Big Dreams With Big Data! Use of Clinical Informatics to Inform Biomarker Discovery

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As the complexity of biomedical data increases, so do the opportunities to leverage them to advance science and clinical care. Electronic health records form a rich but complex source of large amounts of data gathered during routine clinical care. Through the use of codified and free-text concepts identified using clinical informatics tools such as natural language processing, disease phenotyping can be performed with a high degree of accuracy. Technologies such as genome sequencing, gene expression profiling, proteomic and metabolomic analyses, and electronic devices and wearables are generating large amounts of data from various populations, cell types, and disorders (big data). However, to make these data useable for the next step of biomarker discovery, precision medicine, and clinical practice, it is imperative to harmonize and integrate these diverse data sources. In this article, we introduce important building blocks for precision medicine, including common data models, text mining and natural language processing, privacy-preserved record linkage, machine learning for predictive modeling, and health information exchange.

INTRODUCTION

Adoption and meaningful use of electronic health records (EHRs) has continued to increase, spurred by federal mandates in the United States. These electronic systems collect vast amounts of clinical data either as structured elements (vital parameters, laboratory data, etc.) or unstructured clinical notes. In addition, these data are intended to facilitate effective clinical decision support (CDS), as defined by HealthIT.gov (1) as systems or processes that “(provide) clinicians, staff, patients or other individuals with knowledge and person-specific information, intelligently filtered or presented at appropriate times, to enhance health and health care.” These data, currently used primarily for clinical care and administrative purposes, hold tremendous potential for advancing biomarker discovery and personalized medical decision-making at point of care in patients with chronic diseases like inflammatory bowel diseases (IBD).

In parallel with widespread EHR adoption, there have been tremendous advancements in computational biology techniques with proliferation of standardized genetic platforms and sequencing technologies, explosion of multomics approaches, along with streamlined analytic pipelines, facilitating pooling of research data across populations. However, such efforts have relied on carefully curated cohorts with research teams manually identifying patients from clinical care by review of individual charts to identify eligible individuals, which requires significant personnel support and is resource intensive. Moving forward, utilizing the EHR to curate large disease-based cohorts in a short amount of time with modest resources, carefully performing automated detailed disease phenotyping utilizing text mining and natural language processing (NLP), and then integrating these diverse “big data” sources through privacy-preserved linkage can promote effective and efficient discovery research, rapid translation and integration, and adoption at point of care. In this chapter, we discuss important concepts of clinical informatics, a rapidly evolving field at the cross-section of information technology and healthcare, required to facilitate such advancement. Figure 1 summarizes the approach to precision medicine using EHRs.

COMMON DATA MODELS

An intrinsic limitation to any big data approach is the issue of data quality in terms of volume, variety, velocity, and veracity (2–4). Hence, to make EHR data usable across formats and institutions, it is critical to develop a common data model with the use of standard terminology. Each type of data has an associated terminology that enables the vocabulary to be operationalized within the context of the EHR. These terminology systems have unique data formatting, coding, domain coverages, and hierarchical relationships between a specific instantiation, such as amoxicillin capsule 250 mg, and a concept, such as penicillin.

Table 1 shows the common EHR data sources relevant to CDS. Precision medicine is developing a new vocabulary related to genetic conditions, which has yet to be standardized in the EHR. Genetic test results should follow relevant data standards, such as Logical Observation Identifiers Names and Codes, Health Level 7 Genomics, Human Genome Variation Society, that contain information about test findings and potential risk; yet, this is a challenge since these standards are not adopted by all laboratories. The rapid evolution of tests makes this challenging for the field of genetics, posing challenges for discrete data retrieval of
Figure 1. Key tools for precision medicine using electronic health records.

