Deep Learning from Heterogeneous Sequences of Sparse Medical Data for Early Prediction of Sepsis

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Abstract: Sepsis is a life-threatening complication to infections, and early treatment is key for survival. Symptoms of sepsis are difficult to recognize, but prediction models using data from electronic health records (EHRs) can facilitate early detection and intervention. Recently, deep learning architectures have been proposed for the early prediction of sepsis. However, most efforts rely on high-resolution data from intensive care units (ICUs). Prediction of sepsis in the non-ICU setting, where hospitalization periods vary greatly in length and data is more sparse, is not as well studied. It is also not clear how to learn effectively from longitudinal EHR data, which can be represented as a sequence of time windows. In this article, we evaluate the use of an LSTM network for early prediction of sepsis according to Sepsis-3 criteria in a general hospital population. An empirical investigation using six different time window sizes is conducted. The best model uses a two-hour window and assumes data is missing not at random, clearly outperforming scoring systems commonly used in healthcare today. It is concluded that the size of the time window has a considerable impact on predictive performance when learning from heterogeneous sequences of sparse medical data for early prediction of sepsis.

1 INTRODUCTION

Sepsis is a leading cause of hospital morbidity and mortality (Singer et al., 2016). It is also one of the most serious forms of healthcare associated infections, many of which are considered avoidable (Cassini et al., 2016). The survival of sepsis patients is dependent on initiating appropriate antimicrobial treatment as early as possible (Ferrer et al., 2014). To facilitate this, prompt detection of sepsis patients is crucial. It has been shown that early identification of sepsis significantly improves patient outcomes. For instance, mortality from septic shock has been found to increase by 7.6% for every hour that antimicrobial treatment is delayed after the onset of hypotension (Kumar et al., 2006), while timely administration of a 3-hour bundle of sepsis care and fast administration of antibiotics was associated with lower in-hospital mortality (Seymour et al., 2017). Unfortunately, diagnosing sepsis early and accurately is challenging even for the most experienced clinicians, as there is no standard diagnostic test (Singer et al., 2016) and symptoms associated with sepsis may also be caused by many other clinical conditions (Jones et al., 2010).

Clinical decision support tools have the potential to facilitate early intervention in sepsis patients. Fortunately, clinical data that could be used to inform predictions about patients that are at risk of sepsis is already being routinely collected in electronic health records (EHRs). Today, warning scores based on such data are calculated for early identification of clinical deterioration at the bedside. Examples of early warning scores include NEWS (Williams et al., 2012) and qSOFA (Singer et al., 2016). These scoring tools compare a small number of physiological variables to normal ranges of values and generate a single composite score: once a certain threshold is reached, the system triggers an alert (Despins, 2017). However, a serious limitation of early warning scores is that they are typically broad in scope and were not specifically developed for sepsis, which means that many other diseases may also yield high scores. This can cause breakdowns in the training and education process and result in low specificity and high alarm fa-
tigue. Early warning scores, which are used in clinical practice today, are also overly simplistic by assigning independent scores to each variable, ignoring both the complex relationships between different variables and their evolution in time (Vincent et al., 1998; Smith et al., 2013).

Machine learning provides the possibility of overcoming the limitations of heuristics-based warning scores by accounting for dependencies between a large number of (temporal) input variables from EHRs to predict an outcome of interest, such as sepsis. Using machine learning for early prediction of sepsis has been greatly facilitated by the possibility of exploiting agreed upon definitions of clinical sepsis criteria for identifying cases (Delahanty et al., 2019). This allows for the large-scale creation of labeled datasets needed for supervised machine learning. Therefore, the primary focus of machine learning approaches should be on early detection, i.e. to detect sepsis as early as possible prior to sepsis onset.

In this article, the Sepsis-3 definition (Singer et al., 2016) is used for labeling three years of EHR data from a general university hospital population. In contrast, most previous efforts have focused on using data from intensive care units (ICUs) (Mani et al., 2014; Desautels et al., 2016; Taylor et al., 2016; Moor et al., 2019). Data from ICUs tends to be less sparse and hospitalization periods shorter. In this study, a recurrent neural network, in the form of a long short-term memory (LSTM) network, is used to learn from heterogeneous sequences of EHR data for early prediction of sepsis in the non-ICU setting. It is shown that the latter assumption yields somewhat better performance and indicates that missingness is sometimes a valuable indicator that should not always be imputed.

