

Using Machine Learning and the Electronic Health Record to Predict Complicated *Clostridium difficile* Infection

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Background. *Clostridium (Clostridioides) difficile* infection (CDI) is a health care–associated infection that can lead to serious complications. Potential complications include intensive care unit (ICU) admission, development of toxic megacolon, need for colectomy, and death. However, identifying the patients most likely to develop complicated CDI is challenging. To this end, we explored the utility of a machine learning (ML) approach for patient risk stratification for complications using electronic health record (EHR) data.

Methods. We considered adult patients diagnosed with CDI between October 2010 and January 2013 at the University of Michigan hospitals. Cases were labeled complicated if the infection resulted in ICU admission, colectomy, or 30-day mortality. Leveraging EHR data, we trained a model to predict subsequent complications on each of the 3 days after diagnosis. We compared our EHR-based model to one based on a small set of manually curated features. We evaluated model performance using a held-out data set in terms of the area under the receiver operating characteristic curve (AUROC).

Results. Of 1118 cases of CDI, 8% became complicated. On the day of diagnosis, the model achieved an AUROC of 0.69 (95% confidence interval [CI], 0.55–0.83). Using data extracted 2 days after CDI diagnosis, performance increased (AUROC, 0.90; 95% CI, 0.83–0.95), outperforming a model based on a curated set of features (AUROC, 0.84; 95% CI, 0.75–0.91).

Conclusions. Using EHR data, we can accurately stratify CDI cases according to their risk of developing complications. Such an approach could be used to guide future clinical studies investigating interventions that could prevent or mitigate complicated CDI.

Keywords. *Clostridium (Clostridioides) difficile* infection; complications; electronic health records; machine learning; patient risk stratification.

Clostridium (Clostridioides) difficile infection (CDI) is a prevalent condition [1–3] that often arises in health care settings [4]. As a pathogen, *C. difficile* is genetically diverse [1], with some strains more associated with the development of complicated disease, including polymerase chain reaction (PCR) ribotypes 027, 078/126, 056, and 018 [5, 6]. Clinically, the outcome of CDI varies across patients, ranging from complete recovery to mortality [7]. Currently, treatment decisions (eg, antimicrobial selection, admission to intensive care, and use of adjuvant therapies) are not informed by data-driven models of patient risk for complications, constituting a critical need [8].

Current clinical guidelines recommend treatment with vancomycin (125 mg orally 4 times per day) or fidaxomicin

(200 mg twice daily for 10 days) on initial diagnosis. The recently revised guidelines no longer recommend treatment with metronidazole, but data exist demonstrating that compared with vancomycin this antibiotic may be adequate for mild/moderate disease at a lower cost and potentially lower risk of selecting for antibiotic resistance in clinically important bacteria such as *Enterococcus* species [9–11]. Given the complexity in treating patients with CDI, researchers have sought risk stratification models in support of individualized treatments. To quantify patient risk for developing complicated CDI, previous models have used expert-curated sets of patient characteristics [12–15]. In light of the fact that many of the factors driving patient risk for complications are not well understood, we sought a more comprehensive approach.

We used the structured contents of the EHR and a machine learning approach to develop an intelligible predictive model for complicated CDI. Such a model has the capacity to make a prediction on the day of diagnosis and each day thereafter. Though, as the patient's disease progresses, we expect the task to become easier as it shifts from prediction to recognition. We compared the discriminative performance of this model with a previously published, expert-curated model for the same outcome [13]. In parallel, we investigated patient characteristics associated with increased risk for complications. Insights

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from these analyses may guide the development of interpretable machine learning applications for supporting CDI treatment decisions in real time.

METHODS

Study Population

We considered a cohort of 1144 cases of CDI, as described in Rao et al. [13]. This study population included adult inpatients diagnosed with CDI between October 2010 and January 2013 at the University of Michigan hospitals (UM). We linked each case of CDI from this cohort with structured EHR data available in the Research Data Warehouse at UM. We excluded a small fraction of CDI cases for whom data were not readily available. This study was approved by the Institutional Review Board at the University of Michigan.

