Temporal Belief Memory: Imputing Missing Data during RNN Training

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Abstract

We propose a bio-inspired approach named Temporal Belief Memory (TBM) for handling missing data with recurrent neural networks (RNNs). When modeling irregularly observed temporal sequences, conventional RNNs generally ignore the real-time intervals between consecutive observations. TBM is a missing value imputation method that considers the time continuity and captures latent missing patterns based on irregular real time intervals of the inputs. We evaluate our TBM approach with realworld electronic health records (EHRs) consisting of 52,919 visits and 4,224,567 events on a task of early prediction of septic shock. We compare TBM against multiple baselines including both domain experts' rules and the state-of-the-art missing data handling approach using both RNN and long shortterm memory. The experimental results show that TBM outperforms all the competitive baseline approaches for the septic shock early prediction task.

1 Introduction

Multivariate time series data are ubiquitous in real-world dynamic systems such as health care and distributed sensor networks. In many of these systems, measurements are commonly acquired at irregular intervals. For example, many health care systems record large amounts of time series data in electronic health records (EHRs) for each patient's visit; during a patient's visit, the body temperatures are often measured a few times a day while the white blood cells are only measured every other day. As a result of merging such irregular data, real-world multivariate time series data is often plagued by missing values.

Generally speaking, the mechanisms of missing data can be divided into three categories: missing completely at random (MCAR), missing at random (MAR), and missing not at random (MNAR) [Rubin and Little, 1987]. In the domain of EHRs, for example, MCAR refers to missingness that is independent of all observed and unobserved values; one such example is when equipment failed to collect a patient's data. MAR refers to missingness that is independent of unobserved values but depends on the observed values: for example, patients with very good vital signs may not need to undergo certain lab tests. Finally, MNAR refers to missingness that depends on both observed values and unmeasured variables: for instance, a depressed patient might refuse a depression screening.

Ideally, different approaches should be applied depending on the missing mechanisms. For example, if the data is MCAR and the missing rate is low, we can delete the cases with missing values; if the missing rate is high, we might impute missing values with the mean. However, as many realworld datasets often have all three categories of missingness, various missing data approaches have been used, and some common approaches include forward-filling, hot-deck, EM imputation [García-Laencina *et al.*, 2015], resampling [Cismondi *et al.*, 2013], multiple imputation [Galimard *et al.*, 2016] and so on. More recently, Lipton *et al.* showed using *missing indicators* (MI) [Rubin and Little, 1987] to be highly effective for handling temporal missing data [Lipton *et al.*, 2016].

In this paper, we propose a bio-inspired imputation method, temporal belief memory (TBM), that considers the time continuity and captures latent missing patterns based on irregular real time intervals of inputs to handle missing data. More specifically, TBM computes a belief of the last observation over time for each feature and imputes a missing value based on that individual belief in both forward and backward directions. We evaluate the proposed TBM method and compare it with four competitive missing data handling methods - mean-substitution, forward-filling, expert knowledge based imputation rules - and the recent effective approach, missing indicator (MI) [Lipton et al., 2016]. These missing data handling methods are compared on two types of state-of-theart classifiers: RNNs and long short-term memory (LSTM) [Hochreiter and Schmidhuber, 1997]. Our empirical results showed TBM can effectively handle multivariate time series data with a high rate of missing values. It outperformed not only the two baseline methods (mean-substitution and forward-filling) but also the domain experts' rules and MI. TBM and MI tackle missing data from two different perspectives, since the best performance is obtained when we combine the two approaches.

Recurrent neural networks (RNNs), such as LSTM and gated recurrent unit (GRU) [Cho *et al.*, 2014], have been shown to achieve the state-of-the-art results in many real-word applications with multivariate time series data through

deep hierarchical feature construction. Moreover, they can capture long-range dependencies in time series data in an effective manner. RNNs for missing data have been studied in earlier works [Bengio and Gingras, 1995], and applied for speech recognition and blood-glucose prediction [Parveen and Green, 2004; Tresp and Briegel, 1997]. Recently researchers tried to handle missingness in RNNs by concatenating missing entries, or incorporating a time based decay function, or synchronizing different sampling frequencies [Lipton *et al.*, 2016; Che *et al.*, 2016; Neil *et al.*, 2016]. To our knowledge, however, no prior work has addressed the power of TBM, which can systematically model missing patterns in both *forward* and *backward* directions, with RNNs for sequence-labeling tasks.

