Identifying Structure in the MIMIC ICU Dataset

Zad Chin Harvard University Cambridge, MA, 02138 zadchin@college.harvard.edu

Leo Anthony Celi Massachusetts Institute of Technology Cambridge, MA, 02138 lceli@mit.edu Shivam Raval Harvard University Cambrdige, MA, 02138 sraval@g.harvard.edu

Martin Wattenberg Harvard University Cambridge, MA, 02138 wattenberg@seas.harvard.edu

Finale Doshi-Velez Harvard University Cambridge, MA, 02138 finale@seas.harvard.edu

Abstract

The MIMIC-III dataset, containing trajectories of 40,000 ICU patients, is one of the most popular datasets in machine learning for health space. However, there has been very little systematic exploration to understand what is the *natural* structure of the MIMIC-III dataset–most analyses enforce some type of top-down clustering or embeddings. We take a bottom-up approach, identifying consistent structures that are robust across a range of embedding choices. We identified two dominant structures : (1) structure with two clusters clearly distinguished by fraction-inspired oxygen (2) structure with two clusters separated by creatinine. Both of the distinguishing factors were validated as the key features by our clinician co-author. Our bottom-up approach in studying the macro-structure of a dataset can also be adapted for other datasets to provide undiscovered insights.

1 Introduction

Understanding the structure of clinical data can help us identify important insights into disease subtypes and process variation. Within medicine, clustering techniques have been commonly applied to the diagnosis of breast cancer (Chen [2014]), Parkinson's disease (Polat [2012];Nilashi et al. [2016]), mental health and psychiatric disorders (Trevithick et al. [2015]), heart and diabetes diseases (Yilmaz et al. [2014]), among many others.

In this paper, we focus on understanding the natural structure of the MIMIC-III Intensive Care Units (ICU) electronic health record data set (Johnson et al. [2016]) via projection-based clustering methods. MIMIC-III contains timeseries of measures of 40,000 patients in the ICU. With 4,564 citations as of October 2022, it is one of the most commonly used data sets in machine learning and health. Thus, its structure is of inherent interest. However, to date, most works have imposed a top-down approach to finding structure—e.g. imposing a particular clustering method (Shea [2020]; Sharafoddini et al. [2021];Zhang et al. [2021]; Fang et al. [2020]) or creating a task-specific embedding (Beaulieu-Jones et al. [2018]; Galozy [2018]; Chaudhary et al. [2020]).

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Most of the literature identified did not take measurement decisions into account. MIMIC-III data consists of irregularly measured time series and the care of critically ill patients relies heavily on laboratory data, and by extension, the laboratory reference ranges associated with them (Tyler et al. [2018]). However, these laboratory reference ranges are typically created by surveying healthy outpatients (Rifai [2017]). Hence, although using standard scaling in machine learning is computationally convenient, it leads to a significant loss of information (potentially introducing bias and making certain clinical insights inaccessible) and also significant inconsistency in the results.

Moreover, for clinical questions which are already frequently asked and well-defined (e.g. predicting the course of a specific disease or patient outcome), need-driven clustering methods, such as those explored in previous studies, have shown promise for generating insights. However, there are also potentially important clinical questions "hidden" in the data which have never been asked (e.g. the existence of coherent, discrete subtypes of disease with independent progression mechanisms), and in that sense may be more novel. Furthermore, when we first cluster the data and then distinguish the characteristic of the clusters with the clinical labels of interest such as mortality, we are ignoring other possible informative factors that are distinguishing the clusters, which are important for the interpreting the resulting clustering.