Text mining and natural language processing

While several elements handled through common data models are based on structured or codified elements, free text or narratives still dominate in terms of clinically relevant information contained in EHRs. While free narrative is effective and convenient for medical record keeping, its unprocessed form is difficult to search, summarize, or analyze for secondary purposes such as research or quality improvement. NLP is any computer-based algorithm that handles, augments, and transforms natural language so that it can be represented for computation. Because a computer cannot comprehend meaning from a block of text, a series of operations must be defined to transform the data into usable information, which is the essence of NLP. In elegant use of this combination of codified data and NLP to develop an EHR-based cohort, Ananthakrishnan et al. created a cohort of 11,000 patients with IBD within 2 hospitals in Boston (4–6). From among all patients with at least 1 billing code for Crohn’s disease or ulcerative colitis, a chart review revealed a positive predictive value of only 60% with frequent misclassification. Extraction of codified data ascertaining disease complications as well as narrative free text data comprising number of mentions of individual disease names (“Crohn’s disease”) or disease-related terms in clinical notes (“abdominal pain”, “diarrhea”), radiology reports (“ileal wall thickening”), endoscopy (“ileitis” “apthous ulcer”), and pathology (“crypt abscess”) allowed for development of a classification algorithm using machine learning that was able to achieve a positive predictive value of 97%. The addition of free-text data to the codified information not only improved the accuracy of identifying cases but also increased the number of patients who could be classified as having disease. In addition, this approach also allowed identification of phenotypes of disease, such as primarily sclerosing cholangitis, which is limited by lack of specific diagnostic codes or high frequency of use of codes for competing diagnosis (e.g., cholelithiasis), determining status of disease activity in relapsing and remitting disorders, or identifying response to treatment. NLP software is increasingly sophisticated to be able to distinguish positive findings (“has diarrhea”) from negative ones (“does not have diarrhea”), assign specific contexts for occurrence of phrases (“abdominal pain” from “joint pain”), separate personal from family history (“family history of colon cancer”), and search within specific components of the note (such as indication for procedures). Despite the inherent variability in structure and content of EHR data and differences in quality of provider documentation across institutions, disease-defining algorithms created at one institution are portable to other institutions using distinct EHRs and retain their accuracy, which is key for multi-institutional consortia, such as the Electronic Medical Records and Genomics Network. With advances in the field of NLP, detailed phenotyping is feasible, allowing performance of large scale, integrated genome- and phenome-wide studies to promote biomarker discovery and precision medicine.