Model Outcome

As in Rao et al., we considered a binary prediction task, in which CDI cases that became complicated were labeled 1, and 0 otherwise [13]. Building on previous work, we defined CDI as complicated if it led to any of 3 adverse outcomes within 30 days: admission to intensive care, colectomy, or mortality [12, 13, 16]. All cases were independently labeled through chart review by 2 clinicians, and adjudicated by a third if there was disagreement [13]. In this way, cases were labeled complicated only if the complication(s) were judged to have been caused by the CDI and not by other factors. Using data available before the time of prediction, we sought to learn a model that could accurately sort cases from low to high risk for this composite outcome. To facilitate comparison with the model produced by Rao et al., we evaluated predictions made 2 days after diagnosis. However, recognizing that earlier predictions are more useful, we considered 2 additional prediction times: the day of CDI diagnosis and the day after diagnosis. We hypothesized that prediction will be more difficult near the beginning of a patient's infection course, as fewer clinical indicators for mild or severe disease will have manifested. We identified the day of CDI diagnosis using the time-stamped laboratory test result for presence of toxigenic *C. difficile*.

Data Preprocessing

We extracted EHR data describing each patient admission in our study population from the Research Data Warehouse at UM using the patient's medical record number and date of CDI diagnosis. Specifically, we extracted patient demographics (eg, age), patient history within the past 90 days (eg, diagnosis of diabetes within the past 90 days), admission details (eg, scheduled, urgent, or emergency admission), and daily hospitalization details (eg, prescribed inpatient medications) (Supplementary Table 1). To focus more on patient state than clinician suspicion, we removed variables that clearly encoded clinical suspicion of complicated CDI or could act as a proxy for any elements of

our composite outcome, including those related to administration of intravenous metronidazole, sodium chloride bolus, vancomycin enema, vancomycin-resistant *Enterococcus* culture (VRE), and methicillin-resistant *Staphylococcus aureus* culture (MRSA), as well as assignment to medical critical care units/wards. We removed VRE and MRSA cultures, as they are strong proxies for any-cause admission to the intensive care unit.

For every case, we considered data collected on the day of the prediction, in addition to the 2 days leading up to the prediction, with the goal of capturing recent trends. We represented all the data from this time period as a vector of binary features. These data included time-invariant features such as county of residence and time-varying features such as systolic blood pressure. In the case of categorical data like hospital unit/ward, we mapped this information to binary features indicating whether the patient had been in that hospital location during the 3-day period. For continuous vital signs (eg, respiratory rate), we used expert-defined ranges to map the variable to binary features indicating the presence of any low, normal, or high measurements during the 3-day period. We discretized other continuous variables into quintiles, mapping quintiles to a binary feature (eg, having an age in the first quintile was mapped to a binary feature). Some variables exhibited a significant amount of skew (eg, many patients with 0 previous encounters in the past 90 days). In such cases, we grouped any nonunique quintiles together. EHR data are typically not missing at random. Thus, we included additional features encoding missingness for each variable and did not perform data imputation or case-wise deletion.

In addition to these EHR data, we also considered the manually curated set of variables used in Rao et al. [13]. These data included 1 demographic characteristic (age) and information about the current hospitalization (eg, high-creatinine flag and metastatic cancer comorbidity). The curated feature set did not include information about previous hospitalizations, prior CDI exposure, or patient location within the hospital. However, it did include information about cancer diagnoses more than 90 days before the current admission, unlike the EHR model. For vital signs and laboratory results, each feature corresponded to either the minimum or maximum measurement taken within 48 hours of diagnosis. We processed these features like in the EHR model and concatenated them into a binary feature vector to be used as input to the predictive model. This yielded 2 different data representations (curated vs EHR) that could be used in making predictions.

