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Mining clinical text for signals of adverse drug-drug interactions

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ABSTRACT

Background and objective Electronic health records (EHRs) are increasingly being used to complement the FDA Adverse Event Reporting System (FAERS) and to enable active pharmacovigilance. Over 30% of all adverse drug reactions are caused by drug–drug interactions (DDIs) and result in significant morbidity every year, making their early identification vital. We present an approach for identifying DDI signals directly from the textual portion of EHRs.

Methods We recognize mentions of drug and event concepts from over 50 million clinical notes from two sites to create a timeline of concept mentions for each patient. We then use adjusted disproportionality ratios to identify significant *drug–drug–event* associations among 1165 drugs and 14 adverse events. To validate our results, we evaluate our performance on a gold standard of 1698 DDIs curated from existing knowledge bases, as well as with signaling DDI associations directly from FAERS using established methods.

Results Our method achieves good performance, as measured by our gold standard (area under the receiver operator characteristic (ROC) curve >80%), on two independent EHR datasets and the performance is comparable to that of signaling DDIs from FAERS. We demonstrate the utility of our method for early detection of DDIs and for identifying alternatives for risky drug combinations. Finally, we publish a first of its kind database of population event rates among patients on drug combinations based on an EHR corpus.

Conclusions It is feasible to identify DDI signals and estimate the rate of adverse events among patients on drug combinations, directly from clinical text; this could have utility in prioritizing drug interaction surveillance as well as in clinical decision support.

BACKGROUND AND SIGNIFICANCE

Adverse drug reactions result in more than 100 000 deaths annually,¹ with an associated yearly cost of over \$136 billion.² Simultaneously, there is a rise in polypharmacy, which is the use of multiple concomitant drugs to treat medical conditions; one study estimated that 29.4% of elderly patients³ are taking six or more drugs. Drug–drug interactions (DDIs) that lead to adverse reactions are potentially avoidable, if detected early, given that DDIs account for more than 30% of all adverse drug reactions.^{4–6}

New drugs are tested for interactions with existing drugs before market approval using *in vivo* and *in vitro* methods.⁷ However, owing to the sheer number of ways in which drugs can interact,⁸ it is infeasible to test for every kind of interaction. Many DDIs manifest after a certain period of exposure and it takes several exposures for rare

DDIs to occur.⁹ Therefore, postmarketing surveillance is necessary to detect unanticipated interactions that occur when the drug is in use in the general population. The US Food and Drug Administration (FDA) enables such surveillance via spontaneous reporting systems (SRSs) such as the FDA Adverse Event Reporting System (FAERS), and a similar role is served internationally by the World Health Organization's Vigibase. Several studies^{10–13} have successfully inferred DDIs from these sources, overcoming problems of reporting biases¹⁴ and duplicate reporting.¹⁵

Electronic health records (EHRs) are a source of observational data that can complement SRSs and offer the potential for active surveillance.¹⁶ Initiatives like the Observational Medical Outcomes Partnership (OMOP) in the USA and the Exploring and Understanding Adverse Drug Reactions (EU-ADR) project in Europe are focusing on building EHR based surveillance systems. Similarly, the Mini-Sentinel Pilot Program,¹⁷ which is part of the FDA's Sentinel Initiative, uses data of more than 125 million patients across the USA from over 17 data partners for active monitoring of medical products. These projects mainly utilize the structured diagnosis and prescription data from EHRs for identifying single drug adverse reactions. Most efforts aimed at finding DDIs use reported sources for signal detection and use EHRs as a means of validation. For instance, Tatonetti *et al*¹⁰ found 171 new DDIs from FAERS and used the EHRs at Stanford to validate them. Another study by Duke *et al*¹⁸ mined MEDLINE abstracts for hypotheses generation and validated the hypotheses on EHRs.

In addition to structured data, EHRs contain rich information in the unstructured notes taken by doctors, nurses, and other practitioners. By ignoring the unstructured text, we could be missing a substantial portion of adverse events.¹⁹ Many studies^{19–20} have shown that coded information such as the ICD-9 is inadequate to accurately build patient cohorts and there is a considerable advantage in using the unstructured text of EHRs.^{21–22} We argue that such an advantage would also extend to drug safety signal detection. Indeed, there has already been some work^{23–26} demonstrating the discovery of the adverse event profiles for single drugs using unstructured notes.

Therefore, given increasing adoption and access to medical records for research, we expect efforts to shift toward directly mining EHRs with an increased attention to the use of unstructured data for generating hypotheses about drug safety.²⁷ In this paper, we apply data mining methods on the textual portion of EHRs to signal adverse DDIs. To

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our knowledge, this is the first study of its kind. We validate our methods using a gold standard built from existing knowledge bases of DDIs, by applying our methods to two independent large EHR datasets, and by comparing with signaling DDI associations directly from FAERS using established methods. We demonstrate the utility of our method for early detection of new interactions and for identifying alternatives to risky drug combinations. Finally, we publish a database of the rate of adverse events among patients on all combinations of drugs used in our study, based on the EHRs.

MATERIALS AND METHODS

Data sources

Electronic health records

We use the Stanford Translational Research Integrated Database Environment (STRIDE) dataset comprising 9 million unstructured clinical notes corresponding to 1 million patients as our primary source of EHR data. These textual notes span a period of 18 years (1994–2011) and include both inpatient and outpatient notes, that are a combination of radiology, pathology, and transcription reports.

For validation of our results, we use a similar dataset from the Palo Alto Medical Foundation (PAMF), comprising over 50 million outpatient notes corresponding to 1.2 million patients for encounters during 2000–2012. The dataset includes progress, problem visit, procedure, and H&P (history and physical) notes, as well as communication transcripts.