Here, our task is early prediction of septic shock. Sepsis is a life-threatening organ dysfunction caused by a deregulated host response to infection [Singer *et al.*, 2016]. As a leading cause of hospital death in the United States, sepsis affects nearly 30 million episodes and 6 million deaths per year worldwide [Reinhart *et al.*, 2017]. From 2005 to 2014 *Septic shock*, the most severe complication of sepsis, incidence increased from 6.7 to 19.3 per 1,000 hospitalizations, while mortality decreased from 48.3% to 39.3% [Kadri *et al.*, 2017]. Prior studies have indicated that early diagnosis and treatment of septic shock can prevent about 80% of sepsis deaths. On the contrary, over the first 6 hours after the onset of recurrent or persistent hypotension, every hour delay in antibiotic treatment leads to a 7.6% decrease in survival of septic patients [Kumar *et al.*, 2006].

One major challenge associated with early prediction of sepsis/septic shock is its subtle but fast progression at early stages with lack of information. Sepsis has a wide range of potential symptoms, and its common indicators such as infection, fast heart rate, high/low body temperature, and low blood pressure [Polat *et al.*, 2017] are highly likely to progress to other disease. Because of such delicate progressions, variables in the before-shock stage may either be measured infrequently or not measured at all. In result, the duration between two clinical events in EHRs can be long and the missing rate can be very high. For example, the EHRs used in this work were taken from large typical US hospitals, and on average more than 80% of data are missing and several variables' missing rates are above 99.9%. Thus, missing data handling can be a key factor in this early prediction task.

2 Related Work

2.1 The Hodgkin-Huxley Model

Our TBM approach was originally inspired by a biological neural model proposed by [Hodgkin and Huxley, 1952], commonly referred to as the Hodgkin-Huxley model. The Hodgkin-Huxley model describes the electro-chemical information transmission of natural neurons with electrical circuits. It has been shown to realistically model biological neurons, and consistently inspired the advance of artificial neural networks, including deep learning.

Despite the great success of deep learning, the existing artificial neural networks still cannot match the human brain on many tasks. Therefore, some recent research on artificial neural networks returns to the biological roots of neurons and looks at how our brains function. Among them, spiking neural networks (SNNs) have gained increasing attention. SNNs can be seen as *time* dependent neural networks inspired by the Hodgkin-Huxley model and other bio-neuron models, and are often considered as the third generation of computational neural networks [Maass, 1997]. Compared with conventional artificial neural networks, SNNs take advantage of the precise timing of spikes generated by neurons and thus have greater computational power [Gerstner and Kistler, 2002]. Indeed, a probabilistic SNN outperformed deep networks for breast cancer prediction [Hsieh and Tang, 2013].

Although this work has a common ground with SNN in that the neural networks embrace a time concept to process inputs, we focus on the decaying mechanism on how bio-neurons handle missing signals, while most SNNs have concentrated on the spiking mechanism on how bio-neurons fire.

2.2 Missing Data Handling in RNNs

A recurrent neural network (RNN) is a type of deep neural network, designed to learn temporal patterns in sequential data. Although RNNs are theoretically able to find long-term dependencies underlying sequential data, classical RNNs often do not effectively capture them due to the vanishing and exploding gradient problem [Graves, 2013]. As variants of RNNs, LSTM [Hochreiter and Schmidhuber, 1997] and GRU [Cho *et al.*, 2014] overcome these issues by incorporating multiple gating units into an RNN structure. A gating mechanism allows for explicit memory delete and update, and controls the flow of information in hidden units.

