Assessment of the timeliness and robustness for predicting adult sepsis

Yuanfang Guan, Xueqing Wang, Xianghao Chen, Daiyao Yi, Luyao Chen, Xiaoqian Jiang

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HIGHLIGHTS

We report a top-performing algorithm for sepsis prediction
We analyzed the timeliness and robustness for predicting sepsis
Assessment of the timeliness and robustness for predicting adult sepsis

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SUMMARY

Sepsis is a leading cause of death among inpatients at hospitals. However, with early detection, death rate can drop substantially. In this study, we present the top-performing algorithm for Sepsis II prediction in the DII National Data Science Challenge using the Cerner Health Facts data involving more than 100,000 adult patients. This large sample size allowed us to dissect the predictability by age-groups, race, genders, and care settings and up to 192 hr of sepsis onset. This large data collection also allowed us to conclude that the last six biometric records on average are informative to the prediction of sepsis. We identified biomarkers that are common across the treatment time and novel biomarkers that are uniquely presented for early prediction. The algorithms showed meaningful signals days ahead of sepsis onset, supporting the potential of reducing death rate by focusing on high-risk populations identified from heterogeneous data integration.

INTRODUCTION

Sepsis is a life-threatening condition that occurs when the body’s response to an infection causes tissue damage, organ failure, or death (Singer et al., 2016). In the U.S., nearly 1.7 million people develop sepsis and 270,000 people die from it each year (CDC, 2020a). Additionally, over one-third of people who die in U.S. hospitals have sepsis (CDC, 2020b). Internationally, an estimated 30 million people develop sepsis and 6 million people die from sepsis each year (Kumar et al., 2019), and an estimated 4.2 million newborns and children are affected (Demirer et al., 2019). Due to its high mortality rate, fast deterioration, and difficulty in treatment, sepsis has become a major public health concern around the world. It accounts for more than $23.66 billion (6.2%) of total US hospital costs in 2013 (Torio and Moore, 2016).

Death rate of sepsis can be significantly reduced by early diagnosis. It is estimated that for every hour sepsis goes undiagnosed, the death rate increases between 4 and 8% (Kumar et al., 2006; Seymour et al., 2017). Though the sepsis research community has been indefatigably working on establishing models predicting sepsis onset, three important questions remain. First, how far in advance can sepsis be detected and what is the corresponding accuracy? Second, how much data do we need to gather so that we can confidently predict a person will develop sepsis? Third, how does model performance differ among genders, races, age-groups, and care settings? Furthermore, it is yet to be discovered whether novel biomarkers can be revealed by the state-of-the-art machine learning techniques and whether a more rationally weighted strategy could be used to predict sepsis.

In this study, we address the above challenges by taking advantage of a large, heterogeneous cohort of 100,000 patients collected from a various set of care settings. Among the patients, 31,377 patients were septic, and others were matched non-septic individuals based on gender, age, admission type, and length of hospital stay. Previous large-scale studies include Barton et al. that used 3,679 patients with sepsis up to 48 hr of septic onset (Barton et al., 2019), Nemati et al. that used ~5800 patients (Nemati et al., 2018), and Mao et al. that used 4107 patients (Mao et al., 2018). There were other studies that used a less number of patients (hundreds) with sepsis (Le et al., 2019; Schamoni et al., 2019). A recent community benchmark study used 2,921 patients with sepsis for the training set (Reyna et al., 2019). The other large-scale study is Komorowski et al. (47,220 septic patients), and it focused on predicting treatment outcome instead of detecting sepsis (Komorowski et al., 2018). The other large study is Delahanty et al. (2019) which included 54,661 patients with sepsis and examined gradient
boosted trees in predicting sepsis (Delahanty et al., 2019). Overall, the size of this study is satisfactory to evaluate prediction modeling of sepsis, which also allowed us to examine the performance and predictive features up to hundreds of hours ahead of sepsis onset. The validity of this study is further supported by a community-wise benchmark study, DII National Data Science Challenge, which used this data set to benchmark performance, and the results presented here are based on the top-ranking algorithm. We dissected the model performance by genders, races, age-groups, and, most importantly, care settings. Then, we determined biomarkers (including early biomarkers) for predicting sepsis onset, as well as patterns related to forecasting time and segments of the data that are needed to build the model.

