Applying MetaMap to Medline for identifying novel associations in a large clinical dataset: a feasibility analysis

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ABSTRACT

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not in Medline citations, are potentially novel. **Methods** Pairwise associations of ICD-9 codes were independently identified in both the clinical and Medline datasets, which were then compared to quantify their degree of overlap. We also performed a manual review of a subset of the associations to validate how well MetaMap performed in identifying diagnoses mentioned in Medline citations that formed the basis of the Medline associations.

Results The overlap of associations based on ICD-9 codes in the clinical and Medline datasets was low: only 6.6% of the 3.1 million associations found in the clinical dataset were also present in the Medline dataset. Further, a manual review of a subset of the associations that appeared in both datasets revealed that cooccurring diagnoses from Medline citations do not always represent clinically meaningful associations. Discussion Identifying novel associations derived from large clinical datasets remains challenging. Medline as a sole data source for existing knowledge may not be adequate to filter out widely known associations. **Conclusions** In this study, novel associations were not readily identified. Further improvements in accuracy and relevance for tools such as MetaMap are needed to realize their expected utility.

INTRODUCTION

The age of 'Big Data' has arrived.^{1–3} Studies using data collected as part of routine clinical care from hundreds of thousands, or even millions, of patients are becoming increasingly common.^{4–8} Other large datasets (eg, adverse event reports) are also being linked to these clinical data to accelerate discovery,⁹ ¹⁰ leading to new findings of intriguing and potentially clinically relevant associations that could aid in the understanding of disease processes.¹⁰ ¹¹ For example, Tatonetti *et al*¹² recently discovered an association between elevated blood glucose levels and the co-administration of paroxetine and pravastatin, neither of which raised blood glucose when given alone.

Association analysis methods have been widely used to aid in knowledge discovery, where associations are usually determined by finding pairwise relationships among entities that co-occur at a statistically significant rate compared to the overall population. We previously reported on two separate association analyses, one utilizing 1.5 million free text problem list entries from over 300 000 patients⁵ and the other using 41.2 million International Classification of Disease, V.9 (ICD-9) codes from over 1.6 million patients.⁴ The latter dataset represented virtually all possible diagnosisbased clinical associations known to our health system, a large tertiary academic medical center with over 1.8 million outpatient and emergency visits and 44 000 hospital stays annually, derived from over a decade of patient encounters.

In both studies, 4^{45} we noted that a very large number of statistically significant associations resulted from the analyses, making it impossible to identify all novel ones via manual review alone. We also noted that many of the associations, especially the most statistically significant ones, are already widely known. For example, in the study using free text diagnoses, the well-known associations we found included one between obesity and hypertension and one between Turner syndrome and ovarian failure. Lesser known associations that we confirmed with a manual literature review included hypothyroidism and fibromyalgia as well as gout and cardiomyopathy.⁵ An example of a well-known association from the second study using ICD-9 codes included end stage renal disease and kidney transplant, whereas an unusual association with no supporting evidence in the literature was between depression and animal bites (primarily cats).⁴ This latter association was confirmed with a manual chart review.¹¹ However, manually validating the potential novelty of all associations against the literature, from such large-scale analyses, is not feasible. Developing automated approaches that can effectively distinguish known from unknown clinical associations is thus imperative. Such automated methods could be beneficial for a variety of applications, including surveillance of electronic health record data to detect previously unknown or newly arising patterns.

A potential approach to automating the identification of novel associations is through comparing those found in clinical datasets against comprehensive repositories of known associations. The National Library of Medicine's (NLM) Medline/ PubMed database is the world's largest indexed repository of biomedical literature, with over 19 million citations from over 5500 journals.¹³ It might therefore be possible to use Medline as a



To cite: Hanauer DA, Saeed M, Zheng K, et al. J Am Med Inform Assoc 2014;21:925–937. basis from which such a knowledge repository of associations could be assembled and then compared to associations derived from large clinical datasets. Such literature-based discovery techniques, also known as literature mining, are not new.^{14–18} However, prior studies have often focused on specific clinical areas, such as psychiatry^{19 20} or diabetes.^{21 22} It is unclear how well such an approach might work with an 'all versus all' comparison—that is, all patient data from a large health system versus all abstracts in Medline.

Recently, the NLM computationally processed the entire Medline database using their natural language processing (NLP) based named entity recognition software tool, MetaMap.² MetaMap identifies clinical concepts (eg, 'type 2 diabetes') from unstructured biomedical text and then maps them to concept unique identifiers (CUIs) in the Unified Medical Language System (UMLS) Metathesaurus; for example, C0865162 is a CUI for 'diabetes'. These CUIs can, in turn, be mapped to various taxonomies, vocabularies, and ontologies including ICD-9; for example, the CUI above, C0865162, maps to the ICD-9 code '250.0'. MetaMap has been used for a variety of tasks including extracting information from drug labels,²⁴ coding death certificates,²⁵ conducting biosurveillance,²⁶ parsing documents from electronic health records,²⁷ ²⁸ supporting Medical Subject Heading (MeSH) assignments through use of the NLM Medical Text Indexer (MTI),²⁹⁻³² improving information retrieval from Medline,³³ and even assigning ICD-9 codes from clinical text.^{34–36}

We hypothesized that clinical associations derived from administrative ICD-9 codes assigned during patient encounters can be compared against associations extracted from the literature to distill novel associations from known ones. That is, if an association resulting from a clinical dataset is not found in Medline, then it may be potentially novel and warrant further investigation. We also hypothesized that among a collection of associations derived from the same large dataset, clinical associations ranked higher (ie, with smaller p values or larger χ^2 statistics) would be more likely to be identified in the literature than lower ranked ones, based on the assumption that common problems are more likely to have been studied and published. In this study, we developed and empirically validated a set of computational analyses to assess the feasibility of verifying novel associations identified through mining clinical data against the results from mining the literature. Our primary goal in this method development paper is to explore the feasibility of this approach rather than making new clinical discoveries.