Model Training and Evaluation

We trained models using the curated data, the EHR data, and a combination of both. To train and evaluate the predictive models, we separated cases into train and held-out test partitions. We split the cases temporally, training the model on the earliest 80% of the data and evaluating on the most recent

20%. Compared with separating the cases randomly into train and held-out partitions, a temporal split better emulates how a predictive model may perform during prospective deployment.

Given the high dimensionality of our data set and the potential to overfit, we used L2 regularization and k-best feature selection (explained below). We selected the regularization hyperparameter C from 10^{-5} , 10^{-4} , ..., 10^5 using 5-fold cross-validation on the training set with 5 temporally defined folds [17]. We used the same process to fit k , a hyperparameter selecting the number of features to keep, after ranking features based on how related they were with the outcome (calculated with Pearson's chi-square statistic). We optimized k over a range that spanned 25–100 with step size 25, 100–1500 with step size 100, and 1500–4000 with step size 500. The final model was trained using the C and k that led to the best average discriminative performance across training set folds. Applied to the held-out test data, we measured the discriminative performance of the model in terms of the area under the receiver operating characteristic curve (AUROC). We also compared the models' sensitivity, specificity, and average precision. We computed empirical 95% confidence intervals using 10 000 bootstrapped samples of the held-out set. We repeated the process above for each feature representation (curated, EHR, and a combination) and each possible prediction time. This resulted in 5 different models. Figure 1 illustrates our model training and evaluation pipeline. All analyses were performed in Python (Python 3.6), and our code is available at https://gitlab.eecs.umich.edu/mld3/complicated_cdi_prediction.

RESULTS

Patient Characteristics

Of the cases in the original cohort from Rao et al. [13], 26 (<2.5%) could not be matched with data in the Research Data Warehouse. After excluding these cases, the final patient

cohort involved 1118 cases of CDI, 89 (8%) of which were complicated CDI. The median length of stay was 9 days, and median time of CDI diagnosis was on the third day of hospitalization (Table 1).

Model Training

Models were trained on a training set of 894 cases and tested on a held-out set of 224 cases. As in Rao et al., the curated model used 23 features [13]. We extracted 4271 features from the EHR. After feature selection, the EHR models for making a prediction at diagnosis, 1 day after, and 2 days after, retained 3000, 800, and 900 features, respectively. Regarding regularization strength, the EHR models used C values of 10^{-3} , 10^{-4} , and 10^{-3} , respectively.

Model Performance

Two days after CDI diagnosis, the EHR model resulted in better discriminative performance compared with the curated model (0.90; 95% CI, 0.83–0.95; vs 0.84; 95% CI, 0.75–0.92). The EHR model also had greater average precision (AUROC, 0.30; 95% CI, 0.15–0.60; vs AUROC, 0.25; 95% CI, 0.09–0.47) (Supplementary Figure 1). At a threshold based on the 95th percentile of risk for each model, the EHR model had greater specificity (96.7% vs 95.3%) and sensitivity (41.7% vs 16.7%) (Figure 2). Combining the curated features with the EHR features did not improve discriminative performance 2 days after diagnosis (Table 2). Model performance decreased when tasked with making predictions earlier: AUROC, 0.79; 95% CI, 0.67–0.90 1 day after diagnosis and AUROC, 0.69; 95% CI, 0.55–0.83 at diagnosis (Figure 3). Removing several variables clearly encoding clinical suspicion of complicated CDI did not substantially change model performance at any prediction time point (Supplementary Figure 2). Examining the learned coefficients, on the day of CDI diagnosis, high respiratory rate and obtaining an adult blood culture were strongly associated