Structured data sources

To compare the accuracy of DDI signal detection from EHRs with SRSs, we use over 4 million FAERS reports covering the period from 1997 through 2012 Q1. These reports are preprocessed to remove duplicate reports, correct terminological errors, and normalize drugs. Medical Dictionary for Regulatory Activities (MedDRA) V14.1 codes are used in FAERS to code adverse events.

We use the Anatomical Therapeutic Chemical Classification (ATC) and 18 ontologies from the Unified Medical Language System (UMLS) Metathesaurus and BioPortal²⁸ (19 ontologies in total; see online supplementary materials S1) for building a lexicon and for normalizing drugs and diseases. Additionally, we use DrugBank,²⁹ Medi-Span Drug Therapy Monitoring System (Wolters Kluwer Health, Indianapolis, Indiana, USA), Drugs.com,³⁰ the National Drug File–Reference Terminology (NDF-RT), and Side Effect Resource (SIDER)³¹ as sources of drug indications and known DDIs.

Annotation of clinical text

We use sets of terms derived from 19 biomedical ontologies to define drug and event concepts (see online supplementary materials S2) as described in our previous work.^{26–32} For drugs, we include trade names and other forms of the drug from RxNorm, but ignore the dosage of the drug. For adverse events, we identify a SNOMED CT concept most similar to the event and systematically including synonyms and hyponyms via the *is-a* relationship graph. To improve the precision of recognizing drug and event concepts, based on prior work on identifying and removing non-informative terms,^{33–34} we remove terms that occur in common English usage from the automatically compiled term-sets.³⁵ Finally, using frequency based sorting,^{36–37} we manually identify ambiguous terms that belong to more than one semantic group (drug, disease, device, procedure),^{36–38} and we suppress their least likely interpretation. For example, ‘clip’ is more likely to be a device than a drug in clinical text, so we

suppress its interpretation as ‘corticotropin-like intermediate lobe peptide’.

We then use a fast text annotator to process clinical notes for mentions of these concepts and order the mentions by the note’s timestamp, thus forming a concept timeline for every patient (figure 1). The annotator also takes into account negation and family history contextual cues to restrict concepts to positive mentions referring to the patient. For example, if the clinical note mentions ‘no evidence of active colitis’ or ‘family history of cancer’, then the note is not tagged with colitis or cancer. We follow a similar approach for the PAMF dataset. We achieve 74% sensitivity and 96% specificity in recognizing diseases on a gold standard from the 2008 i2b2 obesity challenge³⁹ and our performance varies by condition (see online supplementary materials S3). Drug recognition is done in a similar manner using strings from RxNorm. An independent study at the University of Pittsburgh, which examined the annotations on 1960 clinical notes manually,⁴⁰ estimated over 84% recall and 84% precision for recognizing drugs (personal communication, Richard Boyce).

We focus our study on a set of 14 adverse events based on existing literature,⁴¹ the availability of known DDIs causing them, their population event rate in STRIDE, and our ability to successfully detect their presence in text from EHRs. For interoperability with FAERS, which uses MedDRA codes, we manually assign to each event concept, the most closely matching MedDRA code (see online supplementary materials S4).

Computing drug–drug–event association scores

To compute the association score for a *drug–drug–event* tuple using EHRs, we treat the combination of two drugs as a single drug and use standard methods that measure the disproportionality of the mention of the adverse event, between exposed and comparison groups. Similarly, for FAERS based analysis, we use the Multi-Item Gamma Poisson Shrinker (MGPS)⁴² algorithm to measure the disproportionality of the observed number of reports for a *drug–drug–event* tuple compared to the expected number of reports. We signal an interaction if the association score is greater than some threshold, which is chosen based on the receiver operator characteristic (ROC) curve for the desired sensitivity and specificity.

DDI associations from EHRs

We first construct a 2×2 contingency table (figure 2) where the exposed group comprises patients who have taken both drugs and the comparison group comprises patients who have taken at most one drug (or no drug). Patients are assigned to one of the cells of the contingency table based on the ordering of the first mention of the two drugs and the event in the patient’s timeline (figure 2). For example, if both drugs appear in notes whose timestamp is earlier than the note with the adverse event, the patient is counted into the ‘a’ cell. Any drugs that appear after the first occurrence of the event are ignored. The population event rate among patients on the drug combination can be calculated from this table as $a/(a+b)$.

We calculate an unadjusted OR (UOR) from this contingency table, and use the lower bound of the 95% CI,⁴³ which we denote as $UOR_{0.25}$, as an unadjusted association score. To compute an adjusted OR (AOR), we use propensity score matching (PSM) using a standard caliper of 0.05 (Matching package in R⁴⁴), to match 10 patients in the comparison group to each patient in the exposed group without replacement, followed by conditional logistic regression⁴⁵ (Survival package in R⁴⁶). Once again, we use the lower bound of the 95% CI of the AOR, denoted as $AOR_{0.25}$ as an association score. We compute the

