



The Evolving Use of Electronic Health Records (EHR) for Research



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Electronic health records (EHR) have been implemented successfully in a majority of United States healthcare systems in some form. There has been a rise in secondary uses of EHR, especially for research. EHR data is large, heterogenous, incomplete, noisy, and primarily created for purposes other than research. This presents many challenges, many of which are beginning to be overcome with the application of computer science artificial intelligence techniques, such as natural language processing and machine learning. EHR are gradually being redesigned to facilitate future research, though we are still far from a "complete EHR."

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Introduction: The Rise of EHR in the United States

Electronic health records (EHRs) are, in the most general sense, clinical information systems that collect, store, and present longitudinal electronic data collected during the delivery of health care. The field of radiation oncology, in particular, has benefited from advancements in treatment planning and delivery systems, which can be considered a special type of EHR. After decades of trying to move from a paper-based record system to an electronic one, attention is now turning away from implementation/adoption and moving toward realizing greater benefits from digital records. This article will focus on the evolving impact of EHRs on research in the United States (US).

The last several decades have seen an expanding role for EHRs. The 1960s and 1970s saw development of the first clinical information systems, such as The Medical Record at Duke University, the Computer Stored Ambulatory Record

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at Massachusetts General Hospital, and Health Evaluation through Logical Processing at Intermountain Healthcare in Utah.² In the early 1970s, the Department of Veterans Affairs developed an EHR, now known as VistA, which remains in use today at the Veterans Health Administration. In the 1980s, the National Academy of Medicine (formerly known as the Institute of Medicine [IOM]) began to analyze paper records and EHRs, and ultimately released reports in 1991 and 1997 that made the case for more widespread EHR adoption in the US healthcare system.³ In a subsequent study of medical errors, the IOM concluded that computerized order entry was likely to improve patient safety.^{3,4} These IOM reports also delineated 8 core functionalities of the EHR (Table 1)⁵ and identified barriers to widespread EHR adoption in the US.³

Despite this, EHR adoption rates remained low through the 2000s. Although 48% of office-based physicians reported using any EHR-based system in 2009, a survey published that same year indicated that only 1.5% of US hospitals had a "comprehensive" EHR, 7.6% had a basic system, and 17% had computerized provider-order entry. 6,7 Costs of EHR acquisition and maintenance were identified as the major barriers to EHR adoption. In response to this, the HITECH Act, passed as part of the American Recovery and Reinvestment Act of 2009, provided increased federal funding for health IT and established incentives for "Meaningful Use" of EHR technology.8 President Obama's Precision Medicine Initiative contributed to raising interest in and funding for developing EHRs. Industry-developed systems such as those produced by Epic and Cerner, which focused on optimizing reimbursement, came into widespread use during this time period as well. In response to these and other developments,

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Table 1 Core Functionalities of Electronic Health Records (EHR) Defined by the National Academy of Medicine (Formerly Known as the Institute of Medicine [IOM])

| | Core Functions | Details |
|----|---|--|
| 1. | Health information and data | Key information about patients including diagnoses, allergies, medications, lab results, etc. Needs to evolve over time, as new clinical knowledge becomes available and users' needs change. |
| 2. | Result management | Manage results of all types, including lab test results, radiology procedure results, etc. Facilitates efficiency, improves patient safety, reduces redundant testing, aid communication among providers and between providers and patients. |
| 3. | Order entry/ management | Improves workflow processes by eliminating lost orders and ambiguities caused by illegible handwriting, automates creation of related orders, monitors for duplicate orders, reduces the time to fill orders, and improve clinician productivity. |
| 4. | Decision support | Aid providers in disease diagnosis and management. Improve adherence to evidence- based consensus guidelines and protocols. |
| 5. | Electronic communication and connectivity | Facilitate communication among care partners (eg, lab, pharmacy, radiology), enhancing patient safety and quality of care. Improve public health surveillance. |
| 6. | Patient support | Additional support for patient education. May extend to telehealth. |
| 7. | Administrative processes and reporting | Improve efficiency by reducing delays and confusion for patients; improve access to services with immediate validation of insurance eligibility; more timely payments and less paperwork; faster communication of drug recalls; identify candidates for chronic disease management programs. |
| 8. | Reporting and population health management | Represent clinical data with standardized and manipulatable data format to reduce costs associated with reporting key quality indicators. |