RESULTS
Non-linear model capturing changes over time to predict sepsis
The data were obtained from the DII National Data Science Challenge, which originally came from the Cerner Health Facts database (Table S1). The total patient number is 106 million, and the hospitalization population is 297 thousands. Among the hospitalized patients, we selected patients who were hospitalized for at least 8 hr before sepsis onset, and consequently, anyone that was septic prior to admission was also excluded. Seven thousand seven hundred thirty two septic examples were excluded due to insufficient observation windows. Furthermore, we dropped patients who did not have medication, labs, or events records. Otherwise, missingness can be a confounding factor in studying predictive factors. The negative examples are sub-sampled by propensity score matched to the patients with sepsis based on age, gender, race, admission type, and length of stay. Because the data we will take to predict sepsis are up to 4 hr before sepsis onset, we also truncated the negative examples at their respective matched sepsis onset time of patients with sepsis—4 hr (Figure 1). According to the matched pairs, we truncated the vital data with the same length from admission to sepsis onset (and later to 4 hr, 5 hr, 6 hr, etc ahead of sepsis onset time). Otherwise, the length of hospital stay will bias the analytic results. This step allows a reasonable number of examples for training, as well as a matched population. The selected cases include a mixture of care settings (17,870 urgent care cases, 1,089 from trauma centers, 63,769 from emergency care centers, 20,888 from elective treatment, and 2,675 unknowns). Among them, 31,377 patients were septic.

Other than typical sepsis diagnosis criteria, the data also required the inclusion of patients (septic or non-septic) to be those that must have a minimum of eight hours of hospitalization records. The rationale behind the choice of eight hours is that we attempt to construct a model that can be used in clinical settings to predict sepsis ahead of its onset. Features that are available for constructing models can be categorized into two types: demographic data including gender, race, admission type, admission source, care settings, and age-group (Table S1, Figure 1), and longitudinal data including event time and different types of biometrics, such as albumin, alkaline phosphatase, heart rate, blood pressure (Table S1, Figure 1). Outcome is indicated by a binary label of 1 for septic or 0 for non-septic. This data set followed the Sepsis-II definition (Güll et al., 2017; Obonyo et al., 2018; Patki, 2018) (see challenge webpage [https://sbmi.uth.edu/dii-challenge/usecase.html]). SIRS (systemic inflammatory response syndrome) was defined as meeting the 2/4 criteria within ±3 hr from the point the suspect infection (when the microbiology order was placed). Based on discussion with the Cerner clinical and statistical consultants, concerns were raised regarding retrospectively recalculating a Sepsis-III score or other scores without bias, due to inconsistency of old data with the new definition that leaves many samples in the database unqualified. The clinical and statistical consultant recommended that using Sepsis-II corresponding to the years of collection was more appropriate. We acknowledge that this presents a limitation of this study, in which Sepsis-III was not used. As the data of this long-spanning study were collected during the period of Sepsis-II, we focus on Sepsis-II in this analysis.

We one-hot encoded categorical, demographic information into 43 discrete features (Figure 1, Tables S1 and S2). For the longitudinal data, each time point, which we will term as “record” in the rest of this paper, is represented by a vector representing measurements of diverse biometrics (Figure 1). This results in 144 features for each record. Although the data are produced at a resolution of 30 min, many of the time points have no biometric measurement available (missing data). We started with a base model taking the last eight records, which were recorded a median of 65 hr before sepsis onset—4 hr, ranging from 0 (in case of only one record) to 700 hr. This gives us a total of 1195 features (144X8+43) for each patient. The features are listed in Table S3.

Apart from the above basic predictive features, we included the following clinically relevant features (Figures 1 and 2A, Table S2). First, we recorded the first available measurement for each feature, as well as their
corresponding time of records relevant to sepsis onset. These features were included to approximate the baseline biometric values since these values could be quite different for every patient. Second, we calculated the slope of each biometric by dividing between the difference of the first and last available measurement by the time difference. These features provide information of the changes over time. Third, the mean and standard deviation of non-missing entries are recorded as additional features. This set of additional features profile the whole time series of a patient. Fourth, using the mean and standard deviation of each feature, we normalized observed features and created a new set of features, of which we took the last eight records. When the training set and test

