Early Prediction of Sepsis from Clinical Data Using Single Light-GBM model

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Abstract

— Sepsis is a severe medical condition caused by body's extreme response to an infection leading to tissue damage, organ failure, and even death. The emergence of advanced technologies such as Artificial Intelligence and machine learning, allowed faster exploration of advanced way to recognize sepsis cases. In this paper, we present two main approaches that have been tested using the clinical data. The first method is the combination of survival analysis and neural networks, and the second one is based on booting trees. Our team participated under the name of BERCLAB_UND. The proposed model obtained 0.172 on holdout set and 0.005 on the full test set with ranking of 69.

1. Introduction

Sepsis is a severe medical condition caused by body's extreme response to an infection leading to tissue damage, organ failure, and death. Every 3-4 seconds, at least one person dies because of sepsis worldwide. Sepsis affects about 1.6 million American yearly [1]. As a leading cause of death in the US hospitals, sepsis costs the US about 24 billion USD, 6.2% of the total hospital costs in 2013 [2].

The early detection of sepsis has proven to be a key factor in increasing the efficiency of antibiotic treatment for septic patients. Related studies [3] demonstrated that early detection prevents 80% of death cases caused by sepsis. On the other hand, it was reported that the sepsis mortality significantly increases with the length of stay of the septic patient in the hospital. In other words, delayed recognition of sepsis exacerbates the risk of death of a septic patient by 7.6% every hour [4].

Furthermore, with the emergence of more advanced technologies, there is a significant amount of Electronic Health Records (EHRs) that became available. The EHRs are a systematic collection of data used as health indicators of a patient. The growing availability of the EHRs brought so much interests and opportunities to develop more advanced predictive models to early recognize septic

patients. The electronic health records can be time based features that change over time (time series) like heart rate or blood pressure. Also, they can be static features like demographic information such as age and gender.

The clinical definition of systemic inflammatory response (SIRS) to infection, specifies four conditions which only two of them are sufficient to trigger an alert of sepsis [5]. These conditions are listed as follows:

- 1. Temperature $> 38^{\circ}\text{C} \text{ or } < 36^{\circ}\text{C}$
- 2. White blood cell count > 12,000 per ml or < 4,000 per ml, and > 10% for immature (band) forms
- 3. Heart rate of > 90 beats per minute
- 4. Respiratory rate of > 20 breaths per minute or partial pressure of CO2 of <32 mmHg

The rest of paper is organized as follows: In section II, we outline the task definition of the PhysioNet challenge of this year. The clinical data provided is also discussed and presented. Section III presents the proposed methodology to solve the challenge task. in section IV, we present the obtained results. Finally, we draw the conclusions and some future research directions.

2. Physionet Challenge and Data

Saving septic patients' lives comes with a combination of efforts from different perspective of sepsis. While more advanced instruments allow closer and more accurate follow-up of the septic patient situation, they only can tell the current situation of a patient in a potential risk of sepsis [6].

2.1. Problem definition

The objective of this challenge is to build a machine learning model to predict sepsis 6 hours before the clinical prediction of sepsis.

In most of the time, when hospital patients arrive to the confirmed sepsis shock stage, it's most probably too late to save their lives. Launching the antibiotic culture takes time and is less effective when a patient is in a serious stage of sepsis that became pretty evident.

Medical instruments can tell what have already happened to the patient but alone they cannot tell a lot about the future development of the patient situation. This later information is more relevant for saving infectious patient from death of sepsis.

Each time we detect sepsis patient one hour earlier, we get more 4-8% chance to save the patient's life. With the classical tools of data analysis in the last decade, scientist still face a lot of limitations to detect sepsis early enough [7].

With the emergence of machine learning techniques, a significant progress has been made to improve the current detection tools. [1, 8, 9].

2.2. Related works

The complexity of sepsis is high and depends on so many factors. Several researchers have attempted to propose a predictive model for this purpose, but they are facing some challenges.

Lin Chen, et al. [7, 10] used dynamic EHRs and proposed a generic framework of deep learning to detect septic patients 5 hours earlier. The framework consisted of two experimental setups: one with Convolutional Neural Network (CNN) added before Long Short-Term Memory (LSTM) to extract patterns from time series features (e.g. EHRs). This component will be connected later with a fully connected network. This later component makes use of the static features and extract local characteristics and dependencies in this type of EHRs.

LSTM is very effective in extracting patterns from long sequences. It is always considered as strong candidate to handle sequences like time series signals, videos, or text data

In fact, LSTM [1] is a neural network with a memory component that save the previous inputs and use them along with the current inputs. This is a very efficient strategy to capture the temporal dependencies of the EHRs for sepsis patients. It will help to detect the physiological deterioration in the data earlier than a classical structure of neural network with no memory.

In addition, the CNN [8] is more known for image data rather than temporal data. This structure has achieved great results on many hard topics such as object detection. Temporal data can also be converted into image format and fed to a CNN. This achieve interesting results and shown promising potential of applied CNN on time series.

3. Methodology

In this section we present the data pre-processing stage, features extraction, and an overview of the proposed model.

3.1. Data Features

There are more than 40 health variables used to track the health situation of the patients in this challenge. The 40 columns are classified into three classes: vital signs, lab test, and static variables.

3.1.1. Vitals signs

Most of these features concern the main screening signal that would reflect continuous situation of the patient health. There are 8 vital signs provided in the data and we can cluster them into three main categories: ECG signals, Pulse signals, and Temperature.