Predictive performance is evaluated for healthcare episodes of various lengths, revealing considerably higher performance for shorter vs. longer sequences. The model is also evaluated in terms of earliness, showing that more true positive predictions are made in the time windows closer to sepsis onset.

2 METHODS AND MATERIALS

The EHR data used in this study comes from the research infrastructure Health Bank (Dalianis et al., 2015), that contains EHR data collected from Karolinska University Hospital. This research has been approved by the Regional Ethical Review Board in Stockholm, Sweden under permission no. 2016/2309-32.

2.1 Data Selection

Patients older than 18 years admitted to the hospital between July 2010 and June 2013 were included, and followed until first sepsis onset, discharge or death. Patients were excluded if admitted to an obstetric ward and censored during ICU care. The dataset encompasses 124,054 patients and 198,638 care episodes from a general university hospital population over a three-year period. The incidence of sepsis in the dataset is 8.9%, yielding a very uneven class distribution.
that particular episode. Additionally, if the time following discharge and next admission for the same patient is less than 24 hours, the two are considered to be part of the same care episode. Care episodes may involve stays in several different wards and vary greatly in length, with a median length of around three days.

### 2.2 Feature Selection

Data for each care episode is collected solely from the EHR and comprises data on microbiological cultures and antimicrobial treatment, as well as demographic and physiological data. The included variables were selected by domain experts.

In the study, data defining collection of microbiological cultures and tests from all types of body fluids is used. Data on newly administreted antimicrobial treatment is collected based on ATC (Anatomical Therapeutic Chemical) codes (Nahler, 2009) J01 and J04. Demographic and physiological data is collected for the following 19 parameters: age, body temperature, heart rate, respiratory rate, systolic and diastolic blood pressure, oxygen saturation, supplementary oxygen flow, mental status, leucocyte count, neutrophil count, platelet count, C-reactive protein, lactate, creatinine, albumin, and bilirubin. Most of the variables are numeric, and generally extremely sparse, with a missing rate of more than 90% in some cases.

In addition to the aforementioned variables, early warning scores are also used as aggregated features in the predictive model. The output of the following scoring tools were used: NEWS2 (Williams et al., 2012), qSOFA (Singer et al., 2016) and SOFA (Vincent et al., 1996). This is a common practice in machine learning (Raghu et al., 2017a,b). The model was only allowed to access data that would be readily available in the EHR – or could be computed from it – at the time of prediction.

### 2.3 Care Episode Representation

To account for the temporality of the data, the care episodes are transformed into sequences based on a given window (bin) size. Experiments are conducted using six different window sizes: 1, 2, 3, 4, 6 and 8 hours.

A variable in a time window can either be missing or have multiple values associated with it. For example, body temperature may not have been measured in a given time window or may have been measured multiple times. When multiple values are present in a time window, the "worst" value is chosen. This is defined as the most pathological value for a particular variable and is determined apriori by clinical experts. For certain variables, the most pathological value can be either high or low. For example, in the case of body temperature, a value less than 36 is considered the worst, but if no such value exists, the highest value is chosen instead.

Missing data can be handled in various ways and a fundamental decision concerns whether data for a given variable is assumed to be missing at random or not (Steele et al., 2018). In this study, both assumptions are taken into account. When data is assumed to be missing not at random, imputation is not carried out; instead, missing values are simply assigned an integer value which is not present in the data, and the idea is that the model may learn to treat missingness as a distinct feature. When data is assumed to be missing at random, the following simple imputation strategy was carried out. If a value exists for a given feature in the care episode, it is carried forward to subsequent windows until another observed value is encountered, which is then in turn carried forward and so on. When there is no value for a given feature in a care episode, it is imputed globally: for categorical features, the most frequent value is chosen, while mean imputation is carried out for numeric features. For SOFA, qSOFA, and NEWS2, missing values are not mean-imputed; instead, the score is assumed to be 0 – if missing – at the start of an episode and then carried forward as described above.