METHODS

A high-level overview of the analytic approach used in this paper is illustrated in figure 1. Below, we describe the data sources and each of the analytic steps in-depth.

Description of datasets

Two datasets were used in the analysis. The clinical associations (referred to as 'Clinical') were obtained from the dataset processed for a prior study.⁴ These associations encompass ICD-9 administrative data from both inpatient and outpatient encounters for 1.6 million patients at our health system, and spanned more than a decade. A description of how the clinical associations were computed can be found in the 'Association analyses' subsection, below.

The second dataset (referred to as 'Medline') was provided by the NLM and included all citations in Medline/PubMed as of November 18, 2011. The NLM has named this the '2012 MetaMapped Medline Baseline Results'.³⁷ The dataset contains a total of 20.5 million citations processed by the NLM using MetaMap. It is comprised of 3.3 billion lines of text in the MetaMap Machine Output (MMO) Format, taking over 1.5 terabytes disk space.^{38 39}

Extraction of CUIs

During the named entity extraction process, MetaMap generates a normalized score estimating the degree of match between a candidate term and the Metathesaurus based on four components: *centrality, variation, coverage,* and *cohesiveness.*²³ ⁴⁰ From the Medline dataset, we parsed all CUIs with a 'perfect' MetaMap score of 1000, noting the specific PubMed Identifier (PMID) of the paper(s) from which each of the CUIs was from and whether a CUI was identified in the title or in the abstract of the paper(s). This is referred to as the 'Medline 1000' dataset.

Using CUIs with only a perfect score will exclude alternative ways to express a given concept because it essentially requires an exact match and does not allow for synonomy or slight word variations. Therefore, because it was unclear how much information might be lost by including only perfect mapping scores, we also created another dataset consisting of extracted CUIs where candidate concepts had mapping scores of 600 or higher. This allowed us to broaden the scope of our search to ensure a larger capture rate for concepts, since a CUI mapped to an ICD-9 code might otherwise be 'hidden' by a higher scoring CUI from a different vocabulary. We refer to the second dataset as 'Medline 600'. This threshold was chosen because it was lower, and thus more inclusive, than thresholds commonly used in prior studies (see Discussion).

CUI to ICD-9 mapping

From the UMLS Metathesaurus (V.2013AA), we extracted two vocabularies, namely 'ICD9CM' and 'MTHICD9', in order to map Metathesaurus CUIs to ICD-9 entries. ICD9CM is the clinical modification of the International Classification of Diseases developed by the US Department of Health and Human Services. MTHICD9 provides additional synonyms for ICD9CM terms and was developed for the Metathesaurus by the NLM. Both vocabularies were extracted from the MRCONSO.RRF file that contains concepts, concept names, and their sources (MR: metathesaurus relational; CON: concept; SO: source; RRF: rich release format). For CUIs mapped to more than one ICD-9 code, we included only those that mapped to 50 or fewer distinct ICD-9 codes. This was done because some CUIs mapped to so many different codes that they were considered too non-specific for our analysis. The resulting mapping set contains 36 988 distinct mappings representing 33 999 and 22 198 unique CUIs and ICD-9 codes, respectively. We then processed the CUIs from the MetaMap-prepared Medline citations, retaining only those CUIs that mapped to at least one ICD-9 code.

Association analyses

Leveraging all ICD-9 codes mapped from clinical concepts in Medline citations, we then conducted a large-scale association analysis based on the methodology we had previously developed for discovering clinical associations from patient care data.⁴ In short, we determined the probability and statistical significance of two codes co-occurring in the same patient profile or Medline citation using the χ^2 test. This was done by constructing 2×2 tables for every pair of concepts. For example, in the Medline dataset we determined the χ^2 statistic based on the following conditions: (1) citations that mentioned both diagnosis A



Figure 1 A high-level overview of the analytic approach, including the data sources and processes.

and diagnosis B; (2) citations that mentioned diagnosis A but not diagnosis B; (3) citations that mentioned diagnosis B but not diagnosis A; and (4) citations that mentioned neither diagnosis A nor B. While there are a number of ways to quantify the relationship between two binary variables, the χ^2 test is most popularly used. We used the χ^2 statistic to rank order the associations within each dataset to provide a basis for inferring the 'relative strength' among the associations in the dataset. Note that temporal aspects were not considered in the experiments reported in this paper.

For the Clinical dataset, in accordance with our prior analysis, only codes that appeared at least 30 times and where at least 10 patients shared the same pair of two codes were considered to be potentially associated. For codes under these thresholds a χ^2 test was not performed. By contrast, the association analysis for the Medline dataset considered any pair of codes that appeared together in at least one citation. This was because some citations could represent meta-studies (eg, systematic reviews) as well as studies based on data collected from tens or thousands of patients.⁴¹ These datasets (both 'Medline' and 'Clinical') that included all ICD-9 codes in their original format are referred to as 'Original'.

Additionally, because coding variations may occur among both clinicians and professional coders,^{42–45} and because these variations could affect the nature of associations discovered, we also conducted an additional association analysis by collapsing the hierarchical structure of the ICD-9 taxonomy so that all 'subcategory codes' were merged into their parent 'category code'. For example, codes such as '250.0', '250.23', and '250.80' were converted to simply '250', a high-level code used to designate the entire family of diabetes. These collapsed datasets (both 'Medline' and 'Clinical') are referred to in this paper as 'Simplified'.

