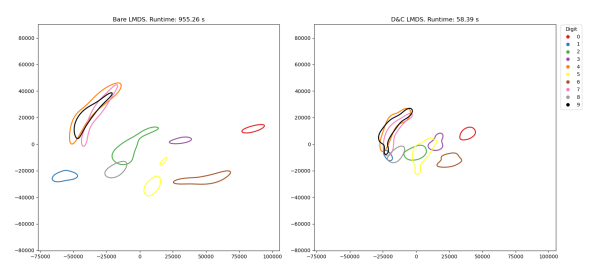


Figure 12: Kernel density estimation of the bidimensional embeddings of a 5,000 points subset of MNIST (Cohen et al., 2017) by bare (left) and divide-and-conquer (right) LMDS. The arguments used were  $k=10,\,\tau=1$  and in divide-and-conquer there also were l=1000 and c=100. We applied a Procrustes transformation to align both configurations. Countour lines are at 70% of the maximum estimated density for each digit and embedding.

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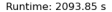
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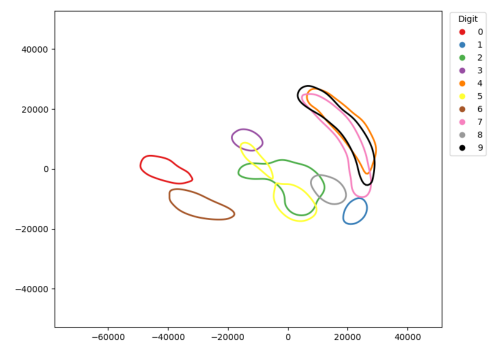


Figure 13: Kernel density estimation of the bidimensional embeddings of the MNIST dataset (Cohen et al., 2017) by divide-and-conquer LMDS. The arguments used were k = 10,  $\tau = 1$ , l = 1000 and c = 100. Countour lines are at 70% of the maximum estimated density for each digit and embedding.

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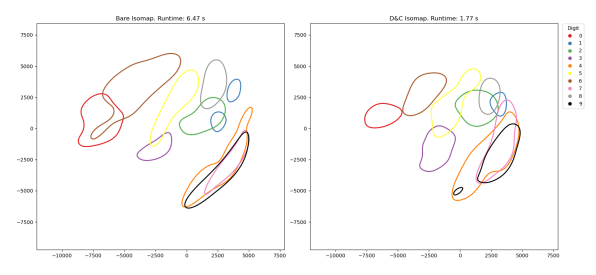


Figure 14: Kernel density estimation of the bidimensional embeddings of a 5,000 points subset of MNIST (Cohen et al., 2017) by bare (left) and divide-and-conquer (right) Isomap. The arguments used were k=5 and in divide-and-conquer there also were l=1000 and c=100. We applied a Procrustes transformation to align both configurations. Countour lines are at 70% of the maximum estimated density for each digit and embedding.

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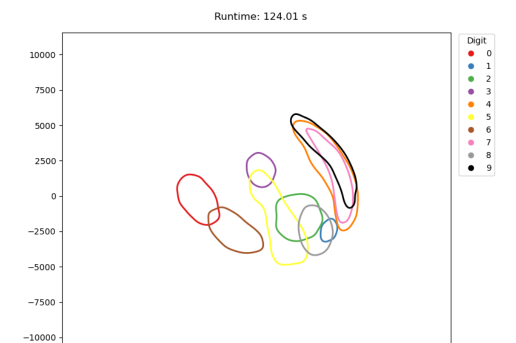


Figure 15: Kernel density estimation of the bidimensional embeddings of the MNIST dataset (Cohen et al., 2017) by divide-and-conquer Isomap. The arguments used were  $k=5,\,l=1000$  and c=100. Countour lines are at 70% of the maximum estimated density for each digit and embedding.

5000

10000

-5000

-10000

-15000

# 8 Annexes

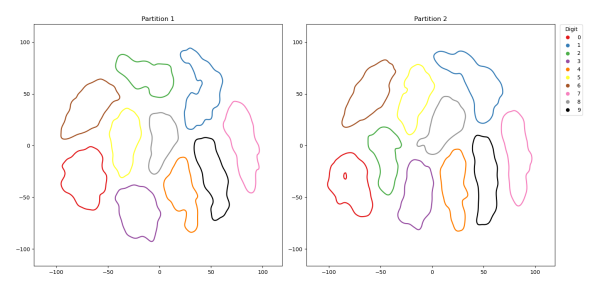


Figure 16: Kernel density estimation of the bidimensional embeddings of two halves of the MNIST dataset (Cohen et al., 2017). Data was randomly ordered before being splitted. The DR method used was divide-and-conquer t-SNE with Perp = 30. Countour lines are at 70% of the maximum estimated density for each digit and embedding.

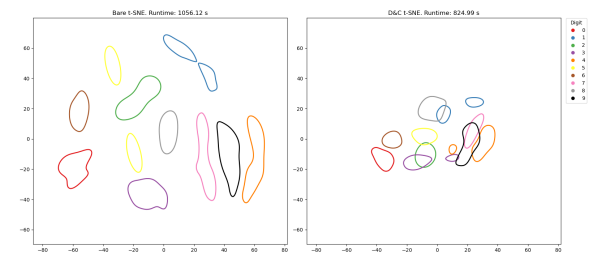


Figure 17: Kernel density estimation of the bidimensional embeddings of a 5,000 points subset of MNIST (Cohen et al., 2017) by bare (left) and divide-and-conquer (right) t-SNE. The arguments used were Perp=30 and in divide-and-conquer there also were l=1000 and c=100. We applied a Procrustes transformation to align both configurations. Countour lines are at 70% of the maximum estimated density for each digit and embedding.

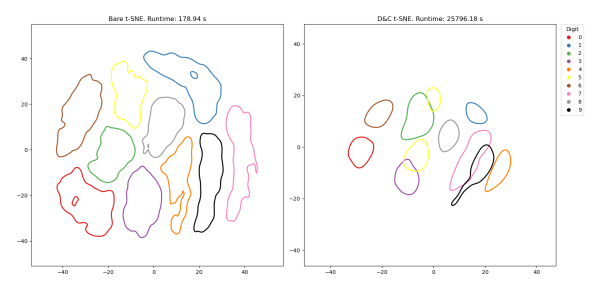


Figure 18: Kernel density estimation of the bidimensional embeddings of the MNIST dataset (Cohen et al., 2017) by bare (left) and divide-and-conquer (right) t-SNE. The arguments used were  $Perp=20,\ n\_iter=100$  and in divide-and-conquer there also were l=1000 and c=100. Countour lines are at 70% of the maximum estimated density for each digit and embedding.