As frontier work for missing data handling for RNNs, [Bengio and Gingras, 1995] proposed an RNN structure for both missing inputs and asynchronous data that randomly initializes missing values and optimizes the filled values through backpropagation. In [Tresp and Briegel, 1997], they demonstrated a modified RNN for missing data handling, combined with a linear error model trained by an expectationmaximization technique. The experimental results show that their method improves performance in the glucose/insulin metabolism prediction task with respect to both conventional RNNs and various linear models.

Recently, [Lipton *et al.*, 2016] showed the effectiveness of missing indicators (MI) with LSTM for the 128 phenotype prediction task using an EHR dataset. In their work, they gave an insight that LSTM is implicitly able to impute missing values based on its memory. About the same period, Phased LSTM [Neil *et al.*, 2016] extends an LSTM unit by adding a time gate to align asynchronous streams, which allows the feature learning only when the time gate is open. On the other hand, [Kim *et al.*, 2017] incorporated a forward-filling operation for missing data into RNN and LSTM, and tested it to a clinical variable prediction task. While all these proceeding approaches show their own achievements, TBM addresses the missing data problem in terms of an observation's reliable period based on the input timing.

Closely related to this work, GRU-D [Che *et al.*, 2016] imputes missing values using a modified GRU, regulated by a temporal decay function with trainable weights. On a wide range of tasks, the authors showed that GRU-D often demon-

strates performance comparable to MI. In their work, they introduce an input/ hidden state decay function, and impute missing values with the input decay and update the previous hidden state with the hidden state decay. Our TBM differs from GRU-D in terms of four aspects: 1) when an input is observed, GRU-D imputes missing values in a forward direction, while TBM bidirectionally updates the imputed values within a reliable time window; 2) GRU-D updates the hidden state with the trained decay rate, while TBM only uses errors backpropagated by RNN models to update its parameters and does not modify an RNN structure; 3) GRU-D's trainable parameters are weight matrices, while TBM's is time; finally, 4) MI [Lipton et al., 2016] is incorporated into GRU-D (the update/reset gates and the hidden state candidate), while in this work MI is not originally incorporated into TBM but they can be combined. In fact, we explored the effectiveness of MI only, TBM only and combining TBM and MI.

3 Methods

In this section, we describe how TBM is derived from the Hodgkin-Huxley model and how TBM's model parameters can be learned during RNN training.

3.1 Bio-inspired Missing Data Handling

The Hodgkin-Huxley model explains how to propagate electro-chemical signals through bio-neurons, using the voltage equation defined as:

$$V(t) = \begin{cases} I * R(1 - e^{-t/(R*C)}) & I > 0\\ v * e^{-t/(R*C)} & I = 0 \end{cases}$$
(1)

where t is the time interval between the current time and the last observed time, V is a voltage, v is the last voltage when the input ceases, I is an input current, R is resistance, and C is capacity. As shown in Equation (1), while input signals continuously come into the pre-synapse of neuron within a specific time interval (I > 0), the neuron accumulates the voltage into its membrane, and if the voltage goes beyond a threshold, the neuron fires and propagates the signal to other neurons. This process is explained by the accumulation function (Equation 1, Top). When the input signal stops to come into the pre-synapse (I = 0), the voltage gradually decreases over time, and if it goes below the threshold, the signal propagation stops, which is performed by the decay function (Equation 1, Bottom). It should be noted that the accumulating/decaying gradients of voltage depend on R * C. As R * C increases, the voltage changes less rapidly; as R * Cdecreases, the voltage changes more rapidly.

For R * C, there are two notable properties. First, when the signal ceases (I = 0), if the neuron has been already activated, the voltage is still valid at least within a specific time window. Second, since each neuron has its own resistance Rand capacity C, each neuron has its own accumulating and decaying rate for voltage. That is, each neuron has its own time-based memory mechanism: how fast it accepts or forgets input signals.