Hence, in our study, addressing previously identified limitations, we explore the macro-structure of the MIMIC-III dataset from a data-driven perspective, in which objects are similar within clusters and dissimilar between clusters restricted to Euclidean dissimilarity. Our approach is to use projections as conventional methods of dimensionality reduction for information visualization in order to transform high-dimensional data into two-dimensional space (Venna et al. [2010]). We adopted various dimensionality reduction techniques, including Principal Component Analysis (Abdi and Williams [2010]), t-Stochastic Neighbor Embedding (Schubert and Gertz [2017]), and Uniform manifold approximation and projection (McInnes et al. [2018]) in projecting our data into a two-dimensional space. Borrowing ideas from persistent homology, we further investigated how persistent our projections are against different methods of standardizations with different clinical ranges and against different model hyperparameters. Once we identified persistent projections, we identified clusterings based on Euclidean metrics. Once clusters are identified, we further investigate the discriminant of the clustering by statistical methods and later validate the discriminant found with statistical methods with a rule-based decision tree. Our approach provides 3 major benefits: (1) It summarizes the dataset in a tractable way, without imposing any labelling, and gives a "representative sample" of patients for physicians. (2) A more informative and comprehensive interpretation of the clusterings rather than coercing a supervised label (such as mortality) to the clustering (3) A pioneering study in how different clinical range affects the structure of the data relating to critically ill patient.

Within the range of our exploration, we find that the dominant structures found either by fractioninspired oxygen (Fi02) or by creatinine levels—both of which were validated as key patient features by our clinical co-author, Dr Leo Anthony Celi, and have been mentioned in another medical journal to be important in clinical decisions (Kang et al. [2017]; Barbateskovic et al. [2019]). Specifically, creatinine is an important indicator for physicians during the earlier stage of admission to the ICU while Fi02 is an important indicator for the later stage of diagnosis and treatments. Moreover, we found a much greater sensitivity to standardization approaches than projection parameters. Our work provides important insights about the MIMIC-III data set and suggests ways to identify robust structures in other clinical data sets.

2 Methods

Our MIMIC-III cohort is the first 48 hours of seven commonly measured features of each of 10,184 patients: The features are heart rate, respiratory rate, creatinine, hematocrit, fraction inspired oxygen, sodium and mean blood pressure. The detail in cohort selection can be found in section A.2 in Appendix. When seeking structure, we explored the effect of (a) different ways to standardize the data and (b) different projection parameters. We identified commonalities between the induced embeddings for further inspection.

2.1 Description of Different Standardization Methods

Different features have different natural ranges and variations. For example, the heart rate normally varies between 60 and 100 beats per minute (BPM); the respiratory rate varies between 12 and 18

Vitals	Valid Low	Valid High	Normal Low	Normal High	Ourlier Low	Outlier High
Heart Rate	0	350	60	100	0.0	390
Respiratory Rate	0	300	12	18	0.0	330
Creatinine	0.10	60	0.59	1.35	0.0	66
Hematocrit	0	75	37	52	0.0	100
Fraction Inspired Oxygen	0.21	1.00	0.21	0.5	0.2	1.1
Mean Blood Pressure	14	330	70	100	0.0	375
Sodium	50	225	135	145	0.0	250

Table 1: Table 1 shows the Valid, Normal and Outlier clinical laboratory values for different features: Valid clinical laboratory values are values that feasibly exist in human physiology. Normal clinical laboratory values are values that are normal for a healthy individual. Outliers clinical laboratory values are the cutoff values in which humans can never achieve laboratory values beyond the stated outliers values. All values are obtained from our clinical co-authors and have been validated.

breaths per minute (BPM). Thus, decisions have to be made regarding how to standardize the data. For each feature d, we consider standardizations of the form:

$$z_d = f((x_d - x_{d,\min}) / (x_{d,\max} - x_{d,\min})),$$
(1)

For the values of $x_{d,\min}$ and $x_{d,\max}$, we randomly sample values between *valid* values—a range that feasibly exists in the human physiology—and *normal* values—a range that that is normal for healthy outpatients, but not necessarily normal or safe for ICU patients. These values are in Table 1. We consider two forms for f: *continuous*, that is, f(x) = x, or *discrete*, that is f(x) = -1 if x < 0; f(x) = 0 if $x \in [0, 1]$, and f(x) = 1 if x > 0. Finally, we include the "standard" standardization $z_d = (x_d - \bar{x}_d)/\sigma_{x,d}$ based on the feature's mean and variance.