EHR utilization has increased dramatically — from 2008 to 2015, usage of any type of EHR increased from 9% to 96% for hospitals, and from 17% to 78% for physician offices. ¹⁰

The Primary Intended Purpose of EHR: Billing

As hospitals and ambulatory care centers are increasingly adopting EHRs, most user-level utilization remains focused on billing. The majority of EHRs in the US were designed to support billing, with clinical work flow and now research as secondary applications. In one single-center study, the majority of physicians accessed the EHR to enter data relevant for the purposes of billing (such as clinical documentation), but only a small minority edited problem lists, allergy lists, or suggested changes to the system. 11 Another study demonstrated that, although there is significant heterogeneity in the percentage of providers who use EHR functionalities like clinical decision support tools, and Meaningful Use objective measures that are critical for obtaining optimal reimbursement are fulfilled for virtually all clinical encounters. 12 A significant proportion of the total time and financial cost associated with EHR-based encounters (eg, as high as 25% for emergency department encounters) is related to billing. 13 Diagnosis and procedure billing codes are usually available for clinical encounters, and are thus a robust source of data that can be mined from EHRs for research purposes.

EHR Secondary Purposes

Secondary use of EHRs, including research and teaching, applies to the use of personal health information for uses outside of direct health care delivery.¹⁴ Ironically, this so-called

secondary use was the primary reason that medical records were originally kept, which was for didactic purposes. ¹⁵ Because research was not the purpose for data creation and the EHR includes sensitive information, research can raise medical ethical, political, technical, and social issues. ¹⁶ For example, we recently showed that private patient—provider and provider—provider communications in the EHR can predict for early treatment discontinuation. ¹⁷ As the volume of and access to health data increases in both the public and private sectors, it is becoming critical to address these issues. The American Medical Informatics Association (AMIA) published a White Paper ¹⁴ describing the need to develop a framework for the secondary use of EHR data, and later a data quality ontology ¹⁸ based on Weiskopf's 5 dimensions of data quality. ¹⁹

The secondary applications of EHR are growing as rapidly as the volume of data collected. First, and perhaps most obvious, is their use in more easily automating commonly performed clinical tasks. The use of EHR has made it significantly easier to gather and transfer a vast array of patient data, and the use of EHR may even allow for easier prospective data collection. The retrospective collection of data can be timeconsuming, error-prone, and may not be suitable for the purposes of clinical trials. EHR can be linked with electronic health recording instruments in order to more robustly track outcomes such as patient toxicity.²⁰ For instance, continuous heart rate monitoring could track and more rapidly detect dehydration from poor oral intake of head and neck cancer patients undergoing chemoradiation therapy, leading to more rapid interventions and reducing hospitalizations. The recent announcement that Fitbit data will be included as part of the All of Us program's data stream is emblematic of the evolution of such data "mash-ups." There have also been early efforts to capture serial patient-reported outcomes and automatically record them in the EHR.2

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Although a great deal of data is housed in the EHR, it is in various and often incompatible formats. The most common formats are, in decreasing order of utility: (1) structured and terminology encoded; (2) structured with local codes or no encoding; (3) unstructured machine-readable text; (4) unstructured scanned text (eg, faxes, PDFs). Radiology and pathology reports, for example, are in machine-readable free-text and generally require natural language processing (NLP) approaches to access the data. Although this data would ideally be prospectively coded in a format that could be easily analyzed in the future, this is not the current reality, and automated methods of retrospectively transcoding and extracting data are needed. Systems that are capable of analyzing and robustly extracting clinical data are of significant value, especially in the current era of widely variable standards for data storage, as well as the multiple EHR systems present in most radiation oncology departments (Table 2).²³

There is no doubt that with the increasing amount of data available to clinicians in the EHR, novel analysis tools may be needed to best make sense of the large amount of data available and to find associations that may not be obvious at first. This is even more apparent in the realm of radiation oncology, where treatment and prognosis not only depend on traditional clinical factors (eg, age, stage, comorbidity, and pathology), but also a wide variety of radiographic and dosimetric data. Furthermore, this data is likely spread across multiple EHR systems for primary records, pathology, radiology, and radiation oncology. The enormous amount of data available in the modern EHR provides a new environment for the application of machine learning (ML) tools and techniques utilizing artificial intelligence. Traditionally, clinicians have been limited to developing nomograms and scoring systems that may only take into account a handful of clinical factors, or be limited in their scope in the interest of ease of use. The use of more sophisticated ML methods may allow for more robust predictive models to be generated using non-traditional variables. These models could ultimately be integrated into the EHR for rapid clinical application.²⁴⁻²⁶

Challenges of Using EHR for Research

There are many challenges of using EHR in research: (1) data quality and validation; (2) complete data capture; (3) heterogeneity among systems; and (4) system knowledge; Cowie et al. describe several examples and potential solutions.