High-level dataset comparisons

The primary objective of this study was to determine whether associations identified in the literature could be used to filter out known associations identified from patient care data, thus revealing potentially novel clinical associations. Therefore, we compared the Medline and Clinical datasets to determine how much overlap existed in terms of the ICD-9 codes used and the clinical associations discovered in each. This overlap was quantified and visualized using Venn diagrams (figures 2 and 3).⁴⁶

We then modeled the probability of an association being in the literature as a function of its ranking in the clinical dataset. Associations in the clinical dataset were ranked according to their χ^2 statistic, and each association was assigned a percentile rank ranging from 1st (highest) to 100th (lowest). For all of the clinical associations assigned to the same percentile, we determined which ones had corresponding associations in the Medline dataset. These results were visualized using bar plots (figure 4). We also modeled the converse—that is, the probability of an association being in the Clinical dataset as a function of its χ^2 ranking in the Medline dataset.

Association-specific comparisons

We also looked at individual associations derived from our clinical dataset to better estimate the utility and reliability of using the literature as a screening tool to validate novelty. We examined this through two steps. First, to determine the reliability of the associations found in both the Clinical and Medline datasets, we randomly selected Clinical associations at several specific levels of significance (eg, 50th percentile, 75th percentile) for which there were corresponding Medline associations. ICD-9 'V' codes were not considered in this part of the analysis because they are often less directly related to specific diagnoses. Two practicing, informatics-trained physicians (DAH and MS), with 14 and 6 years of postgraduate experience, respectively, independently reviewed the titles and abstracts from which the literature-based associations were derived. Each citation was judged to either support the association identified from the clinical dataset or not to support it. Additionally, associations were judged to be clinically 'surprising' if there was no clear explanation for why the two diagnoses might be related. Agreements in the two reviewers' judgments were quantified using the κ statistic.

We then randomly sampled clinical associations for which there were no corresponding associations found in the Medline dataset to determine if they might represent novel associations, and to compare the performance of MetaMap against manual strategies inspecting for novelty. The same two physicians reviewed these associations to determine if they were clinically surprising. They also conducted a manual literature search in Medline in an attempt to find out whether, for each of the associations, at least one citation could be found that provides evidence confirming its validity.



Figure 2 Venn diagrams showing overlap of International Classification of Disease, V.9 (ICD-9) codes found in the Clinical and Medline datasets. Panels A and C represent the original codes, whereas Panels B and D represent the simplified codes. Codes that do not overlap between the two datasets have no chance of becoming a pairwise association found in both sets.

Additional analyses

We performed several additional analyses and report the results in online appendices A, B, and C. Online supplementary appendix A provides several groups of scatter plots exhibiting the relationship between the frequency of an ICD-9 code appearing in each of the datasets analyzed, and the corresponding number of associations to which it belonged. In online supplementary appendix B, we explored the distribution of association rankings as a function of an ICD-9 code's frequency in the original clinical dataset. We selected 15 ICD-9 codes that appeared with varying frequencies in the clinical dataset for this experiment. Finally, in online supplementary appendix C, we explored the potential for using the UMLS relationships as defined in the UMLS relational table MRREL.RRF as another source of associations that could be used to help filter known from unknown associations. Additional methodological details can be found in each of the online supplementary appendices. Note that in this study, we did not explore the use of Semantic MEDLINE^{47 48} because the relationships contained in Semantic MEDLINE are directly derived from MetaMap-processed Medline citations, where are therefore not expected to generate substantially different results from our primary analysis.

All statistical analyses reported in this paper were conducted using *R* V.2.15.3. Venn diagrams were created using the VennDiagram Package for *R*.⁴⁶ While we used the χ^2 statistic to rank the associations, the corresponding p values are also reported to aid in interpretation. For computation of the datasets, we used a 2010 Apple Mac Mini equipped with a 2.66 GHz Core 2 Duo processor and 8 GB RAM, and for storage we used an external 4 TB hard drive connected via USB 2.0.

RESULTS

Characteristics of the datasets

Processing the large dataset of all Medline citations yielded 333.3 million non-distinct CUIs with a perfect score of 1000. About 28.5 million of these CUIs (8.5%) were identified in the titles, with the remaining 304.9 million (91.5%) identified in the abstracts. There were approximately 16.3 million unique citations represented in this dataset. However, only a subset of the CUIs, approximately 23.1 million CUIs representing 5.1 million unique citations, mapped to at least one of 7599 distinct ICD-9 codes. Additional characteristics of the dataset are shown in table 1.

The 'Medline 600' dataset used less stringent criteria for selecting CUIs for inclusion, and thus 3.5 billion non-distinct CUIs were identified, with 348.2 million (9.8%) in the titles and 3.2 billion (90.2%) in the abstracts. This dataset contained 20.4 million unique citations, representing all but 0.4% of the citations included in the original dataset. Only CUIs that mapped to an ICD-9 code were retained, resulting in 148.0 million CUIs, 10.0 million citations, and 9416 unique ICD-9 codes. Table 1 also summarizes the characteristics of this dataset.

Only one ICD-9 code (401.9, 'unspecified essential hypertension') appeared in the top 30 most frequently appearing codes in the Original Clinical dataset as well as the top 30 of the



Figure 3 Venn diagrams showing overlap of pairwise associations found in the Clinical and Medline datasets. Panels A and C display the Original datasets, where as Panels B and D represent the Simplified datasets. The left most area in each panel is the one most likely to contain novel associations not previous described in the literature as they are found in the Clinical dataset but not the Medline dataset.

Original Medline datasets. By reducing the coding variation in the Simplified datasets, six of the top 30 codes in the Clinical dataset can also be found in the top 30 of at least one of the corresponding Medline datasets (table 2). For the Clinical datasets, this list represents the most common diseases (diagnoses) treated by our health system. For the Medline datasets, it shows the most common diagnoses discussed in the literature, based on MetaMap.

Figure 2 shows the overlap of the ICD-9 codes that were included in the datasets being compared. Slightly more than a third (37.1%) of the ICD-9 codes from the Original Clinical dataset were found at least once in the Original Medline 1000 dataset, and only 44.4% were found in the more inclusive Original Medline 600 dataset. When the datasets were simplified by merging subcategory codes, more overlap was evident: four-fifths (79.4%) of the codes in the Simplified Clinical dataset were in the Simplified Medline 1000 dataset, and 84.3% were in the Simplified Medline 600 dataset.