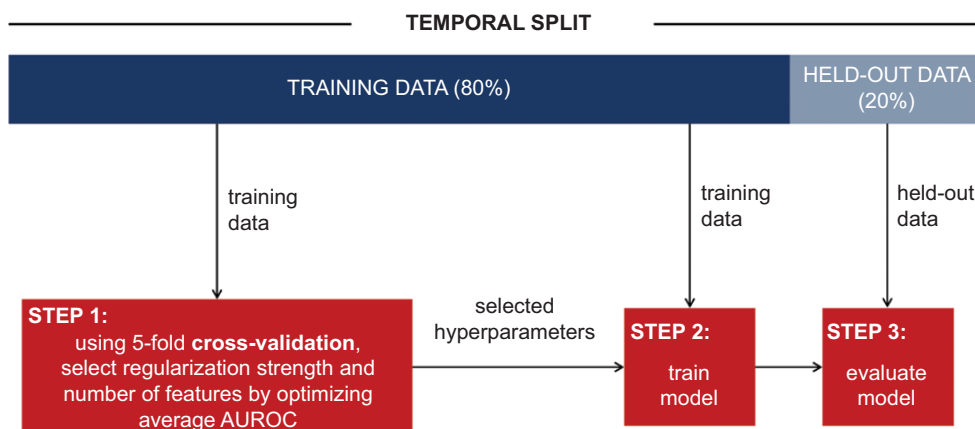


Figure 1. Predictive model training and evaluation flowchart. We split the data 80%/20% temporally, performed 5-fold cross-validation on the training data to select hyperparameters (step 1), and then, using these hyperparameters and all of the training data, learned a model (step 2). The model was evaluated on a held-out set of data (not used in selecting the model). Abbreviation: AUROC, area under the receiver operating characteristic curve.

Table 1. Selected Patient Characteristics of Our Study Cohort

Characteristic	Median (IQR) or No. (%)
No. of CDI cases	1118
No. of patients	966
Age, y	59 (46–69)
Female gender	611 (54.7)
LOS, d	9 (5–17)
Day of CDI diagnosis	3 (2–7)
Charlson-Deyo score	2 (0–3)
Inflammatory bowel disease diagnosed in the past 90 d ^a	20 (1.8)
Solid organ transplant	179 (16.0)
Concurrent non-CDI antimicrobial use	745 (66.6)
Fluoroquinolone use from admission to diagnosis	378 (33.8)
Proton pump inhibitor use	775 (69.3)
Prior CDI within the past year	262 (23.4)
Prior CDI within the past 90 d	182 (16.3)
Failed initial CDI therapy within the past 14 d	22 (2.0)
BMI, kg/m ² ; missing No. (%)	26.3 (22.6–31.6); 131 (11.7)

^aICD-9-CM codes 555 or 556.

Abbreviations: BMI, body mass index; CDI, *Clostridium difficile* infection; IQR, interquartile range; LOS, length of stay.

with development of complicated CDI (Table 3). Two days later, the factors most associated with risk included high and low respiratory rate, low systolic blood pressure, and low blood CO₂. At both time points, normal respiratory rate and young age were associated with protection.

DISCUSSION

Automated patient risk stratification techniques that leverage EHR data have the potential to provide critical support for clinical decision-making. In contrast to risk estimates that depend on manual chart review, risk estimates derived from the EHR can be automatically generated and updated throughout a patient's hospitalization. In the context of CDI, predictive models have been developed for identifying patients at high risk for infection, recurrence, and complications [18–20]. We have previously shown how a model that leverages the structured contents of the EHR can outperform a curated model in predicting CDI, and we have demonstrated how these techniques can generalize across hospitals with different patient populations and different EHR systems [21, 22]. In the context of recurrent CDI, Escobar et al. compared machine learning and curated risk stratification models, concluding that neither approach could reliably predict first recurrence [18]. In this work, we focus on estimating patient risk for a complicated course of CDI.

The task of predicting complicated CDI is difficult, and current treatment guidelines do not incorporate data-driven estimates of risk. To this end, we sought to develop a systematic way to improve our ability to identify complicated cases by considering all structured data available in the EHR. This work represents a first step in that direction. We demonstrate that, using a machine learning approach, it is possible to accurately risk-stratify patients for complications early on in the course of disease.