Figure 1 shows the distribution of care episode lengths with different time window sizes and the episode-wise missing rate, i.e. the percentage of missing values in a care episode. Smaller time windows yield longer sequences and a higher rate of missingness, while larger time windows yield shorter sequences but a somewhat smaller amount of missing data. There is high variance with respect to both sequence length and missingness.

### 2.4 Sepsis Definition

The operational Sepsis-3 clinical criteria are used to define sepsis (Singer et al., 2016; Seymour et al., 2016). To fulfill the criteria, patients are required to suffer from a suspected infection in combination with organ dysfunction (Singer et al., 2016; Seymour et al., 2016). Suspected infection is defined as having any culture taken and at least two doses of antimicrobial treatment newly administered within a certain time period. If antimicrobial treatment was initiated first, cultures had to be collected within 24 hours. If cultures were collected first, antimicrobial treatment had to be started within 72 hours after the cultures. Organ dysfunction is measured by an increase in sequen-
Organ dysfunction is measured 48 hours before to 24 hours after the onset of suspected infection. The baseline SOFA score is defined as the latest value measured before the 72-hour time window and is assumed to be 0 in patients not known to have a pre-existing organ dysfunction.

Sepsis onset time is regarded as the first time-window when both organ dysfunction and suspected infection criteria are met. As a time window is used as a fixed length of time interval to represent the temporality, the particular time window in which sepsis onset occurs is regarded as time zero.

2.5 LSTM Model

A deep learning model learns from the input data features, by using hierarchical representations of input features, from lower level compositions to higher level compositions. This type of abstraction at multiple levels allows learning features automatically using the complex formation of functions, which map the input to the output straightforwardly without any need for human feature engineering. (Goodfellow et al., 2016).

In this study, we use a Long Short-Term Memory based Recurrent Neural Network (LSTM) model (Hochreiter and Schmidhuber, 1997) for deep learning. The choice of model is motivated by the longitudinal nature of the EHR data and the task to predict sepsis as early as possible on the basis of current and past information in a given care episode. LSTM models are particularly suited to this type of sequence labeling task, as it can retain information from previous inputs in their internal memory. It also can learn from a very distant past if the information is relevant by using different gated cells (input, forget, and output) where these cells determine what information to store and what information to erase. From a clinical perspective, this is also relevant since typically clinical measurements and observations closer to the outcome is of higher importance in the care episode.

The PyTorch package (Paszke et al., 2017) was used to implement the model on a Dell R730 server with one Intel Xeon E5-2623 processor with 32 MB cache memory extended with a GeForce GTX 1080 GPU. The server ran the operating system Linux Debian 9.1. One training cycle took approximately 60 minutes.

2.6 Baselines

The LSTM model is compared to two baselines in the form of early warning scores that are commonly used in clinical practice today: NEWS2 and qSOFA (Singer et al., 2016). NEWS2 is an updated version of the National Early Warning Score (NEWS) (Williams et al., 2012) and constitutes an aggregate scoring system for each time window in the episode based on physiological measurements of respiration rate, oxygen saturation, systolic blood pressure, pulse rate, level of consciousness or new confusion, and temperature. qSOFA (Quick SOFA) is another aggregate scoring tool that takes into account altered mental status, respiratory rate, level of consciousness or new confusion, and temperature. qSOFA (Quick SOFA) is another aggregate scoring tool that takes into account altered mental status, respiratory rate, level of consciousness or new confusion, and temperature. A score is originally only present in a time window if the data required for calculating the score is also present. If no score is available at the beginning of a care episode, a score of 0 is inserted in the corresponding time window. Existing values are then carried forward to subsequent time windows until a new value is encountered. In order to convert the score to a binary classification decision, a threshold of 5 is used.
for NEWS2 and 2 for qSOFA. These are standard decision thresholds for NEWS2 and qSOFA (Williams et al., 2012; Singer et al., 2016).

2.7 Experimental Setup

The dataset is split into 80% for training, 10% for tuning and 10% for testing and evaluating the tuned models. The data is stratified in each split using an equal probabilistic distribution with respect to both class and sequence length. Care episodes of a sequence length with five or fewer instances are discarded. Positive cases are care episodes in which Sepsis-3 criteria were fulfilled and comprise data from admission to sepsis onset.