Clinical associations also found in Medline

We hypothesized that many of the associations discovered in the Clinical dataset would have been reported in Medline, achieving a more manageable number of potentially novel associations to be explored further for clinical significance. However, this was not the case according to the results of our analyses. As shown in the Venn diagrams illustrating the overlap between the Medline and Clinical datasets (figure 3), in the original datasets, only 6.6% of the Clinical associations had a correlate in the

Medline 1000 dataset, with slightly more (10.7%) found in the Medline 600 dataset. The simplified datasets, where the ICD codes of the same family were merged into parent categories, displayed higher coverage, with 31.2% of the Clinical associations also present in the Medline 1000 data, and almost half (44.5%) of the Clinical associations also found in Medline 600.

We anticipated that higher ranked Clinical associations (based on larger χ^2 statistics or, conversely, smaller p values) would be more likely to be found in Medline, and this was evident in the bar plots shown in figure 4, where A/C and B/D show the trends for the Clinical datasets, Original versus Simplified, respectively. In figure 4, it can clearly be observed that higher ranked associations in the Clinical set in general were more likely to be found in the Medline dataset. However, our results did not suggest as strong a correlation for the converse. That is, the relationship between the rank of an association found in Medline and the likelihood of finding that association in the Clinical dataset was not as strong (figure 5).

Specific associations

Table 3 shows associations identified from both the Clinical and Medline 1000 datasets at varying levels of significance. As shown in the table, the lower-ranked associations tended to be more clinically 'surprising'. However, our manual review of the citations showed that many of the titles/abstracts mentioning two diagnoses together did not truly suggest an actual association between the diagnoses. That is, the mere mention of two diagnoses in the same citation is not a reliable indicator of a **Figure 4** Bar plots showing the relationship between the rank of an association in the Clinical dataset and the probability of the association appearing in the Medline dataset. (A) Clinical (original) appearing in Medline 1000 (original); (B) Clinical (simplified) appearing in Medline 1000 (simplified); (C) Clinical (original) appearing in Medline 600 (original); (D) Clinical (simplified) appearing in Medline 600 (simplified). The p-values shown represent the significance of the Clinical associations at that percentile.



clinically meaningful association. Table 4 displays associations in the Clinical dataset that were not present in Medline 1000. Again, clinically surprising associations tended to be less significant (lower-ranked), and they were also less commonly found in Medline even with a manual search.

Additional results

The experiments reported in online supplementary appendices A–C revealed several additional insights. First, ICD-9 codes that occur more frequently in the Clinical dataset tend to belong to more associations than those codes that occur rarely (see online supplementary appendix A). Second, for many of the ICD-9 codes, the distribution of association rankings to which each

code belongs demonstrated greater spread in the Clinical dataset compared to the Medline dataset (see online supplementary appendix B). Finally, relationships defined within the UMLS relational table can be an additional resource for known associations that are not found within Medline (see online supplementary appendix C). However, the Clinical dataset still contained many associations not included in the UMLS relational table.

DISCUSSION

The results of the experiments reported in this paper show that while combining literature mining and clinical data mining could aid in the discovery of novel associations, the overlap between the Clinical and Medline datasets was surprisingly low.

Table 1 Characteristics of the datasets used in the analysis							
	Original data	isets		Simplified datasets			
	Clinical	Medline 1000	Medline 600	Clinical	Medline 1000	Medline 600	
Rows of data	41 192 825	23 051 034	147 994 791	41 192 825	23 051 034	147 994 791	
Patients (clinical data)/citations (Medline data) in full dataset	1 620 280	5 069 886	9 975 979	1 620 280	5 069 886	9 975 979	
Patients/citations included in final pairwise comparisons dataset	1 619 785	5 069 347	9 975 705	1 620 271	5 069 851	9 975 966	
Distinct ICD-9 codes	14 499	7599	9416	1195	1073	1133	
Possible pairwise comparisons*	105 103 251	28 868 601	44 325 820	713 415	575 128	641 278	
Actual pairwise comparisons†	3 066 673	1 318 933	2 718 577	315 681	165 391	257 558	
Unique ICD-9 codes meeting criteria to be used in a pairwise comparison‡	8601	7309	9233	1099	1057	1126	

*Possible pairwise comparisons is determined by $(n^2-n)/2$, where n is the number of distinct ICD-9 codes in the dataset.

tA pairwise comparison was only calculated under the following conditions: (1) for the clinical data if (a) each code was assigned to at least 30 patients and (b) the pair of codes were present together in at least 10 patients; (2) for the Medline data if (a) each code was assigned to at least 1 citation and (b) the pair of codes was present together in at least 1 citation. The actual number of distinct ICD-9 codes that were used in the pairwise comparisons in each dataset. Not all possible codes contributed to a pairwise comparison. ICD-9, International Classification of Disease, V.9 Table 2 The 30 most common codes for the Simplified versions of the three datasets