Figure 1. Overview of model construction for selecting patients, predicting sepsis onset, and feature importance analysis

We removed the hospitalized patients without sufficient observation time and selected negative examples according to matched age, gender, hospital stay time, and admission type. Data were truncated by the matched sepsis onset time—4 hr. Input data can be categorized into two sets: demographics and time series records. All demographic features are categorical, and thus we one-hot encoded them into multiple binary features. For time series data, the last eight points of records were included in the baseline model. A variety of additional features including slope, mean, maximal and minimal values for each feature, and first time observations were included as additional features. Diverse base learners were compared for their classification performance. We further tested how much time ahead the model can predict sepsis and the number of useful records for good model performance. Lastly, feature importance was analyzed using SHAP values (the SHAP figure is only for illustration purposes, modified from [https://github.com/slundberg/shap]), and grouped by time points and feature categories. See also Tables S1 and S2.
set are drawn from the same population, the fourth set of features may not add additional value. We included this set of features to account for batch effect and potentially improve the robustness of the model when delivered to new cohorts. Through five-fold cross-validation, we found the above features progressively improved the model performance from area under the receiver operating characteristic curve (AUC) = 0.78178 to AUC = 0.84854 (specificity = 0.77 at 0.8 recall and specificity = 0.827 at 0.7 recall, Figure 2B, p < 1e-5). The partial AUC above 0.8 recall (pAUC80) is 0.128, compared to a random baseline of 0.02 (Figures 2 and 3); the partial AUC above 0.7 recall (pAUC70) is 0.181, compared to a random baseline of 0.045. The most useful additional features are slope and baseline observations, reflecting the importance of capturing changes of biometrics in predicting sepsis.

We further explored a range of base learners and compared their performance in learning information from the above features (Figures 2C and 3A). Overall, we found LightGBM showed the best performance (AUC = 0.84854), followed by gradient boosting trees (0.81904, p < 1e-5) and random forest (0.80312). The other base learners did not show as competitive performance as the above: linear regression (0.7779), Naive Bayes (0.53812), quadratic discriminant analysis (0.58206). LightGBM further demonstrated advantage in training speed (Figure S1). Thus, in the following analysis, we focused on the property of the LightGBM models built with all features described above. Of note, none of the linear or posterior inference models demonstrated satisfactory performance, which corroborates that the relationships between the features are largely non-linear and thus simplistic cutoffs on biometrics to predict sepsis onset are not ideal. This effect prompted us to further categorize and investigate the predictive features in later sections.

Figure 2. Post-processing of features can improve model performance.
(A) Illustration of primary features and additional features: missing values are removed from calculating the mean, standard deviation, first available values, and time for each longitudinal feature.
(B) Performance improvement by adding derived features separately to the base model.
(C) Performance using different base learners on the features. In the boxplots in (B) and (C), center lines refer to median performance, bounds of box refer to the first quartile and the third quartile of the data, whiskers refer to min and max of the data (except outliers), and spheres refer to outliers which are defined as data points below Q1 - 1.5IQR or beyond Q3 + 1.5IQR.
See also Figure S1, Table S2.
Determination of how much time ahead and the number of observations is needed for sepsis prediction in different clinical settings

One critical, unanswered question is how far in advance sepsis can be detected or whether we can prioritize at-risk populations. This data set provides a valuable resource to examine this question for a large time range. To determine this duration, we progressively cut the data toward the beginning of the records from four hours from sepsis onset to 192 hr (8 days, Figure 4A), at which we still have a reasonable number of patients with sepsis (2579 patients with sepsis and 8679 total patients, Figure S2). Certainly, for days of records, we may no longer directly predict sepsis but more focus on prioritization of at-risk population. Of note, the control examples are cut off correspondingly from the end by record hours. As expected, the

Figure 3. Evaluation by specificity and pAUC
Specificity and pAUC at 80% recall for different (A) base learners, (B) feature sets, (C) age-groups, (D) admission types, (E) gender, (F) race. See also Figure S8.

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Figure 4. Performance comparison in different time and population settings

(A) We first tested how many hours ahead we can predict sepsis onset by progressively cutting off data according to record time.

(B) Model performance by cutting the records according to how many hours ahead of sepsis onset to be included. The performance drops as we attempt to make predictions earlier, i.e., more hours ahead of sepsis.