Unlike continuous and frequent signals processed by bioneurons, real-world multivariate time series data such as EHRs are characterized by discrete and sparse data. Therefore, to apply the Hodgkin-Huxley model to our data, we need to reformulate the voltage equation. The accumulation function (Equation 1, Top) models bio-neurons' continuous behaviors toward input and thus better suits for high frequency data such as audio or video. The decay function (Equation 1, Bottom) is designed to model when there is no input, how the voltage V gradually decreases over time. Similarly, for EHRs, we can treat each observation as the initial voltage and model how our confidence on its value gradually decreases over time. Next, we describe how to combine this decay function mechanism with RNNs to address the data missingness.

3.2 Temporal Belief Memory

Figure 1 shows the architecture of a temporal belief memory (TBM), whose output connects to the input of LSTM. A TBM is a memory module that consists of two gating units (a missing gate m and a belief gate b), which collaboratively enables imputation of missing values based on beliefs of observations over time. The missing gate m indicates whether a value is missing (set to 1) or present (set to 0), and the belief gate b decides whether the last observation carries over to the imputed value \tilde{x} based on the temporal reliability of the last observation. When an input is observed (m = 0), the observed value directly passes to the input of LSTM and updates the last observation, x_l , with the current value. When an input is missing (m = 1), the belief gate b computes the belief of x_l based on the time interval t between the current time and the last observed time. If the belief is greater than a threshold, it imputes the missing value with x_l ; otherwise, it sets the missing value to x_m , which is the mean value of observations for each feature.

Once TBM imputes the missing values, LSTM is trained utilizing inputs with imputed values. The LSTM units are described as

$$i_{t} = \sigma_{i}(x_{t}W_{xi} + h_{t-1}W_{hi} + b_{i})$$

$$f_{t} = \sigma_{f}(x_{t}W_{xf} + h_{t-1}W_{hf} + b_{f})$$

$$\tilde{c}_{t} = tanh(x_{t}W_{xc} + h_{t-1}W_{hc} + b_{c})$$

$$c_{t} = f_{t} \odot c_{t-1} + i_{t} \odot \tilde{c}$$

$$o_{t} = \sigma_{o}(x_{t}W_{xo} + h_{t-1}W_{ho} + b_{o})$$

$$h_{t} = o_{t} \odot tanh(c_{t})$$

$$(2)$$

where W is a weight matrix, b is a bias, σ is a sigmoid activation function, tanh is a hyperbolic tangent function, and \odot denotes an element-wise vector product. In these equations, i_t , f_t , and o_t indicate the input, forget, and output gate at time t, while x_t , \tilde{c}_t , c_t , and h_t denote the input, memory cell state candidate, memory cell state, and memory cell output at time t, respectively. Although an LSTM is given as example in this section, TBM is scalable to variants of RNNs such as simple recurrent networks and GRUs.

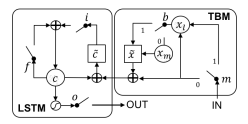


Figure 1: Temporal Belief Memory with LSTM.

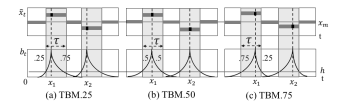


Figure 2: Belief propagation modes: (a) TBM.25: 25% of backward and 75% of forward belief interval, (b) TBM.50: equal weights of backward and forward belief interval, and (c) TBM.75: 75% of backward and 25% of forward belief interval.

Belief Gate

As described above, the decay function: $V(t) = v * e^{-t/(R*C)}$ (Equation 1, Bottom) can play a pivotal role to impute missing values. To reformulate it, we reinterpret a voltage V at time t as a temporal *belief* of an input value at current time t and then impute missing values based on the current belief.