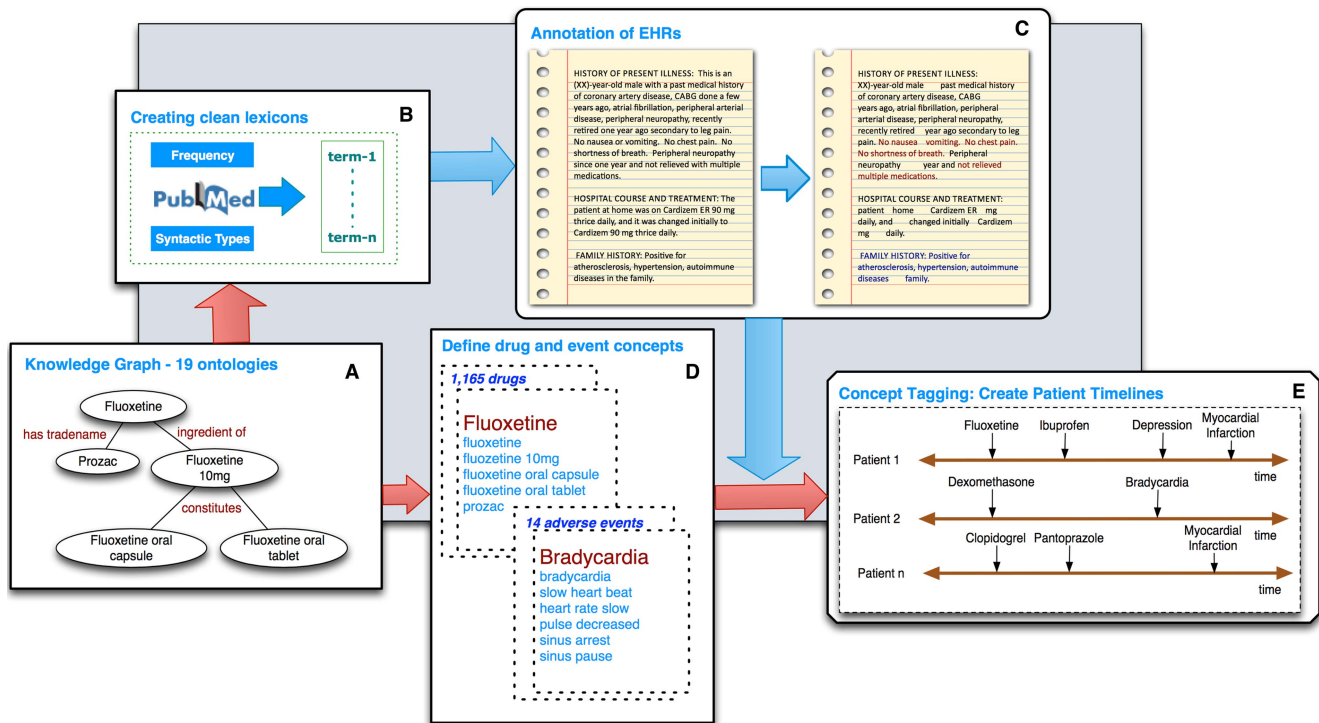
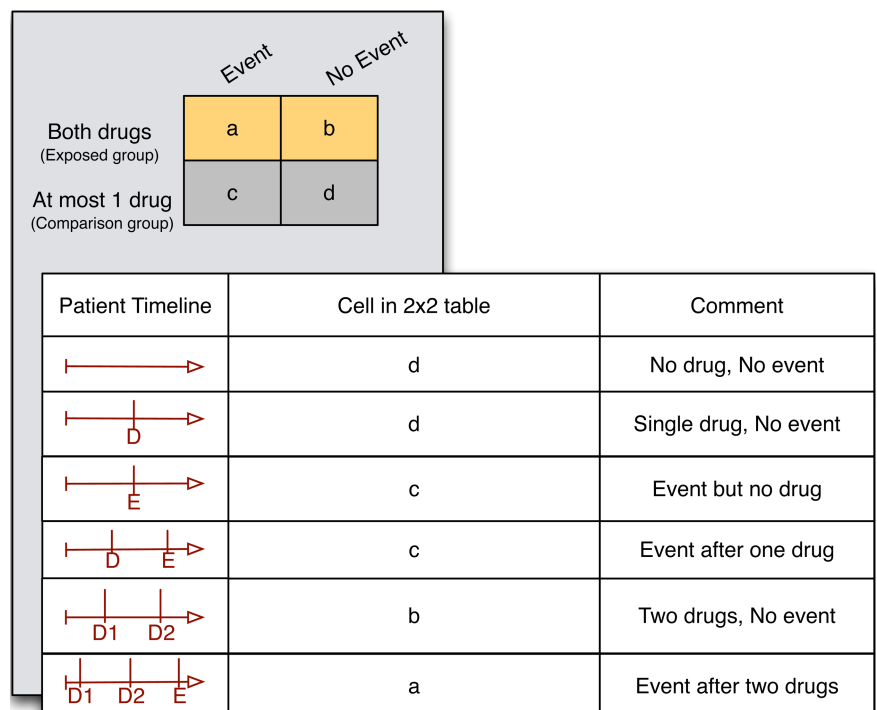


Figure 1 The annotator workflow. (A) The annotator uses a lexicon of approximately 5.6M terms derived from the Unified Medical Language System (UMLS) and BioPortal, as well as trigger terms for NegEx and ConText. (B) It uses term frequency and syntactic type information from Medline to prune the set of strings into a clean lexicon. (C) It then uses the lexicon for exact string matching on the textual notes, followed by negation detection (red) and family history detection (blue). The output is a list of positively mentioned terms recognized in the text. (D) UMLS and BioPortal terms are used to define concepts (a set of terms), making use of the relationships in the ontologies to expand the set. (E) Each note is tagged with a concept if any one of the defining terms appears in the note as a positive mention. The concepts are ordered by the note's timestamp, creating a concept timeline for each patient.

propensity score using age, gender, race, note count, drug count, and disease count. For patients taking at least one of the drugs, we use the average age at the time of drug exposure whereas for patients not taking either drug, we use the average

age of the patient in our dataset. Drug count and disease count serve as proxies for the overall health of the patient,^{27 47} and including the note count serves to adjust for the quantity of data for each patient.

Figure 2 Assignment of patients to various cells of the 2x2 contingency table. The portion of the timeline after the first occurrence of the event is ignored. D, drug; E, event.