(C) Illustration of selecting last $n$ records for model input.

(D) Performance improves as more records ahead of sepsis were included.

(E) Model performance by patients from different admission types.

(F) Model performance by different age groups.

(G) Model performance by different genders.

(H) Model performance by different races. In the boxplots in (B) to (H), center lines refer to median performance, bounds of box refer to the first quartile and the third quartile of the data, whiskers refer to min and max of the data (except outliers), and spheres refer to outliers which are defined as data points below $Q1 - 1.5IQR$ or beyond $Q3 + 1.5IQR$. See also Figures S2, S3, and S8.
performance of the model drops as we include less proximate information of the records from 0.84854 (4 hr ahead) to 0.70818 (192 hr ahead). At 12 hr ahead, the performance was 0.8078, at 24 hr ahead, the performance was 0.78836, and at 48 hr ahead, the performance was 0.77628 (Figure 4B). These performances indicate that meaningful models can be established for identifying sepsis at least 1-2 days ahead of onset.

We next examined the minimal number of longitudinal records that are needed to predict sepsis well. The rationale behind this analysis is to infer the amount of information we lost if we limit our view to the number of records we look at, as clinicians might focus on the 1-2 most recent records. We thus started with taking one, two, three, and progressively increasing the number of biometric records we include in the model (Figure 4C). We found that until the six records, the model still makes meaningful additions in performance. We focused on examining the models without additional features so that the effect of derived features from the entire time course will not affect this evaluation. Using the last record only, the model had an average AUC of 0.73382, using the last two records, the average AUC was 0.74764 (p < 1e-5, compared to using the last one); using the last three records, the average AUC was 0.75974 (p = 1e-5 compared to using the last two); using the last four records, the average AUC was 0.76654 (p = 0.0001, compared to using the last three); using the last five records, the average AUC was 0.7707 (p = 0.0281, compared to using the last four); and using the last six records, the average AUC was 0.77964 (p < 1e-5, compared to using the last five), and using the last seven records, the average AUC was 0.77762, which is slightly worse than last six records (p = 0.1797, Figure 3D). Certainly, all these are significantly worse (p < 1e-5) than taking into account additional features (AUC: 0.84854), which capture the dynamics and overall profiles of patients.

Sepsis prediction model performance is robust across care settings, age-groups, genders, and races

Previous meta-analysis has shown that diagnostic tests for sepsis gathered from different admission types and care settings are drastically different in their accuracy, ranging 0.68–0.99 in ICUs, 0.96–0.98 in hospitals, and 0.87–0.97 in emergency departments (Fleuren et al., 2020). It is therefore natural to hypothesize that prediction models may also differ in their accuracy by the data collection source. We thus separated the test set by the data source and evaluated the performance by admission types. Records resulted from elective procedures showed the best performance of AUC = 0.8528 (specificity = 0.780 at 0.8 recall, pAUC80 = 0.132), followed by urgent care (AUC = 0.849, specificity = 0.771 at 0.8 recall, pAUC80 = 0.129) and emergency admission (AUC = 0.8473, specificity = 0.754 at 0.8 recall, pAUC80 = 0.127); records from trauma center (AUC = 0.8343, pAUC80 = 0.129) showed slightly weaker performance (Figures 3D, 4E, and S8).