Simplified Clinical dataset Simplified Medline 1000 dataset Simplifie					ified Medline 600 dataset			
Code	Description	Freq.	Code	Description	Freq.	Code	Description	Freq.
786	Symptoms involving respiratory syststem/chest	323,562	780	General symptom	279,960	89	Interview, evaluation, consultation, and examination	684,766
780	General symptom	286,381	89	Interview, evaluation, consultation, and examination	168,522	780	General symptom	669,453
789	Symptoms involving abdomen, pelvis			Neoplasms of unspecified nature	164,581		Neoplasms of unspecified nature	491,658
719	Joint disorders	226,585	88	Other diagnostic radiology and related techniques	164,304	99	Other nonoperative procedures	452,538
427	Cardiac dysrhythmias	206,499	99	Other nonoperative procedures	151,387	88	Other diagnostic radiology and related techniques	425,485
V72	Special investigations and exams	195,803		Malignant neoplasm	134,119		Malignant neoplasm	370,794
518	Diseases, lung, other	183,058	250	Diabetes mellitus	116,196	279	Disorders of the immune mechanism	316,417
465	Acute infections of upper respiratory tract	171,964	401	Essential hypertension	101,952	E904	Accident due to hunger/thirst/exposure	311,337
V70	General medical examination	171,947	997	Complication affecting body	95,460	759	Congenital anomalies	254,442
724	Back disorders	152,960	402	Hypertensive heart disease	93,712	39	Operations on vessels	215,326
729	Disorders, soft tissues	151,388	405	Secondary hypertension	93,431	87	Diagnostic radiology	203,444
V04	Need for prophylactic vaccination	149,835	403	Hypertensive chronic kidney disease	93,270	042	HIV disease	193,453
V07	Need for prophylactic measures	148,123	404	Hypertensive heart and chronic kidney disease	93,148	250	Diabetes mellitus	187,375
787	Symptoms involving digestive system	141,223	338	Pain	80,803	695	Erythematous conditions	176,305
959	Injury, not otherwise specified	139,420	042	HIV disease	78,847	782	Symptoms involving skin, other tissue	172,965
V67	Follow-up examination	139,155	278	Overweight, obesity and other hyperalimentation	70,312	338	Pain	172,190
782	Symptoms involving skin, other tissue	138,882	429	III-defined heart disease	69,354	997	Complication affecting body	168,875
V06	Need for combination vaccination	135,206	427	Cardiac dysrhythmias	68,396	401	Essential hypertension	168,857
401	Essential hypertension	133,645	410	Acute myocardial infarction	67,988	799	Morbidity/mortality, ill-defined	162,701
784	Symptoms involving head and neck	129,325	E904	Accident due to hunger/thirst/exposure	67,891	92	Nuclear medicine	159,785
V76	Screening for malignant neoplasms	118,977	311	Depressive disorder	66,878	429	Ill-defined heart disease	158,653
V20	Health supervision of infant/child	114,588	787	Symptoms involving digestive system	66,615	402	Hypertensive heart disease	157,874
785	Symptoms involving cardiovascular system	113,977	414	Chronic ischemic heart disease	65,600	405	Secondary hypertension	157,495
733	Bone and cartilage disorders	112,034	782	Symptoms involving skin, other tissue	62,834	403	Hypertensive chronic kidney disease	157,374
599	Urethra/urinary tract disorders	102,919	39	Actinomycotic infections	61,703	404	Hypertensive heart and chronic kidney disease	156,992
367	Refraction/accommodation disorder	102,395	995	Adverse effects	59,973	368	Visual disturbances	155,964
709	Disorders of skin & sbcutn tissue	101,979		Symptoms involving respiratory system/	58,738	995	Adverse effects	155,379
V05	Need for prophylactic vaccination	98,882	300	Anxiety, dissociative and somatoform disorders	58,397	427	Cardiac dysrhythmias	149,674
530	Esophagus diseases	98,170	759	Congenital anomalies	56,048	268	Vitamin D deficiency	142,584
382	Suppurative otitis media	95,640	277	Unspecified metabolism disorder	55,589	269	Other nutritional deficiencies	142,075
Code	s that appear in the top 30 of the Clinica	dataset a	nd in th	e top 30 of least one of the Medline datasets	are highligh	nted to	show the concordance.	

Considering that many of the concurrent mentions of diagnoses in the same abstract were likely due to chance rather than as a result of true associations, we would have expected these false positives to have increased the amount of overlap. We did find a general trend that higher ranked clinical associations were more likely to also be found in Medline citations (figure 4), but this was not as pronounced as we had expected it to be. Even among the highest ranked clinical associations, and using the more inclusive Medline 600 data, only 27% of the clinical associations were also found in the Medline dataset (figure 4C). Some of this difference may be attributable to the fact that our dataset included historical, deprecated codes that are no longer identified or mapped by current systems such as MetaMap. An example from our previous work was code V72.3 (gynecological examination) which was no longer used after 2005, and was replaced by the more granular codes V72.31 and V72.32.4

Our clinical dataset had 87 420 patients with the older code V72.3, and V72.3 was present in 6005 associations. By contrast, there were 55 212 patients with the newer code V72.31 that was present in 5105 associations.

It is possible that many of the associations found in the patient care data are not clinically meaningful, or are related to one another due to confounding factors such as age or gender. And, many widely and historically known associations might not have been discussed in the literature indexed in Medline (the earliest citation in Medline is from 1809). There may also be an inherent bias in Medline because not every clinical association is necessarily an interesting research topic worth publishing, and the research literature does not necessarily discuss clinical conditions proportionally to their prevalence in the population. Other online resources such as Wikipedia might also provide relevant, supplemental clinical coverage not **Figure 5** Bar plots showing the relationship between the rank of an association in the Medline dataset and the probability of the association appearing in the Clinic dataset. (A) Medline 1000 (original) appearing in Clinical (original); (B) Medline 1000 (simplified) appearing in Clinical (simplified); (C) Medline 600 (original) appearing in Clinical (original); (D) Medline 600 (simplified) appearing in Clinical (simplified). The p-values shown represent the significance of the Medline associations at that percentile.



otherwise included in Medline. Further, our study only utilized titles and abstracts of Medline citations, leaving out the majority of details present in the main body of each publication.