First, we set the v in the decay function to be 1, which means whenever an input is observed, our belief on its value is 1; R * C is combined into one parameter τ , which is conceptually a reliable-time-window variable that indicates how long we can trust the last observation when the current value is not observed. t is mapped to δ_t , the time interval between the last observation and the current time t. Thus we have a rather simple belief function at time t referred to as b'_t :

$$b_t' = e^{-\delta_t/\tau} \tag{3}$$

which models that the belief or the confidence of the last value gradually decreases over time. On the other hand, observations in EHRs are often reliable for a certain period in *bidirectional* ways. For example, if the current heart rate is 100, it is likely to be 100 a few minutes prior as well as a few minutes later. To incorporate the bidirectional nature of observations, we introduce a new parameter β into the equation above and result a new definition b''_t :

$$b_t'' = e^{(-\beta * |\delta_t|)/\tau}$$
(4)

where β is a hyper-parameter that controls a shift of the time window τ and the absolute value of δ_t is the time interval between the last observed time and the current time. Note that backward belief propagation can be implemented by mirroring the forward belief and the use of the absolute value of δ_t would support both the backward and forward cases. The lower part of Figure 2 shows the impact of different β s on the belief functions. Here based on different combinations, we have three settings of TBM: (a) TBM.25: 25% of backward and 75% of forward belief interval, (b) TBM.50: equal weights of backward and forward belief interval, and (c) TBM.75: 25% of backward and 75% of forward belief interval. Finally, we apply the unit step function with threshold h on b_t'' to get the belief gate, $b_t = \theta_h (e^{(-\beta * |\delta_t|)/\tau})$ where if b_t'' in Equation 4 is greater than $h, b_t = 1$; otherwise $b_t = 0$.

Imputation

For a given time t, TBM imputes missing values using the missing gate m_t and the belief gate b_t . We denote an imputed

value at time t with \tilde{x}_t , which is defined as

$$\tilde{x}_t = (1 - m_t)x_t + m_t \{ b_t x_l + (1 - b_t) x_m \}$$
(5)

where x_t is a current value, x_l is a last value, and x_m is a mean value for a feature. When a current value is observed $(m_t = 0)$, x_t takes the current input x_t , while when a current value is missing $(m_t = 1)$, x_t takes the last value only if it is reliable (the output of belief gate is 1: $b_t = 1$). Otherwise, it takes the mean value for the corresponding feature.

Note that by using the unit step function to transform b''_t to b_t , the imputed values will not converge to the mean value within the reliable-time-window, and the variance of the imputed ones will be close to the original values. By contrast, the mean substitution or b''_t without the unit step function often abnormally creates average data points and decreases the variation of the imputed data. This decrease in individual variables is proportional to the number of missing data, and may considerably distort the correlations of variables when the missing rate is high [Cohen and Cohen, 1975].

4 Experiment

4.1 Data

Our dataset constitutes anonymized clinical multivariate time series data, extracted from the EHR system at Christiana Care Health System, Delaware from July, 2013 to December, 2015. Each visit/episode consists of multiple temporal events such as medical readings and interventions. In total, there are 119,857 patients, 210,289 visits, and 10,412,729 medical events. Along with time stamps, identifiers, locations, and description, there are three categories of main attributes as follows:

- Vital signs: systolic blood pressure, mean arterial pressure, temperature, heart rate, respiratory rate, etc.
- Lab results: white blood cell count, Bands, BUN, procalcitonin, platelet, creatinine, bilirubin, C-reactive protein, lactate, sedimentation rate, etc.
- Intervention: oxygen source, change of oxygen source, FiO2, drug administration, intravenous therapy, etc.

Target Population and Labeling

The study population are patients with *Suspected infection* which was identified by the presence of any type of antibiotic, antiviral, or antifungal administration, or a positive test result of Point of Care Rapid, and it consists of 52,919 visits and 4,224,567 medical events. Note that the study population, the aforementioned rules for identifying suspected infection, and the septic shock labeling in next paragraph were determined by two leading clinicians with extensive experience on this subject from Mayo Clinic and Cristina Care Health System.