DDI associations from FAERS

We use the MGPS, which is a commonly applied algorithm endorsed by the FDA for FAERS based disproportionality analysis,⁴⁸ to generate a DDI signal based on FAERS. MGPS is based on a Bayesian framework that accounts for uncertainty of a disproportionality ratio by ‘shrinking’ it towards a baseline case of no association to an extent proportional to its variance. We make use of the EB05 measure, which is the lower 5th percentile of the posterior distribution of the disproportional reporting ratio for our *drug–drug–event* tuple. The MGPS algorithm performs stratification on age, gender, and year of report to adjust for confounding by these variables. Drug combinations with fewer than five reports are ignored and are given a score of 0.

Preparation of gold standard

We use three sources to define our list of drugs: RxNorm (4993 drugs, counted as unique ingredients), DrugBank (6711 drugs), and ATC (4406 drugs). We limit our study to 1165 drugs present in all three sources. We use known interactions from DrugBank and Medi-Span as positive interactions in our gold standard. Using our concept definitions to identify adverse events from the interaction monographs, we form drug–drug–event relations and manually validate them. For estimating the false discovery rate, we simulate a set of negative interactions by generating random drug–drug–event tuples, and removing any known interactions according to DrugBank, Medi-Span, or Drugs.com. We also remove tuples for which the event is an indication (from Medi-Span, DrugBank, Drugs.com, UMLS, and SIDER) for either drug individually.

RESULTS

Characteristics of datasets

We include 1165 drugs and 14 adverse events in our study and the defining term sets are available as online supplementary materials S2. Out of 1.04 million patients, there are 565 998 patients (53% female) in the STRIDE dataset, with at least one drug or event concept from our study mentioned in their records. Out of the 1165 drugs, 858 (73.6%) appeared at least once. Similarly, out of 1.2 million patients, there are 969 511 patients (54% female) in the PAMF dataset, covering 1048 (90%) drugs. Table 1 shows the number of patients corresponding to each adverse event in STRIDE and PAMF datasets.

Evaluation using gold standard and an independent dataset

DrugBank contains 10 906 DDI monographs and Medi-Span contains 40 475 DDI monographs. Together, there were 46 434 interactions that result in 6346 drug–drug–event tuples (13.66%) corresponding to our set of drugs and events (figure 3). Of these, 849 (13.4%) interactions (comprising 443 drugs and 14 events) satisfy the support threshold of exposure of at least 100 patients in the STRIDE data and these form our set of positive test cases. Table 1 shows the number of positives test cases per event. For each event, we generate as many negative test cases as there are positives (see Methods), thus resulting in 849 randomly generated negative test cases.

We use ROC curves that show all possible values of sensitivity and specificity that can be achieved by our association scores on our gold standard by varying the threshold parameter, as a measure of performance. Using $UOR_{0.25}$, the area under the ROC (AUC) curve is 71.4% and we obtain a specificity of 81.27% and a positive predictive value (PPV) of 72.78% at a sensitivity of 50%. We found that acute renal failure (ARF), nephrotoxicity, hypokalemia, and hyperglycemia did not

Table 1 Characteristics of datasets and gold standard by adverse event

Event	No. patients in STRIDE	No. patients in PAMF	No. positive test cases
Total	565 998	969 511	849
Cardiac arrhythmias	88 555	109 601	65
Acute renal failure	32 197	28 994	15
Bradycardia	22 906	24 162	52
Hyperglycemia	19 189	41 574	47
Neutropenia	14 322	6783	13
Hypoglycemia	11 150	28 320	43
Pancytopenia	8718	2850	2
Hypokalemia	8405	10 356	44
Hyperkalemia	4973	7691	142
Parkinsonian symptoms	3541	4605	9
Nephrotoxicity	1460	639	83
Rhabdomyolysis	1378	1948	30
QT prolongation	1260	1441	150
Serotonin syndrome	674	1511	154

We consider 14 adverse events for our study; our gold standard contains a total of 849 positive test cases and 849 randomly generated negative test cases. PAMF, Palo Alto Medical Foundation; STRIDE, Stanford Translational Research Integrated Database Environment.

perform well (AUC ≈ 0.5 , see Discussion). On removing these events (1320 remaining interactions), the AUC improves to 78.3% and the use of $AOR_{0.25}$ further increases the AUC to 82.3% (figure 4). Additionally, specificity improves to 94.24%, with a PPV (precision) of 89.67% at 50% sensitivity (recall) (see online supplementary materials S5 for precision–recall curves). Overall performance suggests that the unstructured text contains relevant information for signaling DDIs. We also observe differential performance when we look at event specific ROC curves (figure 5).

Of the 1320 interactions in our gold standard, 1132 interactions contain enough support (exposure of at least 100 patients) to signal an interaction in the PAMF dataset, and using $AOR_{0.25}$,

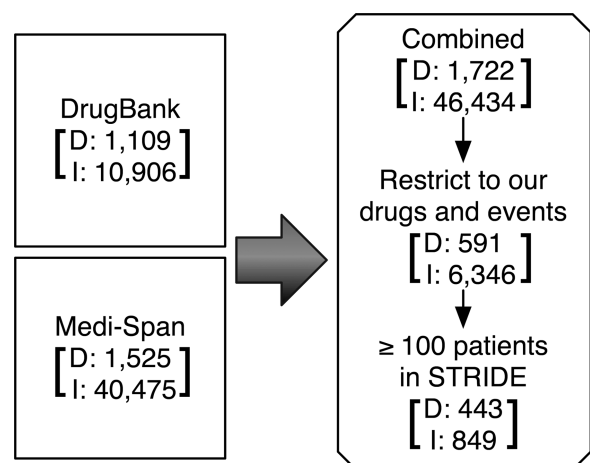


Figure 3 Preparation of Gold standard. We use known interactions from DrugBank and Medi-Span having at least 100 patients on the drug combination in the Stanford Translational Research Integrated Database Environment (STRIDE) as the true positives in our gold standard. The number of drugs (D) and interactions (I) at each stage are specified.