We did find a trend that the lower-ranked associations tended to be more clinically surprising, with reasonable inter-rater agreement between the two human reviewers (tables 3 and 4). However, the low inter-rater agreement on whether or not citations were in support of the associations (table 3) demonstrates that identifying supporting scientific evidence is a challenge even among trained clinicians. Further, the fact that the reviewers were able to locate citations in support of 15 out of the 25 associations that were not identified in our Medline data (table 4) suggests that many relationships may be described in the literature in a manner that is not readily interpretable by computational tools such as MetaMap. While consolidating ICD-9 codes of the same disease category resulted in improved coverage, it might come with a price of potentially losing clinical meaning, especially when concepts that were combined should truly remain distinct. In the future, it may be beneficial to group codes according to their clinical relatedness, rather than hierarchically, as has been done for disorders such as stroke^{49 50} or depression.¹¹

Of course, there may be clinically valid associations that simply have not yet been reported in the literature, as we had discussed in our prior work.¹¹ As an example, table 4 exhibits a clinical association between vitamin D deficiency and nonspecific swellings or lumps. The citations we manually reviewed did not seem to directly support this association, but two of the citations did describe a potential relationship between lumps/ nodules and malnutrition/malnourishment,⁵¹ ⁵² the latter of which could result in a vitamin D deficiency.⁵³

It is also worth comparing our current study to other similar work. Holmes *et al*, for example, used MedLEE, another

medical NLP tool,⁵⁴ to extract concepts and map them to ICD-9 codes from Medline abstracts, Wikipedia articles, and discharge summaries for several rare diseases.⁵⁵ The study also used administrative data consisting of ICD-9 billing codes. This approach was able to identify associations found in the clinical data not found in the published literature, and vice versa. The authors also noted that ICD-9 is limited in its coverage of concepts, which likely limited their ability to detect additional associations, just as its use was likely a limiting factor of our study.

The literature-based discovery approach has also been used to identify other clinical relationships. For example, Vos et al^{20} used the lack of citations in Medline as a source of information to identify novel associations between psychiatric and somatic disorders. Experts reviewed candidates to determine which had clear explanations for their relatedness, or could not be readily explained (thus suggesting truly novel relationships). Another similar study extracted concepts from free text patient records and mapped them to ICD-10 codes in order to discover disease associations.¹⁹ The findings were then linked to the Online Mendelian Inheritance in Man (OMIM) resource which describes genetic disorders and their phenotypes.⁵⁶ An experienced clinician manually reviewed the top candidates to identify 'interesting' associations that were not previously known. In our study we only used ICD-9 codes from administrative data, but future work could include codes derived from clinical documents using NLP tools, as some other studies have done.

MeSH concepts from Medline have also been used to aid in the discovery of associations between clinical concepts.⁵⁷ For example, Avillach *et al*⁵⁸ used MeSH concepts to confirm associations with adverse drug events, defined as 'drug safety signals'. In the study, a threshold of an association being mentioned in three or more citations was used, whereas in our study we included even a single citation mentioning both diagnoses.

Table 3 Clinical associations also found in the 'Original' Medline 1000 dataset

CD-9 code	Description	ICD-9 code	Description	Association p value derived from Clinical dataset	Clinically surprising?	Number of Medline citations with both concepts mentioned	citations	Citation could b interpreted as describing an association?
lst percentile	(highest ranked associations)							
290.0	Senile dementia	331.0	Alzheimer's disease	<5×10 ⁻³²⁴	No	1139	11138345	No
							2614500	Yes
							7056516	No
344.61	Cauda equina syndrome with	596.54	Neurogenic bladder	<5×10 ⁻³²⁴	No	11	11880062	*
	neurogenic bladder						16813905	No
							3400548	No
184.4	Malignant neoplasm of vulva	233.3	Carcinoma in situ, unspecified female genital organs	<5×10 ⁻³²⁴	No	1	2082867	*
396.2	Mitral valve insufficiency and aortic	424.1	Aortic valve disorders	<5×10 ⁻³²⁴	No	2	11163732	Yes
550.2	valve stenosis	727.1		23/10	NO	2	9665226	Yes
362.21	Retrolental fibroplasia	769	Respiratory distress syndrome	<5×10 ⁻³²⁴	No	92	12709796	*
502.21		705	in newborn	< <u>J</u> ×10	NO	52	19568962	*
							15716610	*
5th percenti	le							
268.9	Unspecified vitamin D deficiency	782.2	Localized superficial swelling,	3.8×10 ⁻⁶⁸	Yes	3	11233710	No
200.9	(268.9)	702.2	mass, or lump	5.0×10	165	J	21806909	No
	(200.3)		mass, or rump				9339283	No
558.1	Gastroenteritis and colitis due to	789.0	Abdominal pain	7.1×10 ⁻⁶⁸	No	1	8140765	Yes
550.1	radiation	705.0		7.1.410	110		0140705	105
279.5	Graft-versus-host disease	E888.9	Accidental fall	2.4×10 ⁻⁶⁷	Voct	1	9250172	No
					Yes‡ *			No *
131.9	Trichomoniasis		Pyelonephritis	3.0×10 ⁻⁶⁷		1	9286064	
078.19	Viral warts	622.11	Mild dysplasia of cervix	8.4×10 ⁻⁶⁷	No	1	7559948	Yes
0th percenti	e							
276.52	Hypovolemia	403.9	Hypertensive renal disease	9.8×10 ⁻²³	No	73	11602456	Yes
							17113396	Yes
							365408	No
255.41	Glucocorticoid deficiency	788.3	Functional urinary incontinence	4.8×10 ⁻²²	Yes	1	9836036	Yes
413.9	Angina pectoris	596.51	Hypertonicity of bladder	7.0×10 ⁻²²	Yes	1	17689623	No
286.9	Coagulation defect	344.00	Quadriplegia	8.0×10 ⁻²²	*	4	11403538	No
	-						16958632	Yes
							3508705	No
229.9	Benign neoplasm of unspecified	354.0	Carpal tunnel syndrome	4.9×10 ⁻²¹	Yes	1	1672719	*
	site							
5th percenti	le							
416.9	Chronic pulmonary heart disease	783.41	Failure to thrive	4.6×10 ⁻⁶	No	24	12597677	Yes
	, , ,						8165079	Yes
							2657582	Yes
351.0	Bell's palsy	742.59	Congenital anomalies of spinal	4.7×10 ⁻⁶	No	1	18756840	No
			cord					
302.72	Psychosexual dysfunction with	576.2	Obstruction of bile duct	4.6×10 ⁻⁶	Yes	1	16402030	No
	inhibited sexual excitement							
259.9	Endocrine disorder	368.00	Amblyopia	4.6×10 ⁻⁶	Yes	1	15105955	Yes
611.72	Lump or mass in breast	759.6	Hamartoses	4.7×10 ⁻⁶	No	1	19737912	*
	tile (lowest ranked associations)							
571.2	Alcoholic cirrhosis of liver	617.9	Endometriosis	0.99	Yes	1	8834254	*
153.9	Malignant neoplasm of colon	487.1	Influenza with other respiratory	0.99	Yes	3	12889684 18544745	No
			manifestations				20813181	No No
454.0	Verience using of lower states to	601.0	Other stepie downstitie and	0.00	Vee	1		
454.0	Varicose veins of lower extremities with ulcer	691.8	Other atopic dermatitis and	0.98	Yes	1	3442079	No
F0F C		626.0	related conditions	0.00	N	22	2120005	N
585.6	End stage renal disease	626.0	Absence of menstruation	0.99	No	23	3130865	Yes *
							16619340 9593608	Yes
							7172000	165