Supervised models depend heavily on the accurate label of the training dataset. However, acquiring the true label (i.e., septic shock and non septic shock) can be challenging. Although diagnosis codes, such as International Classification of Diseases, Ninth Revision (ICD-9), are widely used for clinical labeling, solely relying on ICD-9 can be problematic as it has been proven to have limited reliability due to the fact that its coding practice is used mainly for administrative and

Feature	Missing rate	Feature	Missing rate	
Procalcitonin	0.9998	FiO2	0.8046	
CReactiveProtein	0.9994	MAP	0.7735	
SedRate	0.9992	DistolicBP	0.7214	
Bands	0.9892	SystolicBP	0.7204	
BiliRubin	0.9793	PulseOx	0.6369	
Lactate	0.9723	RespiratoryRate	0.6261	
WBC	0.9347	HeartRate	0.6064	
Platelet	0.9341	OxygenSource	0.1267	
BUN	0.9332			
Creatinine	0.9331	Infection	0.9438	
ChangeOxygenSrc.	0.9137	Inflammation	0.6964	
Temperature	0.8125	OrganFailure	0.7661	
Mean			0.8184	

Table 1: Missing rates of 23 features from Cristiana Health Care System EHR, of which every instance and every feature contains at least one missing value. The first 20 features are clinical readings, and the last three features are the early stages of sepsis.

billing purpose. Indeed, it has been widely argued that ICD-9 codes cannot be used for establishing reliable gold standards for various clinical conditions [Ho *et al.*, 2014]. More importantly, ICD-9 cannot tell when septic shock occurs at event level, which is essential for our task. On the basis of the Third International Consensus Definitions for Sepsis and Septic Shock [Singer *et al.*, 2016], our domain experts identified septic shock as having received vasopressor(s) or having had persistent hypotension (i.e., systolic blood pressure less than 90 mmHg or mean arterial pressure less than 65 mmHg for more than 1 hour) and enabled to diagnose septic shock at event level.

When applying both ICD-9 and our clinical rules, we identified 1,869 shock positive visits and 23,901 negative visits. Given the imbalanced ratio of positive and negative shock visits, we further conducted a stratified random sampling on shock negative visits while keeping the same underlying distribution of age, gender, ethnicity, length of stay and the number of records in both positive and negative visits. As a result, the final dataset has 3,738 visits (1,869 positives and 1,869 negatives) and 145,421 events.

Missing Data Analysis

Each visit consists of irregular multivariate time series events with missing values and missing attributes, because different attributes are measured at different events. For example, vital signs are measured every 8 hours while lab values are measured only every 24 hours. Hence there may not be available readings for lab results when a new event is created for vital signs. Table 1 shows the missing rates of 23 features in our final dataset. On average, the missing rate is 81.84%.

Experiment Setup

To evaluate the proposed TBM framework, we conducted a series of experiments to test its effectiveness for early septic shock prediction using two types of classifiers: RNN and LSTM. For each classifier, we explored three TBM imputation modes: TBM.25, TBM.50, and TBM.75, described in Figure 2, and compared these against two widely used baseline methods: *Mean* and *Forward*, and the rules suggested by domain experts, Expert.

- *Mean:* fills all the missing values with the mean value for the corresponding feature, which is zero in our case since the data is standardized.
- *Forward:* fills the missing values with the last observation until the next value is observed.
- *Expert:* is defined by the domain experts; it fills the missing values with the last value within the fixed length of forward time window (8 hours for vital signs and 24 hours for lab tests), and fills the remaining ones with the mean value for the corresponding feature.

Additionally, our TBM modes are also compared against and with missing indicator (MI) given its effectiveness [Lipton *et al.*, 2016]. When applying MI, we need to decide how to fill-in the missing values. In [Lipton *et al.*, 2016], they used MI with zero-filling, forward-filling, and hand-engineered-filling, respectively. In our comparison, we combined MI with *Mean*, *Forward*, and *Expert*, respectively. Also, we explored combining the three TBMs with MI to see whether combining them together would further improve our results.