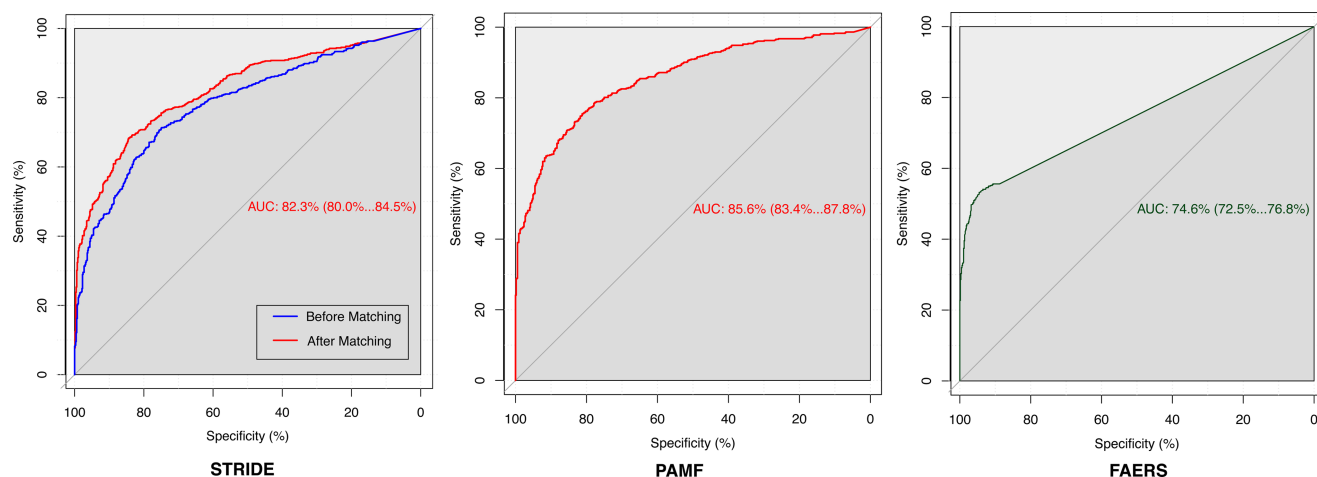


Figure 4 Performance on the Stanford Translational Research Integrated Database Environment (STRIDE), Palo Alto Medical Foundation (PAMF), and FDA Adverse Event Reporting System (FAERS) datasets as evaluated by the gold standard: receiver operator characteristic curves showing sensitivity and specificity levels that can be achieved by varying the threshold. Performance improves after propensity score based matching (red curve in STRIDE and PAMF). For STRIDE, we use our gold standard of 1320 interactions on 10 adverse events. 1132 out of 1320 interactions have enough support for signaling from PAMF. FAERS uses all 1320 test cases, and test cases without enough reports in FAERS were given a score of 0.

we achieve an AUC of 85.6% (figure 4), with a specificity of 95.8% and a PPV (precision) of 92.7% for a sensitivity (recall) of 50% (see online supplementary materials S5).

Evaluation using FAERS

Of the 1320 interactions, 879 did not satisfy the support threshold for FAERS ($EB05=0$); 293 of these 879 were known true interactions, illustrating that several interactions can be missed in FAERS. The $EB05$ measure on FAERS achieves an AUC of 74.6% on the gold standard (figure 4), showing that performance of our method on EHRs seems to be comparable with that of current methods on FAERS—as assessed by our gold standard. Additionally, on running our workflow on STRIDE using data up to each year between 2000 and 2011, we find that the interaction between amiodarone and haloperidol known to cause QT prolongation is signaled using STRIDE data as early as 2007. FAERS reports for this interaction started appearing in 2009, possibly correlated with a research publication at that time by Bush *et al.*,⁴⁹ demonstrating the ability of EHRs to complement FAERS for early detection.

Population rate of adverse events

Using our set of 1165 drugs and 14 events, there are 569 398 *drug–drug–event* tuples with at least 100 patients on both drugs in STRIDE. We publish the 2×2 contingency tables along with population event rate and $UOR_{0.25}$ for these tuples in STRIDE as online supplementary materials S6; this population event rate information has potential utility in prioritizing DDIs in several settings. Table 2 shows the drug combinations with the highest event rates in STRIDE, for each event from our gold standard.

Utility of association scores

We compute $AOR_{0.25}$ for those tuples that are highly likely to represent an interaction. To choose these tuples, we use a three-step strategy. We first obtain thresholds that result in 95% specificity for $UOR_{0.25}$ on STRIDE (threshold=4.7) and $EB05$ in FAERS (threshold=1.5) as assessed by our gold standard. We then compute $EB05$ using FAERS for all 569 398 tuples; we already have $UOR_{0.25}$ for these 569 398 tuples. Finally, we compute $AOR_{0.25}$

for those 9306 tuples that satisfy the $UOR_{0.25}$ and $EB05$ thresholds for 95% specificity (see online supplementary materials S6).

Based on $AOR_{0.25}$, a threshold of 1.1 gives us 95% specificity with respect to our gold standard. Of the 9306 tuples, 5983 satisfy this threshold and have a high likelihood of representing true interactions. We also find that 49 of these 5983 tuples (see online supplementary materials S7) were added to Drugs.com in the period between May 2011 and November 2012 (STRIDE data is up to March 2011). Therefore we believe that EHRs can be used for hypothesis generation to identify new DDI signals.