Twelve different versions of the dataset are created based on six different time window sizes (1h, 2h, 3h, 4h, 6h, and 8h) and two approaches to handling missing values: with imputation and without imputation. In each time window, the model outputs a probability score concerning the presence or absence of sepsis in the patient based on current and previous information in the care episode.

The models are evaluated using a number of predictive performance metrics. In order to assess the models globally, without a specific decision threshold, we used Area Under the Receiver Operating Characteristic curve (AUROC) and Area Under the Precision-Recall Curve (AUPRC). AUROC is an overall measure of discrimination and can be interpreted as the probability that the model ranks a true positive higher than a false positive. As the data is highly imbalanced, AUPRC is considered the primary metric for model selection.

The hyper-parameters of the LSTM model are presented in Table 1. To tune these parameters, fifty points were chosen at random in order to more effectively search the space instead of doing a grid search in some restricted hyperparameter space. Oversampling method was used to make the distribution even (50% positive, and 50% negative) in each minibatch. The model with the best AUPRC on the tuning set was selected and evaluated on the test set.

Earliness is evaluated in one of two ways: (i) calculating AUROC and AUPRC at different time points before sepsis onset, and (ii) looking at the (median) prediction times relative to sepsis onset (in hours) for true positives. In order to evaluate earliness in terms of (ii), class label predictions need to be made using a decision threshold. We use a standard threshold of > 0.5 and only allowed the model to make a single positive prediction per care episode, retaining the first one and ignoring predictions in subsequent windows. Since earliness calculated in this way depends solely on true positives, it should not be analyzed in isolation, but in conjunction with more conventional performance metrics like sensitivity and positive predictive value. We therefore report F1-score, which is the harmonic mean between precision (positive predictive value) and recall (sensitivity), along with the median earliness in hours. Three evaluation settings are used to calculate earliness according to (ii) depending on the time period in which the evaluation is carried out: (a) <24 hours prior to sepsis onset, (b) <48 hours prior to sepsis onset, or (c) at any time in the care episode. The motivation for the more conservative evaluation settings is to reduce the effect of extremely early predictions, which perhaps should not be considered realistic from a clinical point of view. We also report the temporal distribution of true positive predictions.

2.8 Experiments

A series of experiments were carried out with the use of an LSTM model for early prediction of sepsis in a general hospital population and compared to existing early warning scores in the form of NEWS2 and qSOFA. The experiments are centered around the use of different time window sizes for representing the temporally evolving EHR data. The size of the time window affects (i) sequence length and (ii) missingness. The impact of these factors on the predictive performance of the LSTM model was investigated and two approaches are compared to dealing with missing values. The predictive performance of the LSTM models and the baselines is evaluated using a number of different metrics and the evaluation is carried out in various configurations and time points prior to sepsis onset. Naturally, particular attention is

<table>
<thead>
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<th>Name</th>
<th>Values / Range</th>
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<tr>
<td>alpha</td>
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</tr>
<tr>
<td>beta one</td>
<td>0, 10^{-3}</td>
</tr>
<tr>
<td>beta two</td>
<td>0, 10^{-3}</td>
</tr>
<tr>
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<td>neurons</td>
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<td>drop out</td>
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<tr>
<td>classification function</td>
<td>log-softmax</td>
</tr>
<tr>
<td>optimizer</td>
<td>Adam optimizer</td>
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</table>

Note that the evaluation includes an assessment of the models both in terms of their general predictive performance and in terms of one possible way of employing the model in a clinical setting.
The following five experiments are conducted. The first three experiments look at the predictive performance of all twelve models in general, without using a specific decision threshold. The last two experiments use a standard threshold of $>0.5$ and only take into account the first positive prediction, as described earlier. Post-analyses are also conducted using the best overall model.

**Experiment 1: Different Time Window Sizes.** The impact of using different window sizes is evaluated on the predictive performance of the resulting model. The choice of window size has a substantial impact on the sequence length distribution and the amount of missing values in each time window, as seen in Figure 1.