To summarize, we used two classifiers: RNN and LSTM to compare the following twelve methods from four categories: the Base (*Mean*, *Forward*, *Expert*), the Base-MI (*Mean*+MI, *Forward*+MI, *Expert*+MI), the TBMs (TBM.25, TBM.50, TBM.75) and the TBMs combined with MI (TBM.25-MI, TBM.50-MI, TBM.75-MI). For both RNN and LSTM, we use one hidden layer with 30 hidden neurons and 32 maximum sequence length. We use the Adam optimizer [Kingma and Ba, 2015] with the batch size 30, and adopt early stopping with 7 patience after minimum 10 epochs.

Our evaluation metrics include accuracy, recall (sensitivity), precision (positive predictive value (PPV)), F1 score, and area under the ROC (receiver operator characteristic) curve (AUC). Accuracy, F1-score and AUC are widely used to measure the prediction performance for machine learning approaches. In the realm of medical science, researchers commonly refer to sensitivity (recall) and PPV for the annotation performance. Therefore, we include the metrics for both machine learning and medical science domains. In the learning process, we split data into 80% for training, 10% for validation, and 10% for test, and conduct 5-fold cross validation.

4.2 Results

Three-hour-before & overall shock prediction

Table 2 shows predictive performance results. The first column is the classifier, the second column is the missing data handling method, columns 3 to 7 present our evaluation metrics for the three-hour-before shock prediction, and the last column presents the AUC score for the overall shock prediction (i.e., 0-24 hour-before shock prediction).

For each classifier, Table 2 can be divided into four subsessions: Base, Base-MI, TBM, and TBM-MI. For each subsession, the best results are marked in bold. Also, for either classifier, the highest score per metric across all the models is underlined. Finally, the best model across all methods and two classifiers are labeled with *. In the following, we analyze predictive performance within each classifier and then compare across them.

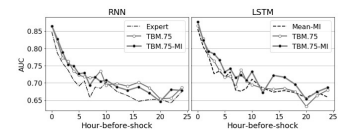


Figure 3: Septic shock prediction for 0-24 hour-before-shock, compared the best baseline, TBM.75, and TBM.75-MI

For RNN classifier, Table 2 shows that for the three Base methods, the *Expert* outperforms the other two on all measures except for the precision. For the Base-MI methods: there is no clear winner but in general *Mean*-MI and *Forward*-MI split the lead. For TBM-based model, TBM.75 outperforms the other two on all measures except for the precision; finally, for the three TBM-MI methods, TBM.50-MI outperforms the other two except for the precision and overall. Across the four categories using RNN, the best performance (underlined) is generated using the TBM-based models. In general, based on the best results from each category, we have Base \approx Base-MI < TBM-MI < TBM.

Next, for LSTM, Table 2 shows that for the three Base methods, the *Expert* outperforms the other two on all measures except for the precision and overall. For the Base-MI methods: *Forward*-MI outperforms the other two on all measures except for the recall and overall. For TBM-based model, TBM.75 outperforms the other two on all measures except for the precision; finally, for the three TBM-MI methods, TBM.75-MI outperforms the other two except for the recall. Across the four categories using LSTM, the best performance (underlined) is generated using the TBM-MI models. In general, based on the best results from each category, we have Base \approx Base-MI<TBM-MI for LSTM.

Across all 12 methods and two base classifiers, the best results come from LSTM+TBM-MI. On both classifiers, the TBM models outperform not only the popular applied baselines and the domain experts' rules but also the state-of-theart approach MI. However, we assume that TBM and MI tackle missing data handling from two different perspectives so that our best performance is generated by combining the two approaches using LSTM.

Detailed analyses on early prediction

We also evaluate TBM's early prediction capacity comparing to other baseline methods based on the 0-24 hour-before shock prediction task. In particular, we compare the best TBM (TBM.75) to the best *Base* (*Expert* for RNN and *Mean*-MI for LSTM) every hour between 0 to 10 hours and every two hours between 12 and 24 hours. Figure 3 (Left) shows that TBM.75 outperforms *Expert* throughout the time for RNN. Interestingly, when combining TBM with the previous state-of-the-art method, MI, TBM.75-MI does not outperform TBM.75. For LSTM (Figure 3, Right), TBM.75-MI outperforms *Mean*-MI except for the 12 hour-before shock prediction, while TBM.75 is comparable to *MEAN*-MI.