These were selected from the area of overlap in figure 3A. The lower-ranked associations tend to be more clinically surprising. Many of the citations found in the literature did not actually suggest a true association after manual review. Whether an association was clinically surprising was independently determined by two physicians (κ statistic 0.84; 95% CI 0.63 to 1.05). The κ statistic for agreement on whether the citations found by our approach supported the association was 0.55 (95% CI 0.30 to 0.79).

+For associations with more than three citations, three were randomly selected for this analysis.

*The abstract stated, 'donor stem cells become tolerant to host antigons and fall to cause GVHD'. The word 'fall' was coded into the ICD-9 code for a fall. Not only was this the wrong context for a fall, but the word 'fall' in this abstract is actually a typographic error and should have been 'fail'. ICD-9, International Classification of Disease, V.9; GVHD, graft-versus-host disease; PMID, PubMed Identifier.

Table 4 Clinical associations not found in the Original Medline 1000 dataset

ICD-9 code	Description	ICD-9 code	Description	Association p value derived from clinical dataset	Clinically surprising?	Citations found (PMID)†
1st percentile	e (highest ranked associations)					
719.46	Joint pain, lower leg	848.9	Sprain and Strain, unspecified site	<5×10 ⁻³²⁴	No	21549978 9343643
172.1	Malignant melanoma of skin and eyelid	190.3	Malignant neoplasm of conjunctiva	<5×10 ⁻³²⁴	No	21478094 10811089
250.71	Diabetes with peripheral circulatory disorders	337.1	Peripheral autonomic neuropathy	<5×10 ⁻³²⁴	No	20724598 2779736
153.0	Malignant neoplasm of colon	230.4	Carcinoma in situ of rectum	<5×10 ⁻³²⁴	No	21125511 6894080
309.28	Adjustment disorder with mixed anxiety and depressed mood	724.2	Lumbago	<5×10 ⁻³²⁴	No	21665125 18673099
25th percenti	le					
535.50	Gastritis and gastroduodenitis	787.6	Fecal incontinence	4.8 x10 ⁻⁶⁸	*	16712555
518.0	Pulmonary collapse	876.1	Open wound of back	9.8×10 ⁻⁶⁸	No	17554992
474.0	Chronic tonsillitis and adenoiditis	558.9	Non-infectious gastroenteritis and colitis	7.6×10 ⁻⁶⁷	Yes	12080166
377.39	Optic neuritis	432.1	Subdural hemorrhage	7.8×10 ⁻⁶⁷	Yes	
307.59	Eating disorder	564.01	Slow transit constipation	9.9×10 ⁻⁶⁸	No	19139750 10925980
50th percenti	le					
600	Hyperplasia of prostate	747.61	Gastrointestinal vessel anomaly	1.89×10 ⁻²³	Yes	
296.20	Major depressive affective disorder	734	Pes planus	3.5×10 ⁻²³	Yes	
164.8	Malignant neoplasm of mediastinum	427.89	Cardiac dysrhythmia	3.7×10 ⁻²²	No	15284266 21387697
227.0	Benign neoplasm of adrenal gland	788.41	Urinary frequency	5.3×10 ⁻²²	Yes	
250.51	Type 1 diabetes with ophthalmic manifestations	959.5	Finger injury	1.9×10 ⁻²¹	*	18820219
75th percenti	le					
217	Benign neoplasm of breast	569.85	Angiodysplasia of intestine with hemorrhage	4.7×10 ⁻⁰⁶	Yes	
362.31	Central retinal artery occlusion	836.0	Tear of medial cartilage or meniscus of knee	4.7×10 ⁻⁰⁶	Yes	
373.32	Contact and allergic dermatitis of eyelid	656.90	Unspecified fetal and placental problem, affecting management of mother, unspecified as to episode of care or not applicable	4.7×10 ⁻⁰⁶	Yes	
110.5	Dermatophytosis of the body	246.2	Thyroid cyst	4.7×10 ⁻⁰⁶	Yes	1607406
011.90	Pulmonary tuberculosis	701.1	Keratoderma, acquired	4.8×10 ⁻⁰⁶	Yes	9828554
100th percen	tile (lowest ranked associations)					
459.9	Circulatory system disorder	684	Impetigo	0.99	*	22642914
171.9	Malignant neoplasm of connective and other soft tissue	333.1	Essential tremor	0.99	Yes	
189.0	Malignant neoplasm of kidney	795.5	Nonspecific reaction to test for tuberculosis	0.99	Yes	20623161
225.3	Benign neoplasm of spinal cord	558.9	Non-infectious gastroenteritis and colitis	0.99	Yes	
378.31	Hypertropia	722.10	Displacement of lumbar intervertebral disc without myelopathy	0.99	Yes	

These were selected from the left-most area in figure 3A. Associations become more clinically surprising as their ranking decreases. Whether an association was clinically surprising was independently determined by two physicians (κ statistic 0.75; 95% Cl 0.48 to 1.01). A manual search for citations by each physicians revealed potential associations that were not detected with the automated approach using MetaMap.