- ECG signals: The main columns related are the heart rate, systolic blood pressure and diastolic blood pressure. The severity of sepsis is very correlated to the number of beats per minute. Many studies have shown that the more sepsis get worse, the more we observe ECG abnormalities. This can be explained by loss of excitability in cardiac tissue during the sepsis.
- Pulse signals: Respiration rate, O2Sat, and EtCO2 (End tidal carbon dioxide)
- **Temperature:** Symptoms of sepsis include: a fever above 101°F (38°C) or a temperature below 96.8°F (36°C) heart rate higher than 90 beats per minute. So, tracking the temperature is very important for the early detection task. In fact, the predictive model uses a certain number of the previous data points from the current time and try to detect any significant shift on temperature (drop or rise).

3.1.2. Lab tests

The role of laboratory tests is very significant for early detection of sepsis. Since the main definition of sepsis is the body's systemic inflammatory response to a bacterial infection [11]. The spread of bacteria in the blood (bacteremia) make it a big indicator of sepsis infection. About 25 lab tests have been provided in the data for sepsis detection. These features include: White blood cells count, Blood urea nitrogen, Lactic acid, partial thromboplastin time, Leukocyte count, Platelets.

3.1.3. Demographics features

There were some studies that have shown that sepsis mortality has a little thing to do with demographics. Especially age and length of stay. In our current model, we still find this is not the case. More results will be shown on the next paragraphs [12]. The main demographic features in the data are: gender, age, and length of stay. Surprisingly, we found out in our model, that the age is a critical factor to predict sepsis risk.

3.2. Data Pre-processing & Features

Before the learning step, the clinical dataset we have for the challenge contain a lot of inconsistencies. For our model we have performed many preprocessing techniques to ensure the consistency of the data and create new features. The processing pipeline can be described as follow:

- (1) Handling missing values: Several lab tests features have up to 90% of missing values. The data resolution is hourly based, and it is difficult to perform lab experiments every hour for every patient. Which explain the high ratio of missing values in lab test columns. However, the values in these columns are very important and removing them would not be the best idea. In order to handle missing values:
 - We have performed interpolation which fill a missing value with the mean of the two consecutive nonmissing values.
 - 2. The remaining missing values are filled using backward and then forward values.
- (2) Lag features: The purpose of this features is to capture the long- and short-time dependencies. We performed different sliding windows with mean of the last 1,2,..,9 hours.
- (3) **Data binning**: In order to reduce some variance in the signal columns. We create new features that aggregate the signal value into ranges and intervals. We have based the data binning on the max and min of each signal. There is an option to define range limits using Random Forest. We expect to explore it in future work.
- (4) Count Encoding of Categorical features: applied mainly on the demographics features such as: age and gender.

3.3. Model Overview

While some research works [10,11] showed a great ability to predict sepsis in infectious patients. It was still limited since many of these simulations could not predict earlier than 6 hours. As the time frame increases, the accuracy decreases significantly.

In this section, we outline two approaches to predict sepsis patient no earlier than 12 hours and no later than 6 hours.

3.3.1. Weibull Time-To-Event- RNN model

Early detection is the typical application of Survival Analysis (SA); it has been used in medicine and early detection of catastrophic events.

However, SA despite being able to fit perfectly and handle the censored nature of data used to estimate Time-To-Event (TTE) problems, it still remains a statistical mod-

elling approach and not able to keep up with the advanced approaches of deep learning.

Likewise, the fundamental idea of deep learning architectures still does not take into consideration the censorship aspect of the data as is needed TTE problems. As a result, deep learning itself would not be able to provide high performance on such problems. In this paper, this model could not show a huge performance in the challenge. But we believe it is very promising and deserve to be mentioned. It will be the subject improvement in future work.

In Weibull Time-To-Event- RNN model (WTTE-RNN) [13] model we assume that TTE variable follow Weibull distribution governed by two parameters: alpha and beta. The idea is to estimate the TTE distribution instead of directly estimating the TTE variable. This approach was not very successful, it is still vague how the algorithm is learning in the data. We needed to show more clarity in the model and that's why we thought about boosting trees. Light-GBM was the best candidate.

3.3.2. Light-GBM

This approach is a gradient boosting learning framework that uses tree-based learning algorithm. The difference between light-GBM and other tree-based algorithms is that it grows tree leaf-wise while other algorithms grows levelwise.

In order to solve the class unbalance in the training, we went for a cross validation of 5 stratified folds. The index sampling made patient wise not data point wise. This later would cause a huge data leak and biased training.

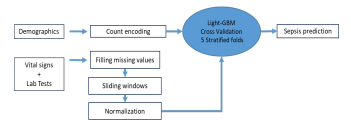


Figure 1. Model Overview

3.4. Results

In this section, the obtained results are presented as well as their analysis. Figure 2 presents the feature importance. From this figure, it can be seen that age is a crucial demographic factor to identify sepsis patients. Besides, lab tests come also on the top of the the critical factors such as Blood urea nitrogen, platelets and partial thromboplastin time. These results are aligned very well with the biological description of sepsis. On the official results, our

team, BERCLAB_UND, got 0.005 as normalized utility score with a ranking of 69.

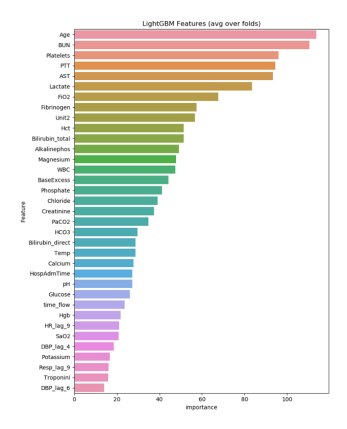


Figure 2. Feature Importance provide better model clarity.

3.5. Conclusion

In this paper, a single Light-GBM model to predict sepsis is presented and implemented. The proposed model could beat a lot of approaches that has been tested in the challenge. The strength of the proposed model in this paper is in the simplicity and speed of processing. It will be much more easier to implement it in the medical devices with lower capacity. Further improvement of the normalized utility score of this model will be explored.

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