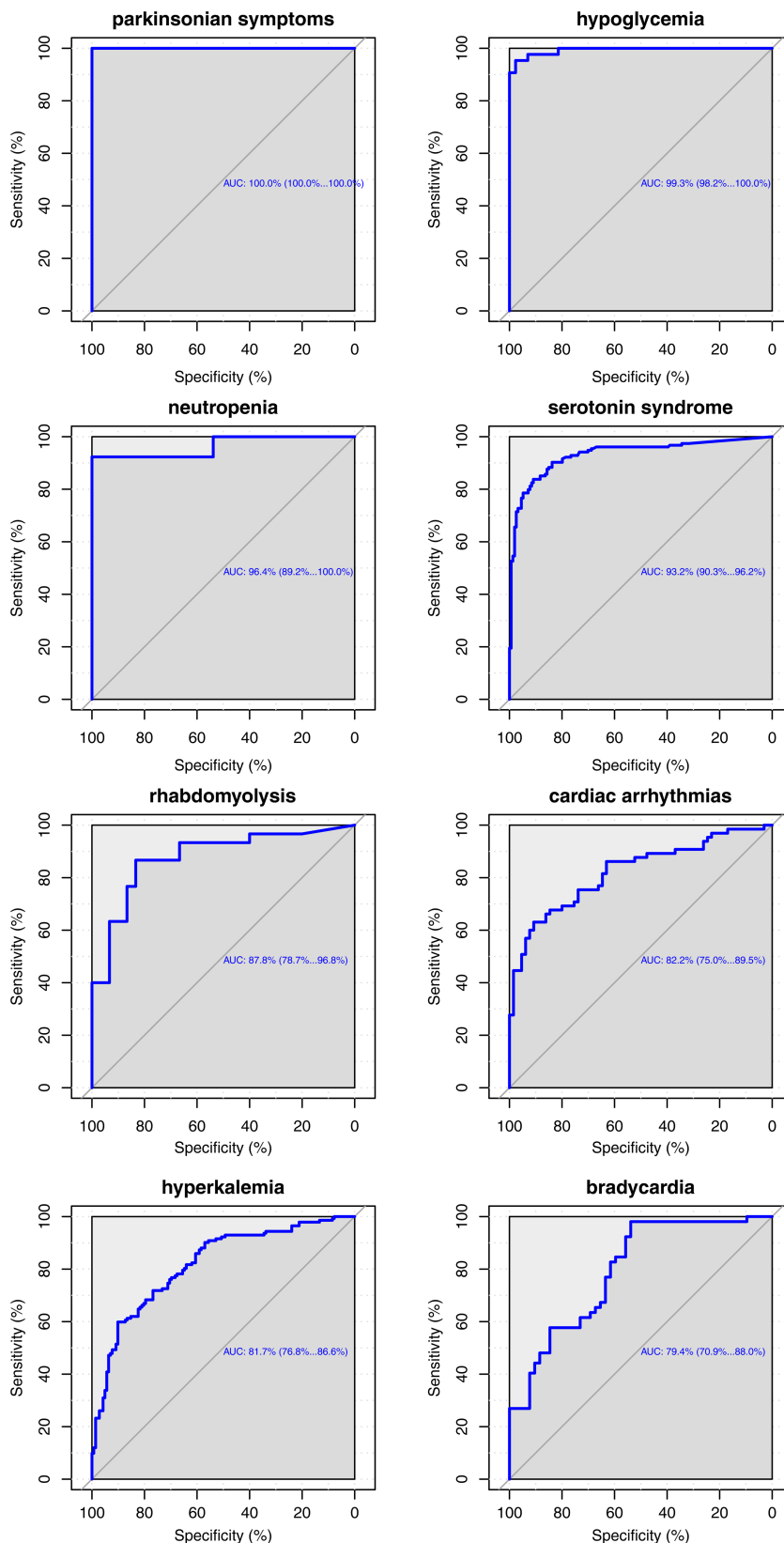
Table 3 shows $AOR_{0.25}$ for the combination of statins with calcineurin inhibitors, tacrolimus, and cyclosporine A, associated with rhabdomyolysis. Using a threshold of 1.1 (95% specificity), the combination of tacrolimus and statins is relatively less associated with rhabdomyolysis. This is in agreement with a study by Lemahieu *et al.*⁵⁰ in 2005, which argues that patients exposed to statins and cyclosporine A are at an increased risk of rhabdomyolysis, whereas tacrolimus is safer. This result illustrates a potential utility of the EHR-derived association scores for clinical decision support—to choose between several viable treatment alternatives.

DISCUSSION

The early identification of DDIs is important and testing for interactions between all drugs by experimental methods is infeasible. Significant research exists on generating models for metabolic interactions of drugs,⁵¹ mainly for interactions related to CYP enzymes. There is also work on predicting interactions based on molecular similarity with drugs already known to interact.⁵² Although these methods are successful at finding new interactions, their results are limited by their modeling assumptions and the limited mechanistic understanding of known interactions. A data-driven way to identify interactions is by analyzing the effect of drugs in the general population via post-marketing surveillance.

Most existing methods for such surveillance use SRS databases to identify interactions, and use coded information present in EHRs for validation and prioritization. We have shown that it is possible to identify significant (drug–drug–event) associations and compute their event rates, directly from

Figure 5 Event-wise performance in the Stanford Translational Research Integrated Database Environment (STRIDE): receiver operator characteristic (ROC) curves showing sensitivity and specificity values for various thresholds on the gold standard test cases using STRIDE. Using such curves, event specific thresholds can be chosen. Hyperkalemia, acute renal failure, nephrotoxicity, and hyperglycemia did not perform well. This could be due to our inability to accurately tag notes with these concepts, or due to the gold standard itself (see Discussion). The area under the ROC curve for pancytopenia has a very large variance due to an insufficient number of tested interactions.



the unstructured text of EHRs. This can serve as an active monitoring tool for signaling unknown interactions for new and existing drugs; advancing phase IV surveillance of drugs and meaningful use of EHRs simultaneously.