These issues arise because real world data is messy. Many conditions are poorly defined or have conflicting descriptions that are used by different providers, and change over time. For example, there are multiple cancer staging classification systems (eg, American Joint Committee on Cancer, International Federation of Gynecology and Obstetrics, D'Amico risk classification) that have each evolved over time. Even for a seemingly simple lab measurement within one hospital system, there are multiple related data points (eg, orders, billing codes, results) and these are updated at different times in different forms (eg, billing codes, discrete numerical values, PDF, or scanned document).

Of course, these data points, formats, and values change over time and the EHR or billing codes are updated. Furthermore, labs have different data than medications, procedures, etc. There are many copies of medical information stored in multiple places being accessed and modified by multiple people simultaneously. Which version of the record reflects the "truth," and how are conflicts resolved? In recognition of the fundamental problem of provenance (who created the data, who recorded the data, who interpreted the data, etc.) the Office of the National Coordinator of Health Information Technology has introduced several "challenges" to improve upon the *status quo*. ²⁷

There is a standard format called Digital Imaging and Communications in Medicine for storing and transmitting medical imaging data. But there is not yet a widely agreed upon standard format for other EHR data, or even for oncology care. It is notable that the American Society of Clinical Oncology is undertaking an effort to define the Minimal Common Oncology Data Elements to facilitate interoperability and improve data quality for cancer patient care and research.²⁸ Nevertheless, records are often shared by sending consultation and follow-up notes by mail, fax, or electronic mail. In oncology care, there are times when multiple specialists (radiation oncology, medical oncology, surgical specialties, pathology, radiology) are concurrently caring for the patient from different clinics/hospitals. Even when specialists work within the same health system, radiation oncologists, radiologists, and pathologists often use additional EHR systems that are not seamlessly integrated with the health system's primary EHR, creating additional boundaries to communication.

Overview of EHR Applications for Research

The use of EHR for research has rapidly become ubiquitous. One obvious and unique benefit is the ability to study real-world real-patient outcomes in near-real-time. Analyzing existing data also tends to be less expensive and more convenient than creating a human-curated dataset, whether prospectively or retrospectively. There seems to be a near infinite potential for future applications of EHR in research, especially as the types of collected data and ability to extract information from records continue to improve. For example, in the growing field of radiomics, technology improvements are creating both new MRI sequences and calculating new quantitative features from scans. Already, current applications include observational studies (drug utilization, natural history, risk factors), safety surveillance (post-marketing safety surveillance), regulatory work (safety surveillance, pharmacovigilance)¹; and clinical research (hypothesis generation, performance improvement, comparative effectiveness²⁹).

As the use of EHR for research becomes more popular, EHR are gradually being adapted and redesigned to facilitate research more easily. Attempts are in progress to establish standards in data structure and display; common data elements, such as the North American Association of Central Cancer Registries and STandards for Oncology Registry

Table 2 Approximation of the Current State for the Average Electronic Health Record (EHR)