We went on to evaluate the performance separated by different age-groups and overall found minimal difference. Performance is lower for the young age group, possibly due to smaller sample size, <20 (AUC = 0.833, specificity = 0.731 at 0.8 recall, pAUC80 = 0.124), 20–30 (AUC = 0.836, specificity = 0.754 at 0.8 recall, pAUC80 = 0.119), 30–40 (AUC = 0.840, specificity = 0.752 at 0.8 recall, pAUC80 = 0.124) and generally performance increases as the patient population gets older, 40–50 (AUC = 0.847, specificity = 0.780 at 0.8 recall, pAUC80 = 0.123), 50–60 (AUC = 0.849, specificity = 0.763 at 0.8 recall, pAUC80 = 0.129), and 60–70 (AUC = 0.844, specificity = 0.762 at 0.8 recall, pAUC80 = 0.126), 70–80 (AUC = 0.847, specificity = 0.777 at 0.8 recall, pAUC80 = 0.128), >80 (AUC = 0.851, specificity = 0.776 at 0.8 recall, pAUC80 = 0.131) (Figures 4F, 3C, and S9). There was no difference between females (AUC = 0.850, specificity = 0.776 at 0.8 recall, pAUC80 = 0.128) and males (AUC = 0.846, specificity = 0.764 at 0.8 recall, pAUC80 = 0.127) (Figures 4G, 3E, and S9). The model performs slightly worse for Asian (AUC = 0.836, 2% of total population, p = 0.0263 and p = 0.1129; respectfully, specificity = 0.755 at 0.8 recall, pAUC80 = 0.117) than African American (AUC = 0.852; specificity = 0.788 at 0.8 recall, pAUC80 = 0.130, 18% of total population) and Caucasians (AUC = 0.844, specificity = 0.763 at 0.8 recall, pAUC80 = 0.127, 73% of total population, ), possibly related to a much smaller population size for Asian (Figures 4H, 3F, and S9).

Game theory-based feature analysis reveals important players in predicting sepsis onset

The above described model integrates discrete demographic and clinical features. This allows us to investigate the important factors predicting sepsis. A technical challenge in finding independent contribution is addressed by a recent advance in game theory application: an improved SHapley Additive exPlanation (SHAP) analysis, which substantially improved the speed of calculation and made the Shapley values feasible to obtain for large-scale feature analysis (Lundberg et al., 2018; Shapley, 1988). Mimicking the process of finding out the contribution of players in a football game, the SHAP analysis assigns the independent contribution of each of the features considering the existence of other features. This feature contribution analysis can be carried out for the prediction of an individual patient (Figure S3 for examples).
or summarized for each feature. Compared to direct correlation analysis, SHAP analysis is more capable of addressing confounding features as a consequence of shared patterns with another important feature.

We implemented SHAP analysis with LightGBM and identified the most important features for predicting sepsis. The top features are the most recent records of heart rate and respiration rate, followed by temperature and end-tidal CO2, i.e., parameters related to cardiovascular and respiratory functions (Figures 5A and S4, Table S3). This is not a circulation in feature analysis, as we are restricting the data input far before the actual sepsis diagnosis; especially, as will be shown later, these factors remain to be important when we are looking ahead much time. This indicates that the model is capable of capturing and integrating early signs of sepsis before a diagnosis using the SIRS criteria can be made. Being in the coronary care unit was identified as an important risk factor among the demographic information (Figures 5B, S4, and S5, Table S3).

Because SHAP values are directly additive, we can combine the features in several ways to either eliminate the time factor in feature analysis. Specifically, we combined the SHAP values of longitudinal data by the type of measurement (Table S4, Figure S6). The top features are a group that related to heart rate (HR_monitored (4.4e-3), HR (2.9e-3), HR_Apical (9.4e-4), peripheral pulse (2.3e-4), pulse (1.7e-4)), a group related to respiration function (respiration rate (2.7e-3), end-tidal CO2 (2.8e-4), oxygen saturation/SPO2 (2.2e-4), partial pressure of CO2/PCO2 (1.4e-4), partial pressure of oxygen/PaO2 (1.1e-4), tidal volume (1.3e-4)), a group related to body temperature (temperature (6.6e-4), oral temperature (3.6e-4)), important biometrics (white blood cell (2.0e-4), Braden scale (1.9e-4), Glasgow coma score (1.7e-4)) (Figure 5C). Again, the majority of the parameters are closely related to respiratory function and heart rate and lastly body temperature, which by itself is a sepsis diagnosis criterion.

We next combined the absolute shap values of longitudinal data by the proximity to sepsis onset—4 hr, in order to study the relative contribution of all features related to a specific record. As expected, the most recent record is most predictive of sepsis, showing an average SHAP value of 3.6e-4, and it declines gradually as the records go...
further away from the onset of sepsis (Figure 5D). Records from fourth to eighth records ahead of sepsis are of almost the same SHAP values, indicating that these biometrics represent certain baseline characteristics that put the patients at risk. Yet, at least the second (1.7e-4) and third (1.1e-4) records should be seriously considered, as the SHAP values are at a similar scale for the most recent records. This result is consistent with the progressive cut experiment we carried out in the previous sections and supports that longitudinal analysis of biometric data is necessary to establish accurate models for sepsis prediction.