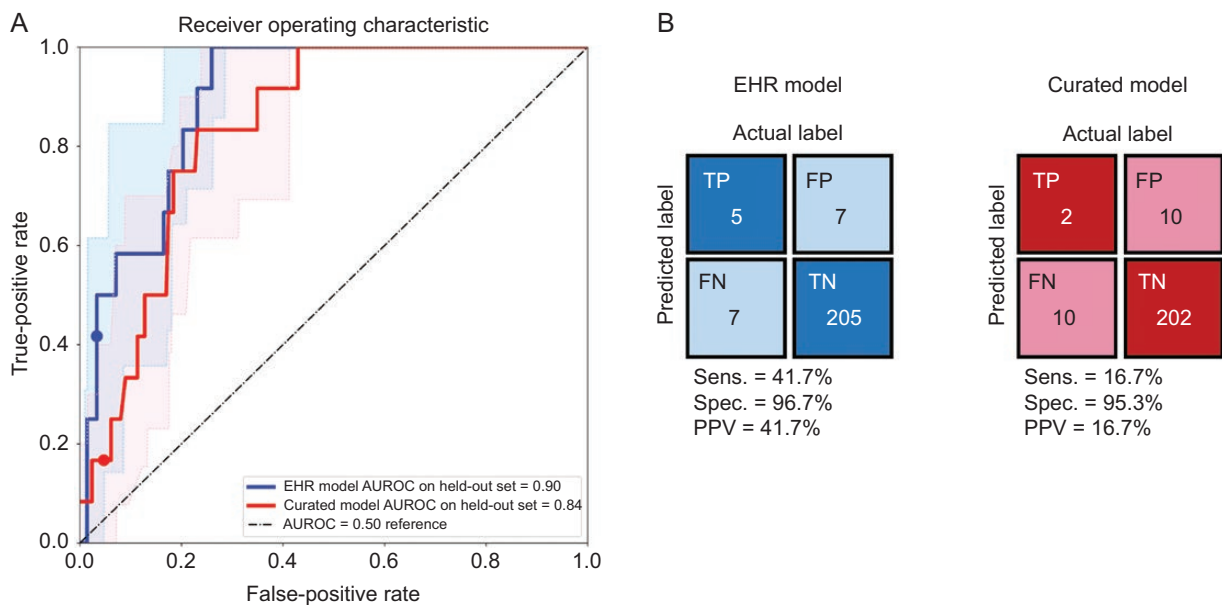


Figure 2. Discriminative performance (A) and confusion matrices (B) of predictive models for a complicated course of CDI 2 days after diagnosis using either electronic health record (EHR)-based or curated features. Shaded regions in (A) represent empirical 95% confidence intervals. The EHR model dominates the curated model for nearly all values of specificity (1-FPR). At a threshold based on the 95th percentile of risk for each model (marked with dots), the EHR-based model yields better sensitivity, specificity, and positive predictive value. Abbreviations: CDI, *Clostridium difficile* infection; FN, false negative; FP, false positive; FPR, false-positive rate; PPV, positive predictive value; TN, true negative; TP, true positive.

Table 2. Comparison of EHR, Curated, and Merged Models for Complicated CDI 2 Days After Diagnosis

Model	No. of Features	AUROC on Held-Out Set (95% CI)
Curated	23	0.84 (0.75–0.92)
EHR	900	0.90 (0.83–0.95)
EHR + curated	923	0.88 (0.81–0.95)

Abbreviations: AUROC, area under the receiver operating characteristic curve; CDI, *Clostridium difficile* infection; CI, confidence interval; EHR, electronic health record.