2.2 Description of Different Projection Parameters

We consider two types of projections, t-Stochastic Neighbor Embedding, t-SNE, (Van der Maaten and Hinton [2008]) and Uniform manifold approximation and projection, UMAP, (McInnes et al. [2018]). Previous works on projections using nonlinear and stochastic methods show that they are sensitive to the hyperparameters of the techniques (Wattenberg et al. [2016]). For t-SNE, we vary the perplexity, that parameter that impacts the resulting projections the most. We selected a range of values, with perplexity = $\{5, 30, 50, 80\}$. For UMAP, the different parameters we explored were min_dist and n_neighbors, where we tested n_neighbors = $\{5, 15, 50, 100\}$ and min_dist = $\{0.1, 0.25, 0.5, 0.99\}$. (Note: we also explored using PCA, but the linear projection did not reveal substructure in the data—see Appendix Figure 3.)

2.3 Embedding Similarity Metric

Each standardization and choice of parameters in projection resulted in an embedding of the data set in a two-dimensional space. To compare the similarities between the embeddings, we tested a total of 42 different settings: 5 discrete randomized ranges and 100 continuous randomized range for both t-SNE and UMAP, where the randomized clinical range is scaled with $e \sim \text{Unif}(0,1)$ from valid range and normal range identified in 1, with the formula below:

$$x_{\min-random} = x_{\min-valid} + e(x_{\min-normal} - x_{\min-valid})$$
(2)

$$x_{\text{max-random}} = x_{\text{max-normal}} + e(x_{\text{max-valid}} - x_{\text{max-normal}}), \tag{3}$$

, UMAP n_neighbors = 5, 15, 50, 100, UMAP min_dist = 0.1, 0.25, 0.5, 0.99 and t-SNE perplexity = 5, 30, 50, 80 for both continuous and discrete range, resulted in a total of $10 + 10 + 3 \times (16) = 42$ different methods tested. For each of the methods, we compare the labelling that we get for each method of standardization and metrics, and compute a similarity matrix between them (put simply: How many points stay at the same clusters comparing two different methods). We visualize the similarity matrix with a heat map for clarity.

3 Results

Methods	Continuous Valid	Continuous Normal	Discrete Valid	Discrete Normal	Standard Scaling
UMAP					
T-SNE					

Table 2: Table 2 shows the UMAP and T-SNE under different standardization schemes identified in Section 2.1. *UMAP*: The Continuous-Valid and Discrete-Normal schemes have separable clusters (colored in red and blue). The Continuous-Normal and Continuous-Standard Scaled schemes have interesting substructures but no clear clustering. The choice of feature range for standardization has important effects on the structure of the resulting projections. *TSNE*: TSNE Projections for different standardization schemes. Similar to UMAP Projections, the Continuous-Valid and Discrete-Normal Schemes have separable clusters (colored in red and blue). Discrete Valid in particular shows interesting substructures. Similar to what we found in UMAP: The choice of feature range for standardization has been found to have a significant impact on the resulting projections.

Across the different standardization and projection parameters, we find two major structures.

Figure 1 shows the similarities between the embeddings across the different standardization and projection approaches for 100 randomized ranges respectively. We see that embeddings cluster into two main groups, distinguished by

- Fraction Inspired Oxygen, for the data that are continuously standardized with valid range or randomized continuous range
- Creatinine, for the data that is only discretely standardized against the normal range

Examples of projections from each group are given in figures in Table 2, as well as the associated timeseries. For embedding that shows interesting structures and visually identifiable clusters, we aim to uncover the important features that result in the particular structure. Statistically, we observed that for the Continuous-Normal scheme, we observe that cluster 1 (the small blue cluster identified in Table 2 column 2 of row 1 and row 2), has a mean and median Fi02 = 0.21 across 24 hours while the other clusters have FiO2 values maintained at a mean of 0.8 and a median of 0.6 for the first 5 hours at maintained at a mean and median of 0.5 after the 5th hour, as shown in the Figure 4a. For the Discrete-Normal scheme, we observe that cluster 1 (the small blue cluster identified in Table 2 column 5 of row 1 and row 2) a mean and median creatinine ≈ 0.8 mg/dL across 48 hours while cluster 2 has creatinine > 0.8mg/dL across 48 hours, as shown in the Figure 4b.

Separation Based on Features We further validated our result using the Simple Decision Tree Classifier to investigate whether there are simple human-understandable rules that can be inferred from this structure that may reliably predict whether a data point may lie in a particular cluster. We observe that a single-layer decision tree is sufficient to find a distinguishing feature between the cluster that cleanly separates more than 97% of the points inside a cluster.