Identification of predictive features for sepsis onset depending on how much time ahead we make the predictions

To investigate whether the predictive features will change as we change the numbers of hours ahead of sepsis as the input data, we separately calculated the feature importance at each time point from 4 hrs ahead of sepsis onset till 192 hrs ahead of sepsis onset. At each hour cutoff, we carried out 5-fold cross-validation and correspondingly generated five sets of SHAP values for each feature. We then took the average of the five SHAP values as the importance for each feature. The SHAP values are additive, while the contributions of standard deviation, mean, slope, and others may be either negative or positive (depending on the correlation direction). Thus, we next mapped the processed features back to their original features and took the sum of the absolute value for each group to represent the original feature importance. For example, the time series, the mean, the standard deviation, the normalized values of HR will be grouped into one single feature importance for HR. This process was repeated for the models of 4 hrs ahead till 192 hrs ahead (Figure 6A). For each time point cutoff, we took the top 5 features, and this gave us a total of 7 features across all time points: HR, monitored HR (HR_Monitored), intensive care setting, respiration rate, hematocrit (HCT), Caucasian, and diastolic blood pressure (DBP).

We found that although the majority of these top features appear to be important throughout the time course (e.g., HR, respiration rate), others showed dynamics in their importance in predicting sepsis. For example, HCT only becomes an important predictive feature when the time cutoff is above 136 hr. DBP has a stable increase in its importance as we require further time ahead to make predictions. The importance of intensive care setting quickly drops as the time cutoff approaches sepsis onset (around 48 hr) (Figure 6B). These observations support dynamic changes of predictive features for sepsis onset, from predicting which population is at risk to which individuals’ biometrics reflect potential sepsis onset. As in clinical settings, it is impossible to know ahead of time how far away a patient is to sepsis onset; this result suggests the importance to consider both early features and late features mentioned above in analysis.

DISCUSSION

In this study, we presented the top-performing algorithm in the DII National Data Science Challenge for sepsis prediction, involving the largest sepsis detection study to date (with over 30,000 patients with sepsis). This data set gave us the unique opportunity to dissect the model performance, depending on a variety of care settings, genders, races, and age-group factors. We found being Asian is the only factor that negatively affects model performance, while performances of different care settings do not differ.
statistically. This result is rather surprising considering that physicians’ diagnosis for sepsis was reported to differ in accuracy in different care settings (Fleuren et al., 2020).

Furthermore, this large-scale data collection allowed us to estimate the number of useful records to make reliable predictions. We found the records are useful until the six records prior to sepsis onset—4 hr. This has important clinical decision implications, highlighting the importance to examine longitudinal changes of a patient in determining his/her risk in developing sepsis. Indeed, we found that the slope of biometrics was one of the top contributing processed features. Additionally, we step wisely estimated the predictability by time ahead of onset by progressively removing the records prior to onset and thus provided a reference to confidence of the predictions. We found even back to 24-48 hr ahead, the performance of the model remains strong, and up to eight days, we can still prioritize at-risk population, which supports the notion of early detection of sepsis to reduce death rate.

Respiratory function, HR, and body temperature as expected are the three major features for early sepsis prediction throughout the time course, even much earlier than a formal sepsis diagnosis. Previous studies have pointed to the importance of HR variability for prioritizing patients at high risk of sepsis (Aboab et al., 2008; de Castilho et al., 2018; Tang et al., 2009). We found that a fast HR is a strong indicator of sepsis. Similarly, fast respiration rate is indicative of sepsis onset. For very early time points, variation of HCT appeared as a strong, novel biomarker for sepsis onset prediction.

We would like to point out several limitations and future directions of the study. First, one potential limitation is the usage of Sepsis II criteria in this study. The primary reasons we chose Sepsis II instead of Sepsis III are the relative amount of data available and the clinical usage of this model. Sepsis III requires organ failure (SOFA criterion). Validation of the performance using similarly collected Sepsis III data is an important next step. Second, although the data set is large and the data source is heterogeneous in this study, it will be informative to further validate the models using independent cohorts. Third, while this study focused on cross-validation evaluation, new insights might be brought in if perspective data can be collected and used for validation.