The development and validation of EHR-based risk stratification models for predicting complicated CDI could eventually help clinicians tailor treatments to individuals. On the day of CDI diagnosis, a patient’s estimated risk for complications could serve as an adjunct, easily obtainable resource for clinical decision support. Treatment decisions such as whether to use high-dose vancomycin or perform a loop ileostomy with antegrade vancomycin infusions [23] often do not occur until complicated CDI has already set in. In severe cases, early aggressive therapy can positively impact the course. However, invasive treatments such as enemas (fecal microbiota transplantation or vancomycin) and surgery are optimally used in only select patients, and such decisions lack the rigorous guidelines associated with initial treatment.

In addition to potentially guiding more drastic and invasive treatment, accurate risk prediction for complicated CDI could guide initial antibiotic therapy. The new US guidelines on treating CDI promote vancomycin or fidaxomicin over metronidazole [24]. In the inpatient setting, this shifts nearly all cases to vancomycin therapy due to the expense of fidaxomicin. This, in turn, could increase VRE selection pressure and disrupt the microbiome, leading to subsequent infections [25]. As even

more narrow-spectrum CDI treatments in the pipeline become available [26], a model that could enable targeted vancomycin or fidaxomicin use in a cost-effective way could help alleviate these potential adverse consequences.

Though identifying effective interventions that prevent or mitigate complicated CDI would require extensive additional studies, tools like the one developed here play an important role. The statistical power of cohort studies and randomized controlled trials (RCTs) is often limited when the primary outcome is infrequent. Risk stratification tools can help target patients with a higher risk of adverse outcomes and increase the feasibility of such important trials.

Based on thousands of variables, the EHR-based model provided better risk estimates than one that relied on a curated set alone. In this study, we sought to test the feasibility of a machine learning approach to generate early and accurate predictions of a patient’s risk for complicated CDI; reassuringly, the model features with highest weight align with those a clinician would identify. Even if such models do not identify new risk factors, they consider a much larger set of patient characteristics than any 1 clinician can consider simultaneously. As the amount of data collected in hospitals continues to increase, it will be important to equip clinicians, who are already faced with many competing priorities, with the tools necessary to identify patterns and distill the data into actionable knowledge. For example, operating in the background, a model for predicting complications could alert the care team if a patient’s risk increases unexpectedly. In parallel, if many patients in the same hospital unit are identified as being at heightened risk, this could inform hospital unit/ward cleaning practices or strategic transferring of patients to private rooms.

These results should be interpreted in the context of several limitations. First, given that these are merely associations,

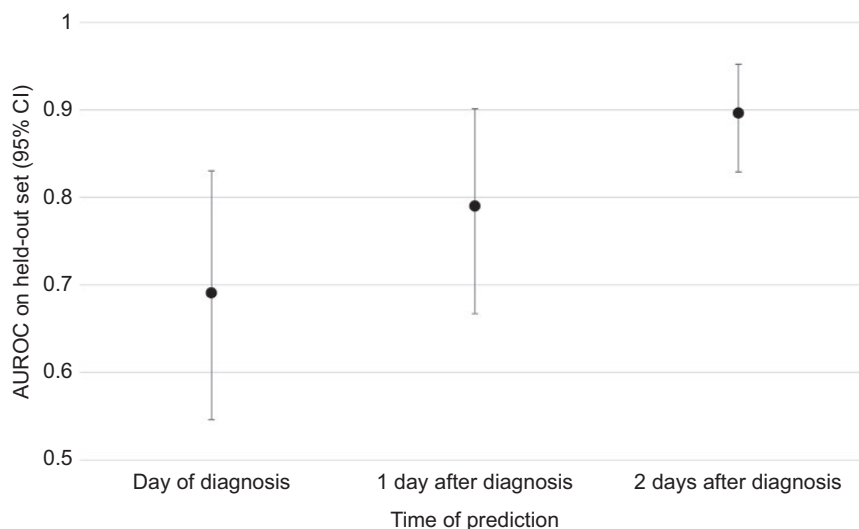


Figure 3. Discriminative performance of electronic health record–based complicated CDI model across time. As more data become available and are integrated into the model, the model achieves better discriminative performance. Error bars represent empirical 95% confidence intervals. Abbreviations: AUROC, area under the receiver operating characteristic curve; CDI, *Clostridium difficile* infection; CI, confidence interval.