**Experiment 2: Handling Missing Values.** Two very different approaches are investigated to handling the large amount of missing values in EHR data on a care episode level. In one approach, data is assumed to be missing not at random and therefore no imputation is carried out; instead, the model is allowed to learn to treat missingness as a distinct feature. In the other approach, data is assumed to be missing at random (i.e., for a particular set of features) and is evaluated using a simple imputation strategy based on carrying forward existing values and globally mean-imputing values that are absent at the beginning of a care episode. These two approaches to handling missing values are evaluated for each of the six time window sizes, i.e., with different degrees of missingness.

**Experiment 3: Performance at Different Time Points.** In this experiment, the predictive performance is evaluated, in terms of AUROC and AUPRC, at different time points relative to sepsis onset, starting from 24 hours prior to onset. The LSTM models – with different window sizes and approaches to missingness – are compared to commonly used early warning scores in NEWS2 and qSOFA.

**Experiment 4: Evaluation of Earliness.** In this experiment, a closer look is taken at the distribution of earliness, i.e., at which time points true positive predictions are made relative to sepsis onset, for the best LSTM model. This is shown with a combination of overall $F_1$-score for the particular time window size. This post-analysis is carried out with the results obtained from the previous experiments.

**Experiment 5: Performance with Different Sequence Lengths.** In this experiment, the predictive performance is analyzed, in terms of $F_1$-score, on care episodes of different sequence lengths. The purpose of this analysis is to learn how the best model copes with heterogeneous care episodes in terms of length of hospital stay.
Table 2: Earliness performance in median hours before sepsis onset combined with F\textsubscript{1}-score, using test dataset, with and without imputation of missing values, using different time window sizes.

<table>
<thead>
<tr>
<th>Window Size</th>
<th>Without Imputation</th>
<th></th>
<th>With Imputation</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>&lt;24h</td>
<td>&lt;48h</td>
<td>All</td>
<td>&lt;24h</td>
</tr>
<tr>
<td>Med.</td>
<td>F\textsubscript{1}</td>
<td>Med.</td>
<td>F\textsubscript{1}</td>
<td>Med.</td>
</tr>
<tr>
<td>1</td>
<td>6.00</td>
<td>0.65</td>
<td>7.37</td>
<td>0.53</td>
</tr>
<tr>
<td>2</td>
<td>7.24</td>
<td>0.73</td>
<td>8.00</td>
<td>0.57</td>
</tr>
<tr>
<td>3</td>
<td>12.45</td>
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<td>12.57</td>
<td>0.65</td>
</tr>
<tr>
<td>4</td>
<td>7.13</td>
<td>0.73</td>
<td>7.92</td>
<td>0.63</td>
</tr>
<tr>
<td>6</td>
<td>10.09</td>
<td>0.76</td>
<td>11.00</td>
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<td>8</td>
<td>8.73</td>
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<td>9.82</td>
<td>0.67</td>
</tr>
</tbody>
</table>

3 RESULTS

In this section, the results of the above experiments are presented in terms of (i) predictive performance at different time points, (ii) earliness of true positive predictions, and (iii) predictive performance for episodes of different sequence lengths.

3.1 Temporal Analysis

The overall predictive performance without a specific decision threshold, in terms of AUPRC and AUROC, of the different models is presented at different time points relative to sepsis onset – from as early as 24 hours to sepsis onset time. The results, also depicting the performance of the two baselines, are shown in Figure 2.

As can be seen, the performance of the LSTM models – with and without imputation – is vastly superior to NEWS2 and qSOFA. The performance naturally drops further from sepsis onset and does so quite rapidly. The LSTM models without imputation generally performs better with respect to both AUPRC and AUROC. The best results come from using a two-hour time window, without imputation: at sepsis onset, the AUROC is 0.93 and AUPRC 0.73. However, four hours prior to onset, the AUROC has dropped to 0.88 and AUPRC to 0.54. Finally, eight hours prior to onset, the AUROC and AUPRC has dropped further to 0.85 and 0.44, respectively.

It is clear that the size of the window has a significant impact on the predictive performance of the resulting models. For instance, the difference between a two-hour window and an eight-hour window is more than 20 percentage points w.r.t. AUPRC and 5 percentage points w.r.t. AUROC at the time of sepsis onset.

