Validation of Electronic Health Record Detection of Patient Safety Outcomes

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ABSTRACT

Background
Adverse events are common for hospitalized Canadians and lead to worse patient outcomes. We aimed to validate the use of our electronic health record (EHR) to monitor important patient safety outcomes.

Methods
EHR data were abstracted for four high-priority safety outcomes: venous thromboembolism (VTE), hypoglycemia, Clostridium difficile (C. difficile) infection, and prolonged nil per os (NPO) orders. A manual chart review was performed to determine the sensitivity and specificity of the EHR for each patient safety outcome.

Results
The sensitivity and specificity were: 94.3% and 99.2% for C. difficile infections, 34.3% and 88.0% for VTE, 96.9% and 96.3% hypoglycemia, 61.8% and 98.5% for prolonged NPO status.

Conclusion
The EHR is reasonably sensitive and specific in monitoring rates of hypoglycemia, C. difficile infection, and prolonged NPO in medical inpatients. Importantly, validation of EHR data with manual chart review is necessary before using this data to monitor patient safety outcomes.

RESUME

Contexte général
Les événements indésirables sont fréquents chez les Canadiens hospitalisés et entraînent une détérioration de l'état de santé des patients. Nous voulons valider l'utilisation de notre dossier de santé électronique (DSE) pour surveiller les résultats importants en matière de sécurité des patients.
Méthodes
Les données du DSE ont été résumées pour quatre résultats prioritaires en matière d’innocuité : thromboembolie veineuse (TEV), hypoglycémie, infection à Clostridium difficile (C. difficile) et ordonnance prolongée de néant par os (OBNL). Un examen manuel des dossiers a été effectué pour déterminer la sensibilité et la spécificité du DSE pour chaque résultat lié à la sécurité des patients.

Résultats
La sensibilité et la spécificité étaient : 94,3 % et 99,2 % pour les infections à C. difficile, 34,3 % et 88,0 % pour la TEV, 96,9 % et 96,3 % pour l’hypoglycémie, 61,8 % et 98,5 % pour le statut d’ANP prolongé.

Conclusion
Le DSE est raisonnablement sensible et spécifique dans la surveillance des taux d’hypoglycémie, d’infection à C. difficile et d’OBNL prolongé chez les patients hospitalisés en médecine. Il est important de noter que la validation des données du DSE au moyen d’un examen manuel des dossiers est nécessaire avant d’utiliser ces données pour surveiller les résultats en matière de sécurité des patients.

Adverse events are complications caused by healthcare, as opposed to the underlying disease process, that lead to worse outcomes. One in every 18 hospitalized Canadians experience an adverse event, leading to 500,000 additional hospital days per year and $685 million in costs. Importantly, patients who suffer an adverse event have higher mortality.

The Canadian Institute for Health Information (CIHI) calls for healthcare organizations to make the monitoring of patient safety a priority; however, patient safety outcomes, which include adverse events, near misses, and nonadherence to best practices (“omissions”), are notoriously difficult to measure. The gold standard is a chart review, which is too time-consuming and resource-intensive for quality improvement cycles. Staff-reporting systems are common but undercount adverse events. Risk-adjusted hospital mortality rates or reviews of administrative discharge codes are easily obtainable but too crude for quality improvement initiatives. The ideal patient safety reporting system generates data in real-time, requires minimal effort from healthcare workers to be sustainable and accurately reflects meaningful events.

The electronic health record (EHR) allows for the collection of timely data on patient safety outcomes but may lack contextual data about appropriate omissions. We sought to validate EHR detection of patient safety outcomes using chart audit as the reference standard.

Methods
Ethics approval was received from the University of Calgary Conjoint Health Research Ethics Board. The study was funded by the Canadian Institute for Health Information (CIHI) and the University of Calgary Conjoint Health Research Ethics Board. The study was conducted in compliance with the Declaration of Helsinki and the Canadian Tri-Council Policy Statement on Research Ethics.