The decision tree result can be found in Figure 2. For both cases where we observe identifiable clusters, we find a decisive feature that separates the clusters and is consistent with the statistical analysis. For the Continuous-Valid scheme, the most important feature for cluster separation is Fraction-Inspired Oxygen at hour 10, the decisive value being 0.24 and for the Discrete-Normal scheme, it is Creatinine at hour 20, with the decisive value being 1.35 mg/dL. For the Continuous-



Figure 1: Figure 1 shows the heatmap Analysis for 100 different randomized ranges. 5 Observations has been identified: (*Observation 1:*) Different randomized continuous range shows consistency throughout different models and UMAP and t-SNE parameters. (*Observation 2:*)The other inconsistency in the heatmap indicates that discrete methods of standardization are sensitive to the scale of the clinical ranges. (*Observation 3, 4:*) consistency of clustering is observed with different UMAP and t-SNE parameters for continuous range, with the exception of t-SNE perplexity = 5 and 80. (*Observation 5:*) The bottom right dark blue clusters represent that for discrete ranges have consistent labelling across different UMAP and t-SNE parameters, except for t-SNE perplexity - 5, 30 and 80.

Valid scheme, it is clear that patients with lower values of FiO2 are in one small cluster, with the other cluster having larger values.



Figure 2: Figure 2 shows the output of the decision tree. The identified first level features important for separating clusters, which is fraction inspired oxygen and creatinine, are clinically relevant in the early stages of treatment.

Effect of Standardization Methods on Distribution We further investigated why different standardization methods give rise to different clustering and structure by investigating the distribution plot for the identified feature. From Figure 5, we observed that the standardization methods that produce meaningful results are those capture the distribution of the original data closely.

Clinical Relevance of Features In the critical care setting, Fraction Inspire Oxygen (Fi02) describes the population of patients that are on/off respirators, with patients on respirators having higher FiO2. Creatinine, an important feature about kidney function, has been identified by the medical domain expert to be crucial for physicians' clinical decisions in the early stages of patients' treatment outcomes.

The "Typical" Ways To Standardize Lead to Very Different Projections. Critically ill patients have numerous laboratory abnormalities. The laboratory reference intervals that define normal values established by sampling healthy outpatients might not provide insights. While we checked for robust

patterns across a different range of choices as well as across different models, we found out that the typical ways of standardizing, which is based on the feature's mean-variance, as well as standardizing against normal ranges, resulted in the loss of structure in the projection, as shown in figures identified in Table 2 columns 3, 4, and 6. Hence, future research should be more careful in selecting a clinical range to standardize against.

4 Conclusion

In this work, we identified that across seven different combinations of ways to standardize and project the MIMIC-III data, two key structures emerged, one dependent on fraction-inspired oxygen and one dependent on creatinine. While we explored the effect of standardizations in depth, there are additional directions one could explore to identify if other key structures exist. In particular, we always assumed that timesteps aligned in the 48-hour timeseries. Future work could more closely relate outcomes to the topologies we discovered, further explore the substructures in our projections, and explore ways beyond standardization to compare different patient timeseries.

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A Appendix

A.1 Additional Figures



Figure 3: Figure 3 shows the PCA Projections for Continuous Valid Scheme. PCA method resulted in qualitatively similar plots for other standardization schemes. In high-dimensional data like the MIMIC-III dataset, MIMIC-III fails because since all linear projections are orthonormal rotations of the data coordinates, clusters that are linear nonseparable entanglements, such as the Chainlink data (Ultsch [1995]) or overlapping convex hulls, cannot be separated.



(b) Creatinine Timeseries

Figure 4: Figure 4 shows timeseries for different clusters for UMAP. For Continuous-Valid, the clusters are clearly separated by Fi02, with cluster 1 having mean and median of 0.21 across the 48 hours while cluster 2 has FiO2 mean and median approximately of 0.5 after the first 10 hours. For Discrete-Normal, clusters are separated by Creatinine, with Cluster 1 having a mean and median below 1.0 while cluster 2 having a mean below 2.5 and above 2.0 and a median of 3.0 across 48 hours.



