Assignment 3: Unsupervised Learning

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* 1. Problem description

For this unsupervised learning assignment, we are given five single-cell synthetic datasets, each comprising the gene expression profiles of 200 cells with 200 genes. The objective is to create a data analysis pipeline that takes a dataset file as input and performs dimensionality reduction, clustering, and visualization of the results.

* 1. Pipeline implementation
     1. Data pre-processing

As a preprocessing step, data normalization was performed on all 5 datasets. Data normalization ensures that each feature is on the same scale, reducing the impact of outliers and extreme values. Normalization helps improve the performance and reliability of a machine learning model. In this study, we used the *Standardization Scaling,* subtracting the mean of each observation and dividing it by the standard deviation.

* + 1. Dimensionality reduction

**PCA** transforms the original expression data into a new set of uncorrelated variables known as principal components. Each principal component represents a linear combination of the original variables, weighted by their respective loadings. The principal components are sorted in descending order of the amount of variance they explain, allowing us to focus on the most influential components. We determined the optimal number of dimensions for PCA by calculating the proportion of variance explained by each component and plotting the cumulative sum of explained variance. By analyzing the plot, we identified the number of components that capture 95% of the total variance. The selected number of components for every dataset in presented in next table:

|  |  |
| --- | --- |
| **Dataset** | **Number of components** |
| Dataset 1 | 109 |
| Dataset 2 | 113 |
| Dataset 3 | 111 |
| Dataset 4 | 112 |
| Dataset 5 | 110 |

**TSNE** is a non-linear dimensionality reduction technique that aims to preserve local similarities in the data. In our analysis, we used two components to construct the t-SNE reduced data.

**UMAP** is a non-linear dimensionality reduction technique that preserves both global and local structures in the data. In this study, we selected three components for the dimensional reduction process using UMAP. By reducing the dimensionality to three, we aimed to capture the essential characteristics of the data while minimizing information loss.

* + 1. Clustering

In order to cluster the dimensionality reduced data into the optimal number of cell "states" (clusters), we employed Gaussian Mixture Modeling (GMM). GMM is a probabilistic model that assigns each cell to a mixture component, representing a potential cell state. To determine the optimal GMM model, we utilized the BIC criterion. The BIC criterion takes into account both the goodness of fit and the complexity of the model, allowing us to identify the optimal number of components and the appropriate covariance matrix structure. Each cell was assigned a posterior distribution to each state, indicating the probability of belonging to a particular cluster. The number of extracted states was determined to be optimal based on the BIC criterion, ensuring an effective clustering solution. The optimal GMM model for every dataset and for every dimensionality reduction method is presented in next table:

|  |  |  |
| --- | --- | --- |
| **Dataset** | **Optimal GMM model** | |
| Dataset 1 | PCA | Optimal number of components: 4  Optimal covariance matrix type: full |
| t-SNE | Optimal number of components: 3  Optimal covariance matrix type: diag |
| UMAP | Optimal number of components: 6  Optimal covariance matrix type: tied |
| Dataset 2 | PCA | Optimal number of components: 4  Optimal covariance matrix type: full |
| t-SNE | Optimal number of components: 3  Optimal covariance matrix type: tied |
| UMAP | Optimal number of components: 7  Optimal covariance matrix type: tied |
| Dataset 3 | PCA | Optimal number of components: 3  Optimal covariance matrix type: full |
| t-SNE | Optimal number of components: 5  Optimal covariance matrix type: diag |
| UMAP | Optimal number of components: 9  Optimal covariance matrix type: spherical |
| Dataset 4 | PCA | Optimal number of components: 5  Optimal covariance matrix type: full |
| t-SNE | Optimal number of components: 4  Optimal covariance matrix type: spherical |
| UMAP | Optimal number of components: 9  Optimal covariance matrix type: spherical |
| Dataset 5 | PCA | Optimal number of components: 4  Optimal covariance matrix type: full |
| t-SNE | Optimal number of components: 5  Optimal covariance matrix type: spherical |
| UMAP | Optimal number of components: 6  Optimal covariance matrix type: full |

* + 1. Visualization

In the next figures, the clusters inferred for every dataset and for every dimensionality reduction method are presented. In every case, a 2D plot was designed using the first two and most important dimensions of the dataset.

|  |  |
| --- | --- |
| **Dataset** 1 | |
| **PCA** |  |
| **TSNE** |  |
| **UMAP** |  |

|  |  |
| --- | --- |
| **Dataset** 2 | |
| **PCA** |  |
| **TSNE** |  |
| **UMAP** |  |

|  |  |
| --- | --- |
| **Dataset** 3 | |
| **PCA** |  |
| **TSNE** |  |
| **UMAP** |  |

|  |  |
| --- | --- |
| **Dataset** 4 | |
| **PCA** |  |
| **TSNE** |  |
| **UMAP** |  |

|  |  |
| --- | --- |
| **Dataset** 5 | |
| **PCA** |  |
| **TSNE** |  |
| **UMAP** |  |

In the next figures, we present the cell posteriors for every dataset and for every dimensionality reduction method. Cell posteriors represent the probabilities or likelihoods of a cell being in a specific state or category.

|  |  |
| --- | --- |
| **Dataset** 1 | |
| **PCA** |  |
| **TSNE** |  |
| **UMAP** |  |
| **Dataset** 2 | |
| **PCA** |  |
| **TSNE** |  |
| **UMAP** |  |
| **Dataset** 3 | |
| **PCA** |  |
| **TSNE** |  |
| **UMAP** |  |
| **Dataset** 4 | |
| **PCA** |  |
| **TSNE** |  |
| **UMAP** |  |
| **Dataset** 5 | |
| **PCA** |  |
| **TSNE** |  |
| **UMAP** |  |

**Appendix**

The analysis presented in this technical report was performed using Python programming language. The code was executed in Jupyter notebooks, which allows for easy documentation of the code and results.

In order to successfully run the notebooks, the following Python libraries are required: pandas, numpy, matplotlib, seaborn, scikit-learn, and tqdm. These can be installed via pip or conda.

To install the required libraries via pip, simply run the following command in the command line: `pip install pandas numpy matplotlib seaborn scikit-learn tqdm umap-learn `

To install the required libraries via conda, simply run the following command in the command line: `conda install pandas numpy matplotlib seaborn scikit-learn tqdm umap-learn`

It is recommended to use Python version 3.7 or higher for running the notebooks.