Privacy-preserved record linkage

One of the greatest challenges to utilizing big data for research has been data partitioning from diverse sources. In a research network, information from the same individual may be partitioned among several sites such as healthcare providers, sequencing facilities, insurance companies, research institutions. There are mainly 2 types of patient data partitioning across institutions: (i) horizontal partitioning, where different institutions hold information on the same and (ii) vertical partitioning, where different institutions hold information on different attributes. The former one consists of records with the same features, for an overlapping or nonoverlapping set of individuals. Feature values are the same in the case of true overlap, or they can differ when patients switch healthcare systems or receive complementary care in different health systems (e.g., patients cared for primarily at the Veterans Health Administration system but receiving specialty care in another system). In vertical partitioning, there is information about different features for the same individual at different sites. In both situations, patient record linkage is an essential step to combine data in cross-institutional studies. For example, if the truly duplicated records across different institutions cannot be sufficiently removed, the estimation could become biased in the study with horizontally partitioned data. For the case of vertically partitioned data, the genome data of a particular group hosted in a sequencing facility can be significantly enriched by linking the data to EHRs. In addition to linking patient records across research networks, the existing clinical data research networks can link their data to publicly available databases of vital statistics (such as the National Death Index), pharmaceutical databases, etc., allowing comprehensive and simultaneous capture of multiple exposures, health status, interventions, and outcomes. The existing record linkage methods can be categorized into 2 approaches: deterministic and probabilistic (7). If there exist explicit identifiers (e.g., name and social security number) among different datasets, deterministic record linkage methods are used. Probabilistic linkage methods are more complex, as they assign different weights for different discriminative linkage variables to compute an overall score that indicates how likely it is that a record pair comes from the same patient. Furthermore, due to concerns of invasion of privacy, institutions and patients alike may be hesitant to share personal health
| Type of information | Standardization | Opportunities | Challenges |
|---------------------|-----------------|---------------|------------|
| Laboratory          | LOINC, HGVS, HL7 FHIR value sets | Clinical laboratory tests have a mature standardization capabilities via LOINC. LOINC and HL7 genomics groups have started developing standards for genetic tests—that enable standardized discrete coding of some genetic test information | - Not all clinical laboratory tests are encoded with LOINC (still in process in many institutions)  
- Discussions on including genetic text in EHR in a structured way have only recently commenced  
- Significant volumes of tests are performed at external laboratories with processes and results that lack standardization.  
- Laboratory orders are frequently matched in the computer to component results.  
- Genetic test results are not systematically incorporated into EHR in a searchable way. For example, they are nondiscretely stored in the EHR as a scanned PDF document or image at the UCSD Medical Center |
| Medication          | RxNorm, NDC     | Clinical drug names have been standardized using these codes. Dictionaries provide the opportunity to include manufacturer, dosing, and route information | - Categorization is not clean as medications may have multiple indications both on and off label that skew groupings  
- Combination drugs may not neatly fit into clinical groupings  
- Deriving relevance related to effect over time, dosing intensity, or adherence are problematic |
| Diagnosis           | ICD 9, ICD 10, SNOMED-CT | Most institutions adopt ICD system to support both active problem lists and encounter diagnoses. Diagnosis names are interrelated, meaning that terms encoded with other one terminology such as SNOMED-CT, can be converted to ICD through cross-mapping established between the two systems | - Coding is frequently completed by a clinician with time constraints that may not search through the extensive terms for the true best fit (undercoding, miscoding)  
- ICD9 and ICD10 contain level of detail that may deviate from clinical relevance  
- ICD9 is historic and ICD10 current (codes expire and newly develop)  
- Not all codes are billable (irrelevant)  
- Some diagnoses are not encoded (missing)  
- SNOMED concepts are frequently not parsed into terms that support clinically specific workflows  
- IMO updates can impact term groupings and insert clinically mismatched concepts |
information outside the health system. Hence, robust privacy-preserving record linkage tools are clearly needed before this rich environment is ripe for research use. Figure 2 depicts an example of a record linkage system for vertically partitioned data between a hospital and a biobank where DNA data are available.

Health information exchange

One of the limitations of EHR-based research is that data are contained in silos in health systems, which do not interact adequately with each other. While patients move in and out of health systems, their data do not move and get lost in translation. Not only does this disrupt clinical care, but it also impedes phenome-wide association studies and biomarker discovery due to misclassification of clinical data. However, through approaches of health information exchange (HIE), a special case of privacy-preserved record linkage, this barrier may be overcome (2). The Health Information Technology for Economics and Clinical Health Act of 2009 was proposed to promote interoperable health information. HIE initiatives aim at realizing timely and appropriate level of access to the patient level of health information stored in the EHR by healthcare providers through a secure means to exchanging health data among healthcare organizations. Having complete information about disease progression and treatment data at the point of care helps healthcare providers make better treatment decisions and achieve better patient outcomes. Utilizing information collected from different healthcare systems is an important step toward this goal.

HIE covers 3 types of data exchange: (i) Directed exchange that occurs between healthcare providers to complete the planned

| Table 1. (continued) |
|----------------------|-----------------|-----------------|-----------------|
| Type of information  | Standardization | Opportunities | Challenges |
| Radiology            | RadLex, SNOMED-CT DICOM | Standards to capture the key findings and metadata about the radiologic studies exist | • Radiology test related metadata may not be formatted in a structured way using a standard like DICOM  
• Radiology reports are in an unstructured narrative text format. Processing the text to tease out the key findings and mapping them to the standardized codes requires additional efforts/resources that involves NLP |
| Pathology            | SNOMED-CT HL7 (anatomic pathology) | Standards to capture the key findings and metadata about the pathology test exist NAACCR is interested in adopting standard for cancer pathology reporting | • Pathology reports are in an unstructured narrative text format or PDF. Processing the text to tease out the key findings and mapping them to the standardized codes requires additional efforts/resources (NLP)  
• Pathology frequently utilizes standardized nomenclature but does not record data in structured format |
| Clinical evidence and outcomes | OMOP CDM and all terminology systems listed above | EHR data stored in a clinical data warehouse serve a powerful knowledge resource OMOP CDM is recognized as a de facto standard and adopted by many institutions | • There are types of data that are not sufficiently represented by the OMOP CDM such as patient reported outcomes  
• OMOP has not been universally adopted across organizations |
| Procedures           | Terms to represent clinical procedures | Standardized terms that define common clinical procedures and their associated charges | • Process for approving new procedural codes is onerous as a result the library may incompletely represent activity detail  
• Many procedural codes are fairly generic and do not incorporate the level of details that impact outcomes |