Previous studies have reported a range of performance, which are reexamined in this study. First, there is a strong variation in reported accuracies, ranging from ~0.65 to 0.9 (e.g., reported results in (Barton et al., 2019; Gwadry-Sridhar et al., 2011; Liu et al., 2019; Mao et al., 2018; Michelson et al., 2019; Nemati et al., 2018; Taylor et al., 2016)). It is unclear if these differences come from algorithms, populations, the time frame of the data, or care settings. Using the top-performing algorithm in a benchmark study, we dissected each of the above factors using a very large data set and now give an estimation of the expected performance by care settings, genders, age-groups, races, numbers of hours ahead, and number of records. We conclude that the time frame of the data appears as a major influential factor, while predictive features may change along the time course. This information can serve as an important reference for future studies and applications.

Limitations of studies

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Resource Availability

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Yuanfang Guan (gyuanfan@umich.edu).

Material availability

This study did not generate new unique reagents.
Data and code availability
Code is available at [https://github.com/GuanLab/sepsis].

Methods
All methods can be found in the accompanying Transparent methods supplemental file.

SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.isci.2021.102106.

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AUTHOR CONTRIBUTIONS
Challenge data preparation: X.J. and L.C. Implementation of challenge model: Y.G. and X.C. Post-challenge analysis and manuscript writing: Y.G. Figures: X.W. and D.Y. All authors edited and approved the submission.

DECLARATION OF INTERESTS
On behalf of all authors, the corresponding author states that there is no conflict of interest.

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Supplemental Information

Assessment of the timeliness and robustness for predicting adult sepsis

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Supplementary materials

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Methods

Dataset source

Cerner Health Facts® is a database that comprises de-identified EHR data from over 600 participating Cerner client hospitals and clinics in the United States and represents over 106 million unique patients (“Cerner,” n.d.). With this longitudinal, relational database—reflecting data from 2000-2016—researchers can analyze detailed sets of de-identified clinical data at the patient level. Types of data available include demographics, encounters, diagnoses, procedures, lab results, medication orders, medication administration, vital signs, microbiology, surgical cases, other clinical observations, and health systems attributes (see Table S1 for demographics, and Table S3 for complete list of biometrics included in this study).

Machine learning implementation and parameters

As the feature extraction methods have been laid out in the result session, here we will focus on presenting the parameters used in each of the base learners. The final model used LightGBM as a base learner, ‘gbdt’ as boosting type, ‘regression’ as objectives, and number of leaves equal to 150, a learning rate of 0.05, and regulatory alpha as 2. We run the models for 1000 boosting rounds. The above parameters were identified through an intensive grid search during cross-validation. Minor adjustment of these parameters does not affect the performance substantially.

The random forest model used a maximal depth of 10, and a total of 200 estimators. The gradient boosting model used a learning rate of 0.1 and a total of 200 estimators. All other base learners used default parameters and were implemented using the scikit-learn package (Garreta and Moncecchi, 2013).

Cross-validation and performance measurement

We carried out five-fold cross-validation to evaluate and compare the performance of diverse models. Briefly, the dataset was randomly separated into five parts. In each iteration, four of the five parts are used as training examples, and the other part is used as testing examples. The performance was evaluated using the area under the receiver operating characteristic (AUROC).

Challenge final model assembly

In the result session, we focused on presenting the best-performing single model in predicting sepsis. Additionally, we identified a set of models that perform similarly to the best single model, but complement the single model and improve its performance if these models are assembled together. These models are often small variations of the original model.

The first variation is filling in the missing values as -5000 (compared to the original model where the NaNs are not filled and ignored in determining the tree-splitting), and removing the slope features. The second variation is removing the slope features. The next few variations are the above models running on data imputed with missing time points. For example, if between 0.5 and 1.5 hours, the 0.5 hour is missing, we impute this value as additional features. In supplementary Figure S7, we presented the performance of each of the models and their assembled performance.
Evaluation metrics and statistical significance tests

We used three metrics in this paper to evaluate the performance: Area under the Receiver Operating Characteristics curve (AUC) as a global measurement, and specificity at 70% and 80% recall and partial AUC above 70% and 80% recall as evaluations of the predictive values of the models.