Table 1. Definitions of Patient Safety Outcomes

| Term       | Definition                                                                                       | Example                                      |
|------------|--------------------------------------------------------------------------------------------------|----------------------------------------------|
| Adverse event | Complications caused by healthcare management as opposed to the disease itself that results in worse outcomes, including prolonged hospitalization, disability or death. Not all adverse events are errors. | C. difficile infection after antibiotic use. |
| Errors     | An action or omission that has the potential to cause an undesired outcome. Most errors do not lead to adverse events. | Antibiotics are ordered but is not administered to patient. |
| Near miss  | An error that has the potential to result in a poor patient outcome but does not reach the patient or result in harm. | Incorrect medication administered to the patient without resulting in harm. |
| Omissions  | Failure to use an evidence-based intervention intended to reduce adverse events. | Pharmacologic or mechanical venous thromboembolism prophylaxis not ordered for high-risk inpatients. |

C. difficile = Clostridium difficile
Robert and Wachter, 2018.
Table 2. Composite Patient Safety Outcomes and Corresponding Data Definitions

| Patient Safety Outcome | Category | EHR Data Signal |
|------------------------|----------|-----------------|
| C. difficile           | NM       | Order for a C. difficile PCR test |
|                        | AE       | Positive C. difficile PCR test |
| Venous thromboembolism | O        | Missing order for VTE prophylaxis at 48 hours. VTE prophylaxis included VKA order, DOAC order, or SCD order. Patients with an INR > 3.0 or platelet count < 50 x 10^9/L were excluded. |
|                        | NM       | An omission plus a negative radiographic study for VTE* performed 48 hours after admission time. |
|                        | AE       | An omission plus a positive radiographic study for VTE* performed 48 hours after admission time. |
| Hypoglycemia           | AE       | Capillary glucose < 3 mmol/L Serum glucose < 3 mmol/L Activation of “Hypoglycemia Order Set” |
| Prolonged NPO order    | NM       | NPO order for or longer than 48 hours |

AE = adverse event; C. difficile = Clostridium difficile; EHR = electronic health record; INR = international normalized ratio; NM = near miss; NPO = nil per os; O = omission; SCD = sequential compression devices; VTE = venous thromboembolism; VKA = vitamin k antagonist.

*Radiographic study for VTE includes: V/Q scanning, CT Pulmonary Embolism or extremity doppler ultrasound.

Patient Safety Outcome Definitions

Four adverse events were chosen from CIHI based on their prevalence, relevance to internal medicine, and feasibility of measurement. The study team identified relevant contributing errors (near misses and omissions); the near misses, omissions and adverse events were combined into a composite measure that was called “patient safety outcomes.” This was done due to the small numbers of adverse events, the importance of detecting errors even when harm does not occur, and the feasibility of measuring certain events using the EHR. Errors are actions or omissions that have the potential to cause an undesirable outcome; not all errors lead to adverse events and not all adverse events are caused by errors. Near misses and omissions can be seen as process measures that may lead to an adverse clinical outcome for a patient.

A clinical data analyst with expertise in our EHR created data definitions based on the patient safety outcomes in consultation with two clinicians (SR and RK). The data definitions were used to identify patient safety outcomes during an inpatient admission. Table 2 outlines the definitions used to extract patient safety outcomes from the EHR.

i. Clostridium Difficile
   No identified relevant omissions. Near miss was defined as a negative Clostridium difficile stool PCR test. In our center, this test can only be sent in patients having diarrhea who have risk factors including previous C. difficile infection or recent antibiotic exposure. While testing was pending, the patient was placed on contact precaution and moved into an isolation room, representing a burden on patients, healthcare staff, and the hospital system itself. However, this did not demonstrate the harm that would need to be disclosed to the patient. An adverse event was a positive test not otherwise captured in the near-miss definition. Any patients with positive testing done prior to or within 48hrs of admission were excluded.

ii. Hypoglycemia
    No identified relevant omissions or near misses. An adverse event was defined as any day with a recorded capillary or serum glucose less than 3 mmol/L or activation of the EHR Hypoglycemia Order Set.

iii. Prolonged NPO Order
    Adverse events related to prolonged nil per os (NPO) include decreased patient satisfaction, delirium, and electrolyte...
abnormalities. We included patients who did not have diet orders or had NPO entered in the EHR for greater than 48 hours. Forty-eight hours was selected to exclude patients who may have had appropriate, temporary NPO orders while awaiting procedures that were delayed for one day.

iv. Venous Thromboembolism

Omissions were patients without an order for VTE prophylaxis at 48 hours from their admission time. Near misses included patients without an order for VTE prophylaxis at 48 hours into their admission who underwent relevant radiographic testing more than 48 hours after admission which was negative. Patients not on appropriate VTE prophylaxis after 48 hours of admission with positive radiographic results for VTE were considered adverse events. All patients that underwent radiographic testing within 48 hours of admission or had the admission diagnosis of VTE were excluded.