We perform text mining on two large corpora of clinical records to successfully detect DDIs with a simple and fast

approach to text annotation,⁵³ where we sacrifice some accuracy in tagging concepts at the note level in exchange for population level trends (see online supplementary materials S3). We test our results using a gold standard comprising known DDIs as positive examples and an equal number of randomly generated interactions as negative examples. Testing AOR_{0.25} using all unknown

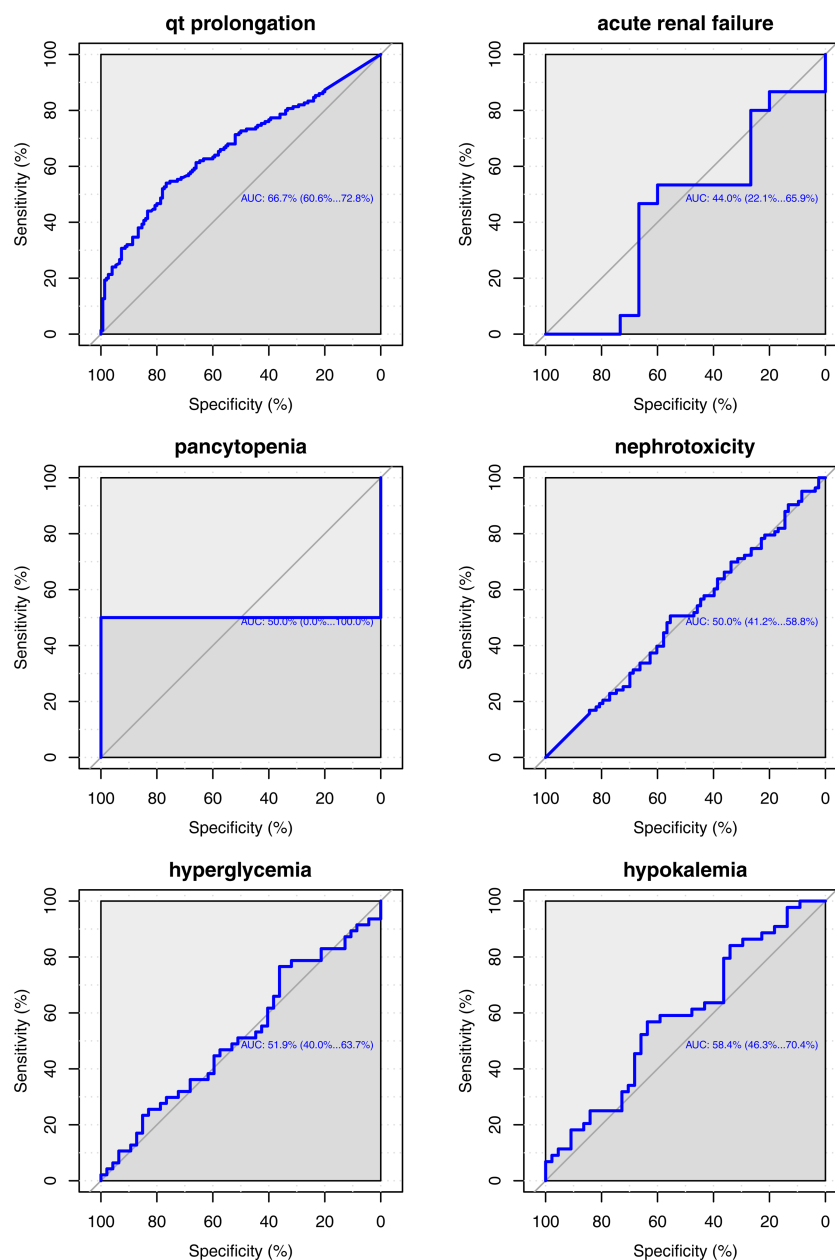


Figure 5 Continued

interactions as negative examples poses computational problems; however, the AUC for UOR_{0.25} using all negatives is almost identical to that using sampled negatives, indicating minimal biases due to sampling. Most interaction studies^{52–54} use similar techniques to build a gold standard and the generation of an ROC aids in comparison of results. We see differential performance per event; for example, the performance of the method for parkinsonian symptoms, hypoglycemia, and neutropenia is significantly better than the others (in terms of AUC). This finding reflects similar findings in the EU-ADR project,⁵⁵ and suggests the need for event specific thresholds.⁵⁶

Our approach did not perform well for four events, perhaps owing to limitations that we discuss here. The performance of our workflow heavily depends on the accuracy of concept recognition in EHRs using lexical approaches. In this study, we use frequency based methods^{36–37} and manual curation to remove ambiguous terms corresponding to concepts—at the expense of reducing sensitivity. Nevertheless, some concepts are hard to

detect in EHRs using lexicon based methods; for example, the occurrence of hypokalemia or hyperglycemia is sometimes stated in quantitative terms as changes in potassium or sugar levels, which would be missed by our annotator. On the other hand, having too few terms reduces our sensitivity; for example, the term set for nephrotoxicity comprised only three terms. Our current method cannot signal DDIs that are dependent on drug dosage and resolving these issues requires advanced natural language processing methods.⁵⁷ We focus solely on the note's timestamp to determine the time of occurrence of a concept; we acknowledge that doing so is based on the assumption that an event described as historical in the note will be mentioned in some previous note as a current event, and this can reduce performance. We do not use drug interaction eras and drug exposures spaced far away in time may cause false associations. We are currently examining the annotations for the utility of the last mention of concepts, sentence-level co-occurrences, and temporal density of mentions to address this question. We use