In general, assuming that data is missing not at random leads to better predictive performance compared to assuming that data is missing at random. However, the difference is is generally smaller compared to the size of the time window.

3.2 Earliness

The median in the distribution of earliness (in hours) for true positive predictions of the models – with different time window sizes and with/without imputation – are shown in Table 2. Earliness results are reported for the three evaluation settings (<24h, <48h and All). Irrespective of evaluation setting, all models are able to predict sepsis more than five hours before onset in half of the cases, with the best median result as early as 24.28 hours prior. We also present the F\textsubscript{1}-scores of the models. F\textsubscript{1}-score is calculated by allowing only one positive prediction in the care episodes where the decision threshold was >0.5. With this particular way of using the model for early prediction, the best result is obtained in the <24h setting with a four-hour time window and imputation of missing values: the F\textsubscript{1}-score is 0.79 and the median earliness is 11.22 hours prior to sepsis onset.

In addition to the median values reported in Table 2, the distribution of true positive predictions with respect to earliness in hour is shown in Figure 3. The distribution is obtained from the overall best model: a two-hour time window and without imputation. When evaluating the entire care episode (All), the median earliness in identifying sepsis is 12.68 hours before sepsis onset. In the more conservative evaluation setting (<24h and <48h), the median is 7.24 and 8.0 hours before onset, respectively.

The distribution of earliness is also shown in Figure 4. Here, each bin represents the successive two hours of prior predictions up to 24 hours. As can be seen, there are relatively more true positive predictions closer to sepsis onset. However, there are also cases of earlier predictions, up to 24 hours prior to sepsis onset. The bin representing earliness >24h in-
Figure 3: The distribution of earliness, using test dataset, measured in terms of hours prior to sepsis onset, for true positive cases in three different evaluation settings: allowing true positives only < 24h or < 48h before sepsis onset, as well as any time (All) prior to sepsis onset. Results are shown for the best model: 2h time window, no imputation.

includes some very early true positive predictions, for example, predictions > 96h prior to sepsis, which can be regarded as outliers.

Figure 4: The distribution of earliness, using test dataset, measured in terms of hours prior to sepsis onset from the best LSTM model (2h time window, no imputation), evaluation setting 'All'.

3.3 Episode Sequence Length

As using different time window sizes has a significant impact on the length of sequences that constitute the care episodes, the predictive performance of the model is also evaluated with respect to sequence length. In Figure 5, the F1-score calculated as a local scope of evaluation (described in section 2.8) at the sepsis onset time (time zero) of our best model is shown. The care episodes in the test set are binned in such a way that each bin contains at least 1000 test instances, ensuring that there is sufficient statistical evidence for estimating performance. As a result, the bins may vary in size: there are, for instance, many episodes of length 1 (n > 1000), i.e. comprising only a single time window (here, 2h), which make up one bin. As seen in Figure 1, the episode sequence length distribution is skewed towards shorter sequences; as a result, the bins encompass more sequence lengths the longer the sequences get.

In general, the model performs better on shorter care episodes. The results are high for sequences comprising fewer than around eleven time windows. The performance degrades quickly with care episodes of length greater than eleven.

4 DISCUSSION

In this study, our aim was not to propose a novel deep learning architecture for early prediction of sepsis. We rather wanted to take a step back and investigate two basic and interconnected assumptions, without empirical or theoretical justification, made by current state-of-the-art models (Futoma et al., 2017a,b; Moor et al., 2019): (i) dividing the temporal EHR data into hourly time windows and prediction times, and (ii) treating missing values in the care episodes as missing at random. We carried out an extensive empirical investigation into these matters and moreover focused on early detection of sepsis in the non-ICU setting, where data is considerably more sparse and heterogeneous. As the choice of window size affects the number of windows in each care episode, we also investigated how the model performs on care episodes of different (sequence) lengths.