Data Abstraction

EHR data were abstracted for patients admitted to Internal Medicine teaching services in Calgary, Canada. No patients were excluded. The teaching service teams were led by an Internal Medicine specialist and included a combination of learners ranging from medical students to General Internal Medicine fellows.

Using an estimated prevalence of 20%, we determined that 156 patients were required to achieve a minimum power of 80% for detecting a change in the percentage value of sensitivity from 0.70 to 0.90 based on a target significance level of 0.05. This sample size was powered to detect an increase in specificity from 0.80 to 0.90. The reviewer received a list of patients randomly selected from each patient safety outcome group and included control patients who did not have a patient safety event. Patient safety outcomes were identified from the paper charts using the same definitions as used to collect the EHR event rates. The paper chart auditor (CH) was masked to the EHR data. These included 1,492 C. difficile outcomes (1,307 near misses and 185 adverse events), 1,657 patients with a VTE safety outcome (1,609 omissions, 38 near misses and 10 adverse events), 809 hypoglycemia outcomes, and 545 patients who were NPO for more than 48 hours. A full chart review was performed on 169 randomly selected patients by a study team member (CH) who was masked to the EHR data (Figure 1).

The chart review included: 77 control patients, 31 who had a hypoglycemic outcome, 34 with a C. difficile outcome, 28 patients with a VTE outcome, and 23 patients with an NPO outcome. The sensitivity and specificity of the EHR data is shown in Table 4. The EHR detection of C. difficile outcomes had a sensitivity of 94.3% and specificity of 99.2%. VTE events were detected with a sensitivity of 34.3% and specificity of 88.0%, and hypoglycemia had a sensitivity of 96.9%, the specificity of 96.3%. EHR detection of prolonged NPO status had a sensitivity of 61.8%, the specificity of 98.5%.

Discussion

Adverse events occur in 5.6% of the over 2,300,000 annual hospitalizations in Canada, and up to 40% may be preventable. Evaluation of patient safety interventions is limited by challenges in measuring errors, near misses, and adverse events. EHRs could

| Characteristic       | All patients (n=7,527) | Controls (n=3,733) | Patient Safety Outcome (n=3,794) | p-value |
|----------------------|-----------------------|-------------------|---------------------------------|---------|
| Male (%)             | 4,168 (55.4)          | 2,029 (54.3)      | 2,139 (56.4)                    | 0.08    |
| Mean age (years, SD) | 59 (18.6)             | 59 (18.9)         | 60 (18.3)                       | 0.02    |
| Median length of stay (days, IQR) | 5.2 (2.8–10.8) | 4.7 (2.5–8.7) | 6.1 (3.0–13.8) | <0.01 |

IQR=interquartile range; SD = standard deviation.
be an important tool in patient safety reporting because EHRs may overcome the time delays and resource intensity of other methods. Previous studies have validated EHR for monitoring quality indicators with sensitivities ranging from 34 to 98.5% and specificity ranging from 88 to 99%.

Fortunately, individual types of adverse events, such as in-hospital VTE, are rare; however, this limits assessment of the impact of a quality improvement intervention, especially when attempting rapid Plan-Do-Study-Act (PDSA) cycles, because multiple events must be accrued to demonstrate a change. This can be overcome by combining adverse events with other types of related errors, such as near misses and omissions, that occur more frequently and are part of the critical path leading to an adverse event.

Our validation study demonstrates that rates of patient safety outcomes including hypoglycemia, *C. difficile* infection, and prolonged NPO status can be measured using the EHR.
with reasonable sensitivity and specificity. Importantly, our study highlights that EHR data must be validated prior to use because the face validity of EHR measures is not consistently demonstrated by chart audit.