Table 2 Interactions in the gold standard with the highest event rates ($a/(a+b)$) in the Stanford Translational Research Integrated Database Environment dataset for each event

Adverse event	Drug1	Drug2	a	b	Rate (%)
Parkinsonian symptoms	Levodopa	Lorazepam	176	235	42.82
Cardiac arrhythmias	Potassium chloride	Lisinopril	1091	1615	40.32
Neutropenia	Paclitaxel	Trastuzumab	140	567	19.8
Bradycardia	Amiodarone	Metoprolol	796	3671	17.82
Hypoglycemia	Glipizide	Lisinopril	367	2160	14.52
Acute renal failure	Hydrochlorothiazide	Ibuprofen	884	8375	9.55
Hyperkalemia	Potassium chloride	Spirolactone	349	3471	9.14
Hyperglycemia	Prednisone	Salmeterol	379	4612	7.59
Nephrotoxicity	Fluconazole	Tacrolimus	85	1208	6.57
Pancytopenia	Mercaptopurine	Azathioprine	15	278	5.12
Hypokalemia	Prednisone	Salmeterol	222	4982	4.27
Serotonin syndrome	Tramadol	Duloxetine	57	1301	4.2
QT prolongation	Amiodarone	Ciprofloxacin	46	2487	1.82
Rhabdomyolysis	Ciprofloxacin	Simvastatin	50	5184	0.96

standard disproportionality analysis adapted to longitudinal data to signal an interaction, however several other measures of interaction have been proposed in the literature^{58–60} and may prove to perform better. To adjust for possible confounding factors, we use PSM to choose a comparison group that is similar to the exposure group and then use matched conditional logistic regression to generate an AOR. We acknowledge that this approach is one of several known methods to use PSM⁶¹ and that other methods could be equally effective. The databases from which the known interactions are derived are not exhaustive and some interactions are based on unsubstantial evidence; for example, several monographs for ARF claim that ‘ARF was reported on rare occasions’. We use several sources of data in this work, each having its own method for coding drugs and events, and the mapping from concepts in one data source to the other is another source of errors; for example, the mapping from our event concepts to MedDRA PTs for use with FAERS. Finally, we note that our association scores do not indicate causality—which means that in some cases, the adverse event might not be caused by a drug interaction in the mechanistic sense but might be associated with the exposure to the drugs. The goal is to provide early warnings of drug combinations that require investigation.

Using standard methods to analyze FAERS, we show its performance on our gold standard as a reference to enable an evaluation of the comparative utility of EHRs. Many known interactions from our gold standard that appear in EHR data do not contain a sufficient number of reports in FAERS, possibly

owing to under-reporting. EHRs may therefore have utility in the early signaling of DDIs. Furthermore, we demonstrate a potential use of EHR derived DDI association strengths to choose between drugs used in combination therapy; for example, the results on the combination of calcineurin inhibitors with statins. Such analysis is not possible with FAERS alone owing to the differential reporting rates for drugs.

EHRs have good longitudinal coverage of patient history, a larger number of measured covariates, and thus are likely to provide an accurate measure of the real world rate of adverse events among patients on a particular drug combination. The pre-computed population event rate information and association scores could find application in clinical decision support. For example, augmenting existing DDI databases with this event rate information could potentially be useful to prioritize interaction alerts in computerized physician order entry (CPOE) systems,⁶² where at present 49–96% of all alerts are overridden.⁶³ Finally, calculating population event rate information could directly serve as a means to enable stage 3 of the meaningful use measures that are to be implemented by 2016.⁶⁴ To this end, we publish the event rates (see online supplementary materials) for 569 398 *drug–drug–event* combinations in our study. We emphasize that the limitations of our method, as highlighted above, must be accounted for when using the estimated rates of events. With the intent of finding novel DDIs, we also publish the adjusted association scores for 9306 interactions that are signaled by both EHR sources and FAERS. However, the number of predictions are still far too many to be experimentally tested; it may be possible to identify the most promising interactions using a weighted score based on features such as the number of affected people and the cost of the drugs, or by searching for a mechanistic explanation as proposed by Bauer-Mehren *et al.*⁶⁵

CONCLUSION

We use data from two independent sites to demonstrate the feasibility of using the textual notes from EHRs for signaling DDIs and to estimate the rate of events among patients on various drug combinations. We develop a gold standard of DDIs that may be useful for detailed characterization of future methods and our evaluations show that we can signal interactions from EHRs with as good performance as established methods on SRSs (FAERS). We find that in some cases we can signal interactions many years before the first report in FAERS, which could expedite the discovery of new interactions and demonstrates the complementarity of our approach. Additionally, we publish a first-of-its-kind resource: a database of the rate of each event for all drug–drug pairs from our EHR corpus, which may be useful to triage alerts in CPOE systems or to identify drug combinations with a lower chance of adverse events. We conclude that the text portion of EHRs can complement existing sources for postmarketing surveillance for DDIs.

Table 3 Potential for clinical decision support

Event	Statin	Calcineurin inhibitor	a	b	c	d	AOR _{0.25}
Rhabdomyolysis	Nystatin	Tacrolimus	21	2324	1357	562 296	0.67
	Atorvastatin		20	1267	1358	563 353	0.96
	Nystatin	Cyclosporine	32	1407	1346	563 213	1.78
	Atorvastatin		37	1331	1341	563 289	2.26

This table shows AOR_{0.25} scores for the association of rhabdomyolysis with the combination of calcineurin inhibitors like tacrolimus or cyclosporine A, and statins. It appears that tacrolimus is relatively safe and this agrees with in vivo experiments. AOR, adjusted OR.

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