The investigation demonstrates that the size of the time window used for dividing up a patient’s longitudinal EHR data has a clear impact on the predictive performance of the model. The size of the time window is hence an important consideration and a design choice that needs to be justified, either empirically or from a clinical point of view. In our non-ICU setting, with heterogeneous data representative of a general hospital population, we found that using a 2-hour time window led to the best predictive performance overall. In an ICU setting, it may very well be that a smaller window leads to better performance since the data is of higher resolution and less sparse. As described earlier, the size of the time window has a significant impact on sparsity per care episode and how often clinical measurements are taken will differ significantly in ICU vs. non-ICU settings. However, as clinical measurements are not taken as frequently
in the non-ICU setting, only considering the "worst" value – as we have done here – for each feature and time window may not be adequate since, in this way, we are discarding potentially valuable information. In future work, a method is needed that better utilizes all of the available data.

An important challenge lies in handling the extreme sparsity (as shown in Figure 1, right) in non-ICU EHR data. In this study, we used 22 features, and the missingness in our data per feature is on average 72.7%, with a median of 87.6% (with a four-hour time window). The variance across care episodes is substantial and could in future be modeled using an end-to-end hybrid attention-based neural architecture (Vaswani et al., 2017), where different attention can be provided to an episode when it is sparse and otherwise. Previous studies in the area consider missingness as a random phenomenon and model it on an evenly spaced grid using the Multitask Gaussian Process (MGP) adapters (Futoma et al., 2017a,b; Moor et al., 2019). However, data in the clinical setting is often missing not at random. In this case, missingness may provide important information about the condition of the patient or the assessment of the treating physician. The meaning of a missing value, and thereby also how it should be handled, may also vary across medical institutions. For instance, if some tests are completed routinely at one location, but only when there is a suspected infection at another, then there will be a significant difference in the predictive contribution of missingness between the two locations. It may therefore be counterproductive to impute missing values in all cases and our preliminary results support this notion. In this study, we experimented with very basic techniques for imputation and modeling missingness as a feature, i.e. using a dummy value. In the future, we plan to represent missingness not at random in a more sophisticated way, for example using advanced Generative Adversarial Networks (GANs) (Li et al., 2019) to model the missing data distribution as not at random. We also plan to explore approaches capable of modeling both data that is missing at random and not at random. Attention-based architectures, as described above, could also be utilized here where both types of missingness can be modeled in an end-to-end manner.

Model evaluation using AUROC and AUPRC is common practice in machine learning, including for the early sepsis prediction task (Futoma et al., 2017a;b; Moor et al., 2019). AUROC and AUPRC provide a global view of predictive performance, measured on a continuum of thresholds values for the classification of patients into sepsis and no sepsis. In order to deploy a machine learning model in a real clinical setting, however, the model should be tuned according to the circumstances and prerequisites of the medical institution in which it will be used. The decision threshold can be optimized according to one or more performance metrics and this choice should be informed by clinical needs – a decision that may depend on, for instance, the tolerance for false positives. In this paper, we used a decision threshold of >0.5 for positive predictions and only allowed one positive prediction per episode. There are many alternative ways of deploying the model, not only using a different decision threshold, but also, for instance, allowing the model to make multiple predictions and silencing the model for some period following a false positive, i.e. according to the attending physician. The results also show that the optimal window size is different when evaluating the model from a general perspective and when employing the model in a specific manner. It indicates that the window size needs to be determined accordingly.

In the future, we will try to modify the neural network-based architecture based on the insights we obtained from this study as described above. We may also explore using a dynamic window size depending, for instance, on the level of sparsity. To enrich our data representation, we will also incorporate free EHR text data with our current structured data, as previous work has shown that it can lead to improved predictive performance for outcome prediction (Henriksson et al., 2015). Multihead attention, in particular, has proven to be the reason for the success...
of state-of-art pre-trained natural language processing embedding models based on BERT (Huang et al., 2019; Alsentzer et al., 2019) for downstream tasks, which can be used in this context. The explainability of the model is a crucial issue here as this can be utilized by different stakeholders (Lipton, 2017). In the future, we will also explore this explainability issue.

5 CONCLUSIONS

We investigated missingness and different time window sizes in extremely sparse EHR data obtained from a Swedish university hospital for the task of early prediction of sepsis using a deep learning-based LSTM model. It was shown that the size of the window has a significant impact on the predictive performance of the models. We also observed that treating missing data as missing not at random can in some cases lead to better predictive performance compared to assuming that it is missing at random.

REFERENCES

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