Figure 5: Figure 5 shows the effect of Standardization on the Distribution of important features. For Fraction Inspired Oxygen at Hour 10, we observe that the continuous valid distribution closely captures the peaks observed in the original feature but in a more normalized manner. For Creatinine we observe that the original feature has a slightly skewed normal curve, which is similarly observed in the discrete-normal case. Our hypothesis is standardization methods that closely capture the distribution of the original data produce meaningful results.

A.2 Data and Cohort Selection

In this study, we used data from the Medical Information Mart for Intensive Care (MIMIC-III) database, which is a large, freely available database comprising de-identified health-related data associated with over forty thousand patients who stayed in critical care units at Beth Israel Deaconess Medical Center in Boston, Massachusetts between 2001 and 2012. (Johnson et al. [2016]). For our study, we sample from the MIMIC-III dataset based on the following criteria:

- Patients with minimum 48 hours of records;
- Patients with age over 18 years old;
- Patients with complete age, gender, ethnicity group, insurance, readmission, and mortality record

A total of 10,184 patients fit the criteria and are considered. For each of the patients in the selected cohort, we have further selected seven laboratory test measurements for their first 48 hours in the ICU. These 7 clinical features are a key indicators of important organ functions and have been commonly mentioned in previous research in the ICU. These clinical features are respiratory rate, heart rate, mean blood pressure, sodium, creatinine, fraction inspired oxygen, and hematocrit (Fialho et al. [2012]Curto et al. [2016]Desautels et al. [2017]).

For each feature, we record its value at each hour of the patients' first 48 hours of admission to the ICU. If a value is not found, we carry forward the value from the hour before. To minimize model variance in our study, we also remove the patients with outliers values, which are values that are physically impossible to achieve by human standards, as shown in Table 1. All in all, we get a data frame that has a dimension of $10, 184 \times 336$.

A.3 A view on Projection Based Dimensionality Reduction Algorithm

For high-dimensional datasets, finding global structure directly is difficult as in high-dimensional, the curse of dimensionality applies, where there is a loss of meaningful differentiation between similar and dissimilar objects observed. Dimension reduction techniques reduce the dimension of the input space to facilitate finding structures in the data. A novel method for improving structure finding in the high-dimensional dataset is using projection-based clustering. A projection is used as a method for visualizing high-dimensional data in 2D space such that the distance and the density of the data are captured. One approach is to use projections as conventional methods of dimensionality reduction for information visualization in order to transform high-dimensional data into low-dimensional space (Venna et al. [2010]). Thus, in our paper, we experimented with 3 unsupervised dimension reduction techniques, namely Principal Component Analysis (PCA), t-distributed stochastic neighbor embedding (t-SNE), and Uniform Manifold Approximation and Projection (UMAP). We projected our data to two-dimensional, which resulted in a scatter plot that we can analyze. Scatter plots generated by a projection method remain the state-of-the-art approach in cluster analysis to visualize data structures. (e.g., Mirkin [2005]; Ritter [2014]; Hennig et al. [2015]). We will discuss each of the dimension reduction techniques mentioned above below.

PCA is the oldest and most commonly used linear projection technique that reduces the dimensions of the data. PCA works by dropping the least important variables and preserving the most valuable variables, projecting them onto a two-dimensional space. However, for very high dimensional data, PCA may fail to find interesting structures and patterns. This is because since all linear projections are orthonormal rotations of the data coordinates, clusters that are linear nonseparable entanglements, such as the Chainlink data (Ultsch [1995]) or overlapping convex hulls, cannot be separated. With this type of projection, it is unavoidable that at some locations remote data are erroneously superimposed in the output space.

In such cases, non-linear stochastic methods like t-SNE and UMAP are particularly beneficial. t-SNE uses points in high-dimensional space to find a faithful representation of those points in a lower-dimensional space. The algorithm is non-linear and adapts to the underlying data, performing different transformations on different regions. The other non-linear method is UMAP, created by McInnes and Healy (McInnes et al. [2018]), uses the data to construct the initial high-dimensional weighted graph with the nearest neighboring points in a local region. Compared to t-SNE it preserves as much of the local and more of the global data structure, with a shorter runtime.