Table 3. Predictive Features for Complicated CDI at Diagnosis and 2 Days After

Day of Diagnosis		2 Days After Diagnosis	
Rank	Associated With Higher Estimated Risk	Rank	Associated With Higher Estimated Risk
1	Obtaining adult blood culture	1	High respiratory rate (>20 breaths/min)
2	High respiratory rate (>20 breaths/min)	2	Low respiratory rate (<12 breaths/min)
3	Low blood CO ₂ (<23 mmol/L; Q 1/5)	3	Low systolic BP (<90 mmHg)
Rank	Associated With Lower Estimated Risk	Rank	Associated With Lower Estimated Risk
1	Normal respiratory rate (12–20 breaths/min)	1	Missing PTT measurement
2	Low RDW (<14.6%; Q 1/5)	2	Normal respiratory rate (12–20 breaths/min)
3	Age (<41 years old; Q 1/5)	3	Missing phosphorous level measurement

Please see [Supplementary Table 2](#) for the model coefficients associated with these features.

Abbreviations: BP, blood pressure; CDI, *Clostridium difficile* infection; PTT, partial thromboplastin time; RDW, red cell distribution width.

additional investigation is required to establish the direction of the true underlying relationships (eg, through RCTs). Moreover, despite the removal of many variables that could serve as proxies for our composite outcome, some model features may not be true risk factors but rather markers for the beginning of complicated CDI itself. This may be elucidated by tracking the evolution of a patient's risk over multiple days of their hospitalization; automated EHR models enable these kinds of future analyses. Second, our analysis is retrospective. As a following step, prospective studies are necessary for understanding how such models perform in real time. Third, the results are based on a small sample size from a single institution. Though this has implications regarding the generalizability of the model, it does not diminish the generalizability of the approach. As patient populations, clinical protocols, and in turn risk factors can vary across institutions, we encourage researchers to focus on tailoring models to the populations in which the model will ultimately be deployed [22]. On a related note, the definition of a complicated course of CDI is not universal. We used the CDC surveillance definition [16], the same definition as in Rao et al., which includes a stipulation that the specific adverse outcomes be attributable to CDI (requiring clinical review). Future work should investigate the appropriateness of definitions of complicated CDI derived solely from the EHR. Fourth, we cannot exclude the possibility that patients may experience the outcome at another hospital, and thus we may have potentially underestimated the extent of complications from CDI. In addition, although CDI testing was recommended only for symptomatic patients during our study period and this was further validated by chart review, some positive CDI tests might reflect asymptomatic carriers. We hypothesize that our model learns, in part, to differentiate asymptomatic carriers from those who will experience complicated disease. Finally, beyond the structured content, the EHR contains data from pathogen genomic sequencing, free-text clinical notes, and/or wearable technology that could provide a more complete picture of a patient's clinical state. For example, the Xpert *C. difficile* diagnostic test has a gene target that can report whether the strain is NAP1 or non-NAP1. In the future, integrating both biological and administrative

data like these has potential for improving models of disease progression.

In summary, we demonstrated how a machine learning approach could be used to learn an EHR-based predictive model for accurately estimating a patient's risk of developing complicated CDI. Our approach leverages thousands of variables that can be readily extracted from the EHR. This approach has many potential applications, including guiding future clinical studies. Prospective evaluation and deployment of models such as these offer important opportunities to aid clinicians in real time and tailor patient therapy.

Supplementary Data

Supplementary materials are available at *Open Forum Infectious Diseases* online. Consisting of data provided by the authors to benefit the reader, the posted materials are not copyedited and are the sole responsibility of the authors, so questions or comments should be addressed to the corresponding author.

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