Base classifier	Method	3 hour-before-shock					Overall
		Acc	Recall	Prec	F1	AUC	AUC
RNN	Mean	0.7061	0.7076	0.7101	0.7077	0.7061	0.6875
	Forward	0.7233	0.7264	0.7378	0.7270	0.7233	0.6968
	Expert	0.7367	0.7529	0.7113	0.7292	0.7368	0.6983
	Mean-MI	0.7306	0.7427	0.7102	0.7256	0.7306	0.6954
	Forward-MI	0.7306	0.7644	0.6792	0.7152	0.7308	0.6956
	Expert-MI	0.7100	0.7416	0.6726	0.6997	0.7102	0.6973
	TBM.25	0.7567	0.7700	0.7368	0.7525	0.7568	0.7146
	TBM.50	0.7511	0.7554	0.7522	0.7221	0.7511	0.7083
	TBM.75	<u>0.7628</u>	<u>0.7815</u>	0.7357	<u>0.7570</u>	<u>0.7630</u>	0.7237
	TBM.25-MI	0.7467	0.7681	0.7168	0.7395	0.7469	0.7195
	TBM.50-MI	0.7556	0.7732	0.7312	0.7504	0.7557	0.7144
	TBM.75-MI	0.7528	0.7700	0.7313	0.7464	0.7530	0.7185
LSTM	Mean	0.7078	0.7088	0.7102	0.7094	0.7078	0.6903
	Forward	0.7156	0.7103	0.7423	0.7233	0.7154	0.6927
	Expert	0.7367	0.7429	0.7312	0.7358	0.7367	0.6918
	Mean-MI	0.7267	0.7346	0.7300	0.7276	0.7266	0.7107
	Forward-MI	0.7300	0.7263	0.7644	0.7395	0.7299	0.7067
	Expert-MI	0.7217	0.7174	0.7378	0.7249	0.7215	0.7058
	TBM.25	0.7556	0.7573	0.7589	0.7569	0.7556	0.7188
	TBM.50	0.7606	0.7725	0.7457	0.7576	0.7607	0.7184
	TBM.75	0.7633	0.7751	0.7456	0.7593	0.7635	0.7245
	TBM.25-MI	0.7739	<u>0.8085</u> *	0.7246	0.7623	0.7742	0.7262
	TBM.50-MI	0.7683	0.8002	0.7213	0.7572	0.7686	0.7322
	TBM.75-MI	<u>0.7839</u> *	0.7900	<u>0.7777</u> *	<u>0.7832</u> *	<u>0.7840</u> *	<u>0.7340</u> *

Table 2: Sepsis shock prediction at 3 hour-before-shock and the overall time (0-24 hours).

5 Conclusion

Missing data pervading in real-world multivariate time series datasets pose significant challenges in deriving robust predictive models for these real-world applications. More challengingly, the categories of missing data in these datasets are mingled or difficult to identify. To address this challenge, we have investigated missing data handling methods with RNNs and LSTMs, and examined early prediction of septic shock using imputed missing values in EHR. We have introduced a bidirectional time-based imputation method, called TBM, which was inspired by bio-neurons' behaviors. Empirical evaluations demonstrate that TBM achieves the best performance in the septic shock early prediction task, outperforming four competitive missing data handling methods.

Rather than use the pre-defined β s like in this study, in the future, we will further optimize TBM with respect to the model hyperparameter β , which adjusts the bidirectional portions of reliable-time-window. Another interesting line of research is to investigate the interpretation of the reliable-timewindow τ , which can suggest a desirable sampling frequency for each feature to clinical and medical experts. Finally, it will be important to evaluate robustness and generalizability of TBM by investigating the septic shock early prediction framework for other EHR datasets such as MIMIC-III and as well as other disease prediction tasks or different prediction tasks on multivariate time-series data suffering a high rate of missing values in other domains.

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