*Opinions for which the physicians differed.

tlf only one citation is listed, it means that only one reviewer found a citation that supported the association. If two are listed, each reviewer found at least one citation to support the association. If none are listed, neither reviewer was able to find a citation to support the association.

ICD-9, International Classification of Disease, V.9; PMID, PubMed Identifier.

Other approaches could also be used to improve the performance of similar data mining methods, some of which could be applied when the original data are collected. For example, if authors were given the option to manually annotate Medline citations with additional coded data (eg, diagnoses, procedures, drugs), the need for complex post-hoc NLP could be diminished. Additionally, development of a curated knowledge base of known associations could be useful in a variety of contexts. Ontologies could also be leveraged to find associations that may not be explicitly defined but may be discoverable through the ontological relationships. There are a multitude of association measures, ⁵⁹ and selecting the right measure to capture a subjective notion of 'interestingness' is a research topic of future investigation. Association measures differ in what they are aiming to capture but many of them use the same quantities, for example, marginals, conditionals, and sizes of intersections.

It is possible that the validity of our study findings is contingent on the performance (ie, accuracy and relevance) of MetaMap. Processing free text is complex and MetaMap, similar to many other NLP-based named entity extraction systems, can introduce systematic errors in its classification of clinical concepts.²⁸ ^{60–63} One recent study, for example, reported an F-measure of 61% when MetaMap was applied to a biomedical corpus including Medline abstracts.⁶⁴ Nevertheless, with the continued support from the NLM, the performance of MetaMap is expected to improve over time,²³ and in certain use scenarios it has been demonstrated that MetaMap outperforms human annotators.⁶⁵ Further, there have been studies showing that MetaMap is able to identify a broader range of concepts than other NLP systems.⁶⁶

It is also worth pointing out that in our experiments we were not attempting to compare the coding accuracy of MetaMap to other named entity recognition systems, nor were we trying to compare the accuracy of MetaMap to the accuracy of ICD-9 coding in clinical encounters. The goal of converting the concepts in the Medline citations into ICD-9 codes was to have a common coding framework with which to compare both the clinical and Medline datasets. Named entity recognition and ICD-9 coding are very different tasks, and often follow different 'rules' for converting text into their respective coded counterparts. The clinically assigned ICD-9 codes themselves represent an abstraction of the diagnoses in clinical text and it may not always be the case that the levels of granularity of codes from a clinical dataset would match those identified from the literature. We attempted to address the potential granularity issue by merging the codes in our 'Simplified' datasets. Still, it is important to note that the two datasets were not created with the intention of being compared in this manner, and the base entities (abstracts vs patients) from which the data were extracted are also not directly comparable.

The identification of falls related to graft-versus-host disease (GVHD) due to a typographic error ('fail' \rightarrow 'fall') in an abstract we reviewed (table 3) demonstrates that more work is warranted to accurately identify concepts in the correct context. Yet, this spurious literature support does not rule out the existence of this relationship, which was found in our Clinical dataset. Indeed, patients with GVHD can experience balance impairments⁶⁷ which could, in turn, result in falls, even if this association is not explicitly mentioned in the citations we analyzed. This general assertion is supported by the results shown in table 4 which demonstrate that in multiple cases experienced clinicians could identify citations supporting the clinical associations while the automated approach was unable to.

In the experiments reported in this paper, we used two distinct score thresholds (1000 and 600) to process the results generated by MetaMap. The use of the MetaMap score of 1000 may have been too restrictive, not allowing the system to detect variations of a concept, and could have excluded many potential concepts that were present in the Medline citations. This is why we also used the threshold of 600, which was lower than scores commonly used in prior studies, to make sure the results achieved were more inclusive. One risk of reducing the threshold to 600 is a higher likelihood of incorrect text classifications yielding more clinically irrelevant associations. Yet, even with that potential loss of accuracy, there were still many clinical associations not identified in the Medline dataset.

In prior literature, studies have either not reported the specific MetaMap scores used²⁴ ²⁸ ⁶⁸ ⁶⁹ or have reported seemingly arbitrary thresholds for including concepts in their work. Thresholds of 700,⁷⁰ 800,^{71–73} 850,⁷⁴ 900,⁷⁵ and 950⁷⁶ ⁷⁷ have all been used in the past, as well as perfect scores of 1000.⁷⁸ ⁷⁹ Further, some studies have used the highest ranking score

among all candidate concepts for inclusion, with no mention of a minimum threshold.^{80–82} While it has been stated that the threshold can 'usually be determined simply by examining MetaMap output for typical text in a given application',⁸³ there appears to be no consensus about the optimal MetaMap score above which results should be retained. This lack of a consensus may affect the generalizability of research using such tools, including our current study.

CONCLUSION

This work demonstrates the potential utility but persisting challenges of using large biomedical knowledge repositories for identifying novel relationships derived from clinical datasets. At a broad scale, additional filtering approaches will likely be needed to reduce the size of the set to a reasonable number for expert review, and the addition of other resources beyond Medline could help to distinguish novel associations from wellknown ones. Improved NLP and concept extraction capabilities would also be expected to play a significant role in improving the performance of such an approach. Informatics researchers should consider the implications of these promising, but potentially limited, data mining approaches when exploring 'big data', and how the findings should be presented and interpreted.

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