AUC is calculated using all individuals by taking various thresholds of false positive rate (FPR, or 1-specificity) and generating corresponding true positive rate (TPR, or recall). Recalls come from TPTP+FN, and specificity come from TNTN+FP. By connecting the points created at different thresholds, we draw a curve, which is the Receiver Operating Characteristics curve, and then we calculated the area under this curve as the AUC values reported in this study. Additionally, we reported specificity at 80% recall (in the paper) and specificity at 70% recall (in the supplementary materials).

Precisions come from TPTP+FP.

We presented and compared the AUC values of two methods in multiple places of this paper, or those of two sets of feature input, or those of two populations. In order to estimate the statistical significance of the differences in AUCs for each pair, we used a non-parametric approach to estimate the significance level. Specifically, we bootstrapped the examples for 10,000 times, and computed the number of times (n) of approach A out-performing approach B, if B is overall the better-performing one, and used \( p = n/10000 \) as the significance values. We used \( p = 0.05 \) as the significance cutoff throughout the paper.

In order to further assess the ability of the model to exclude negative examples while picking up positive ones, we also calculated partial AUC, which is calculated by taking the area under the curve above a specified level of recall. In this study, we used partial AUC at 80% recall (pAUC80) in the paper and partial AUC at 70% recall (pAUC70) as additional information in the supplementary materials.
Figure S1. Training time of different models. Related to Figure 2.
Figure S2. Numbers of total patients and sepsis patients at various hours ahead of sepsis onset. For non-septic patients, we truncated the hours to the same according to their specific propensity score-matched sepsis examples. Related to Figure 4.
Figure S3. Example single patient feature contribution plots. Related to Figure 4.
Figure S4. Top 50 most important features for sepsis onset prediction, separated by the occurrence of the record. HR_monitored was labelled as monitored heart rate in the Cerner data, and HR was labelled as the apical heart rate in the Cerner data. Related to Figure 5.
Figure S5. SHAP importance of demographic features for sepsis onset prediction. Related to Figure 5.
Figure S6. Top 50 most important biometrics features for sepsis onset prediction. For each type of feature, we combined the SHAP values across records in the time series data. Related to Figure 5.
Figure S7. Imputation methods and their effect on model performance. Related to Figure 2.
Figure S8. Specificity and pAUC at 70% recall for different demographic groups and admission types. Related to Figure 3-4.
Table S1. Demographic summary of Cerner data used in this study. Related to Figure 1.
| Aspects of Feature Engineering | Specific Method          | Detailed Description/Explanation                                                                                                                                                                                                 |
|--------------------------------|--------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| categorical data               | one-hot binary encoding  | Transformed into a set where the number of features is the number of total categories                                                                                                                                               |
|                                | numbered categories      | Some base learners (LightGBM) have a default method for categorical data. For each feature, the categories were translated as different numbers and the categorical features were specified.                                               |
| Missing data                   | 0                        | use 0                                                                                                                                                                                                                              |
|                                | Large negative           | use an arbitrary negative number with large absolute value (e.g. -2048, -5000)                                                                                                                                                     |
|                                | NaN                      | use default indication of missing value (numpy.nan).                                                                                                                                                                              |
|                                | Interpolation            | use predicted/interpolated value (e.g. average)                                                                                                                                                                                   |
| Features                       | Last n events            | Extract the last n records before sepsis onset. This ensures rawest information fed into the training step.                                                                                                                       |
|                                | Last t time              | Containing empty time when data is not available. This emphasizes data availability.                                                                                                                                            |
|                                | First available record   | Recording the first available data of each feature, as well as the time of the first occurrence.                                                                                                                                  |
|                                | Slope                    | Calculate the difference between the last available data and divide by the time difference.                                                                                                                                       |
|                                | Fractional max-pooling   | Split the timeline into n parts (time points are rounded to 0.5), and extract the maxpool of each feature for each part. This takes in the full-time length.                                                                           |
|                                | Basic stats              | Mean, standard deviation of the patient data.                                                                                                                                                                                      |
|                                | Count of missing         | Count the number of missing values of each column.                                                                                                                                                                                |
Table S3. SHAP values for all features. Related to Figure 5.
Table S4. SHAP values grouped by longitudinal parameters. Related to Figure 5.