CDM, common data model; DICOM, digitalized imaging and communications in medicine; EHR, electronic health records; FHIR, fast healthcare interoperability resources; HGVS, Human Genome Variation Society; HL7, health level 7; ICD, international classification of diseases; IMO, international medical objects; LOINC, Logical Observation Identifiers Names and Codes; NAACCR, North American Association of Central Cancer Registries; NDC, national drug code; NLP, natural language processing; OMOP, Observational Medical Outcomes Partnership; SNOMED-CT, standardized nomenclature of medicine - clinical trials; UCSD, University of California San Diego.
healthcare services such as sending and receiving laboratory test orders and results, and exchanging patient referral documents, (ii) **Query-based exchange** that occurs when a healthcare provider delivers unplanned services and requires accessing necessary health information about the patient. For example, when an emergency room physician needs to access patient’s disease history, current medications, allergies, etc. (iii) **Consumer-mediated exchange** that lets patients control their health information. In this model, patients grant access to their health information to healthcare providers. However, establishing a sustainable HIE is not a trivial task; there are a number of technical and nontechnical barriers that need to be addressed first. For example, lack of business incentives, specifically concerns on losing patients to other hospitals by making their health data available anywhere, has long been recognized as a factor that makes some healthcare systems hesitant to embrace HIEs. Patients and providers sometimes opt out from HIEs due to privacy concerns. Other recognized challenges are poor data standardization, inefficient processes of sorting through overloaded unselective information of a patient, and difficulties in understanding the shared data in the absence of context when detailed clinical notes are withheld due to privacy concerns.

**Statistical approaches including machine learning**

With this vast amount of data being generated from diverse sources, novel and powerful analytic approaches are needed. Figure 3 summarizes different approaches to analysis. Machine-learning methods consist of computational algorithms to relate all or some of a set of predictor variables to an outcome (8). To estimate the model, they search, either stochastically (randomly) or deterministically, for the best fit. This searching process differs across the different algorithms. However, through this search, each algorithm attempts to balance 2 competing interests: bias and variance. In the machine-learning context, bias is the extent to which the fitted predictions correspond to the true values—that is, how accurately does the model predict the “true” risk of death.
in the population? Variance is the sensitivity of the predictions to perturbations in the input data, that is, how does sampling variability impact the predictions? Even though it is not possible to separately quantify a model’s bias and variance, these 2 values are summarized together by loss functions. Many machine-learning methods can be grouped into different families based on their underlying structure. The 2 largest families are those that amend the traditional regression model (such as regularized methods, including common ridge regression and Lasso) and tree-based methods (such as classification and regression trees), and others including artificial neural networks, nearest neighbors, and support vector machines.

In summary, marrying EHR-based clinical research approaches with advancements in computational biology is immensely promising for biomarker discovery and promoting personalized medical decision-making. One can readily envision this approach being applicable across a wide swath of diseases relevant to gastroenterology, including IBD, chronic liver diseases, and gastrointestinal cancers. All of these diseases have in common varying, and often poor, accuracy of the existing administrative coding-based diagnoses, but can be readily identified in the EHRs using data (e.g., serology, pathology, and endoscopy) that are a routine part of clinical care and that can be mined using clinical informatics tools. Linkage of such disease registries to bio-banked genotyped samples, ensuring appropriate data protection and de-identification, can be enormously valuable to advance scientific discovery. This, however, is contingent on standardization of reporting methods and attributes and the ability to receive structured data from outside sources. Once these data are standardized across formats, it can readily be used to populate specific patients attributes in integrated CDS tools directly and seamlessly promoting personalized medical decision-making at point of care.

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