Studies using ICD-9 and ICD-10 codes to detect *C. difficile* infections demonstrate sensitivities of 76 to 81% and specificities of 99.9%.

Our EHR demonstrated similar sensitivity and specificity, 94.3% and 99.2% respectively, for the *C. difficile* patient safety outcome. The chart audit revealed that the EHR did not capture *C. difficile* testing orders entered during patient transfers; for example, patients without an order for testing had a test result reported, and this occurred after a patient had moved locations. It is not clear why this occurs with our EHR.

Eleven percent of patients in our study suffered a hypoglycemic event. The EHR sensitivity of 96.9% and specificity of 96.3% for hypoglycemia is similar to ICD-9 coding, which demonstrates a sensitivity of 88% and specificity >99% compared to chart review.

Our EHR data was not 100% sensitive for hypoglycemia because some nurses did not enter low glucose measurements into the trackable “Vital Signs” portion of the EHR and were instead documenting the episode in a free-text “Nursing Note”. These episodes were treated with juice which does not activate the Hypoglycemia Order Set. This finding highlights the importance of validating EHR data definitions before using the data collected in improvement efforts; the end-users of our EHR were not using the EHR as expected and therefore our data definitions missed these outcomes.

**Table 4. The Sensitivity and Specificity of Patient Safety Outcomes Detected By Electronic Health Record Data Compared to Chart Audit At Our Institution for Patient Admitted to Teaching Units between January 1, 2016 And May 3, 2017.**

| Patient Safety Outcome | Sensitivity | Specificity | Percent Agreement* |
|------------------------|-------------|-------------|---------------------|
| *C. difficile* infection | 94.3%       | 99.2%       | 98.2%               |
| Venous thromboembolism  | 34.3%       | 88.0%       | 76.8%               |
| Hypoglycemia           | 96.9%       | 96.3%       | 96.4%               |
| Prolonged NPO order    | 61.8%       | 98.5%       | 91.1%               |

*C. difficile = Clostridium difficile, NPO = nil per os.*

*Between the two methods (Manual chart review vs. EHR data extraction)*

Our study demonstrates that EHR data is adequately sensitive and specific in certain patient safety outcomes to be used in rapid-cycling quality improvement work. The mismatch between EHR data definitions and chart audit for patient safety outcomes with high face validity, such as hypoglycemia, occurred when EHR users deviated from expected processes. More accurate data could be captured with additional parameters being set on data definitions. This should be done in consultation with a local hospital IT expert for the EHR and medical clinicians. There should be an extensive review of definitions prior to utilization for data extraction. This will often require multiple rounds of refinement to find the optimal data definitions which will be health centre specific. Further, all data definitions used should be validated with a select chart review to ensure that the data being collected is what is intended.

An important limitation of our study is that EHR data does not provide a context for whether a patient safety outcome is an error or if it was appropriate care. For example, a patient may be NPO for a prolonged duration due to a bowel obstruction. Though this prolonged period of NPO may still lead to an adverse event such as delirium or an electrolyte abnormality, it was not an error but a side effect of appropriate care. The trade-off between this level of detail and the feasibility of measurement is a key issue in the collection of patient safety data. One method that can account for this trade-off between context and resources required for data collection is benchmarking; a quality improvement team accepts that the rate of patient safety outcomes may never be expected to be zero due to acceptable indications. However, they aim to reduce the rate of outcomes to an accepted benchmark or track for unexpected changes in patient safety outcome rates using temporal data collection.

This study was meant to bring awareness to the possibility of using EHR data for quality improvement studies in the clinical setting. As the sophistication of EHRs continue to improve, so will the sensitivity and specificity of the data they can track.
Conclusion

Using EHR data to measure adverse event and error rates has advantages over other methods such as chart audit because it requires fewer human resources and can report results more quickly. These characteristics are important in rapid PDSA cycles used in quality improvement. However, these advantages come at the cost of potentially less accurate information. Our study demonstrates that EHR event definitions with face validity, even when developed with clinicians and clinical data specialists, may not capture events as intended. Validation of EHR detection of patient safety outcomes is a necessary step prior to using this data in future quality improvement work.

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