| | Likely to be Found in Structured Data | Likely to be Found in Narrative (+/– Machine Readable) | Unlikely to be Foun |
|---------------------------------------|--|--|---|
| Facts | Name | If female: | Place of birth |
| | Identification numbers | - Number of pregnancies | Sexual history |
| | Date of birth | - Age of menarche | Dietary patterns |
| | Sex | - Age of menopause | Living conditions |
| | Medications as prescribed | Occupation | Other social/ behaviou ral determinants of health |
| | | Substance use | Germline DNA or othe biologic data Photographs |
| Observations (provided | Race/Ethnicity | Symptoms of illness | Gender identity |
| or elicited from | ridee/ Edifficity | Family history of illness and/ or longevity | Medications as taken |
| Observations (obtained | Vital signs | Signs (other than vital signs) | Radiology images |
| by the healthcare practitioner and/or | Most laboratory test results (values) | Physical exam findings | Pathology slide images |
| system) | Walues | Some laboratory test results (esoteric labs, somatic DNA | |
| | | testing) | |
| "Low-level" | Some meaning of laboratory | Meaning of signs | |
| Interpretations | test results (normal/ abnor- | Meaning of signs Meaning of physical exam | |
| P | mal, high/ low etc.) | findings | |
| | a., mgm, 1014 ctc./ | Extended meaning of | |
| | | laboratory test results | |
| | | (incl. genomics reports) | |
| | | Radiology reports | |
| | | Pathology reports | |
| "High-level" | Diagnosis (coded) | Diagnosis (descriptive) | |
| Interpretations | • | Risk for developing a | |
| • | | diagnosis in the future | |
| | | Prediction (ie, will a future action work or not) | |
| | | Prognosis (ie, what is the | |
| | | expected outcome) | |
| Plans | | Diagnostic workup | Survivorship care |
| | | Lifestyle modifications plan | planning |
| | | Treatment plan | |
| | | Risk modification plan | |
| A | 0 1 | (eg, preventative measures) | 1.00 . 1 . 1.00 |
| Actions | Starting and modifying medications | Observation and monitoring | Lifestyle modifications |
| | Removal procedures (eg, excising, eliminating) | Discontinuing a medication | |
| | Additive procedures (eg, device placement) | | |
| | Substitutive procedures | | |
| | (eg, knee replacement) | | |
| | Damaging procedures | | |
| | (eg, radiotherapy) | | |
| Outcomes | Mortality | Success or failure of a plan | Financial toxicity |
| | · | Treatment summarization | Patient reported outcomes (PRO) |
| | | Treatment-related adverse events | Psychological distress |
| | | Time to event (eg, PFS) | Treatment burden (eg, time spent in appointments) |
| | | Complications of natural | |
| | | processes | |

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Entry standards used by national cancer registries; and terminologies for semantic tagging and annotation, such as Systematized Nomenclature of Medicine — Clinical Terms, Logical Observation Identifiers Names and Codes, and RxNorm. ²⁹ But the lack of financial incentives for EHR vendors and users has not facilitated the widespread adoption of standards ²⁹ beyond ICD-9/10 and CPT codes. ³⁰

Example: Natural Language Processing of EHR Data

The vast majority of EHRs were not designed with the intention of research, so it can be both time-consuming and challenging to convert existing EHR data into a format that can be analyzed. For example, the Surveillance, Epidemiology, and End Results³¹ and National Cancer Data Base³² are national databases built by trained and certified cancer registrars who manually enter relevant clinical information extracted from patients' medical charts. Extracting data in a format that can be analyzed can be time- and cost-intensive.

NLP can convert clinical documents into analyzable data elements, turning big data into smart data. NLP techniques are an application from computer science and computational linguistics, and include processing tasks such as tokenization, named entity recognition, and character gazetteer. For interested readers, there are many tutorials online introducing these concepts, including Coursera, Codecademy, and Universities. Most modern NLP approaches use a combination of rule-based and supervised ML approaches.

NLP has been used to extract cancer stage information³⁴; to create oncology treatment summaries³⁵; to automate determination of prostate cancer risk group³⁶; to categorize oncologic responses in radiology reports³⁷; and to mine the EHR for patient-centered outcomes following prostate cancer treatment.³⁰ While these are specific examples that demonstrate a proof of concept, these NLP techniques have the tremendous benefit of being scalable, adapted, and applied to other similar data. In the future, the role of cancer registrars could be quality control and database maintenance, instead of manual data entry.

Within the past few years, unsupervised methods of NLP, especially word2vec and CUI2vec-based approaches, have become increasingly popular. Essentially, these approaches use context within sentences and other clues in the language itself to make determinations, as opposed to classifying to predetermined labels applied by content experts. In the cancer domain, these techniques have primarily been used in the parsing of pathology reports. ^{38,39}

Example: Artificial Intelligence and Machine Learning (ML) Using EHR Data

The use of ML algorithms to generate predictive models in medicine has been increasing nearly exponentially. Jiang et al. queried the PubMed database for deep learning in healthcare; the number of articles increased from about 30 in 2013 to nearly 250 in 2016. ⁴⁰ Supervised, unsupervised, and semisupervised learning algorithms have been used, most commonly using neural networks or support vector machines, and mostly using data from diagnostic imaging. ⁴⁰

ML models have already been used in radiation oncology to predict recurrence patterns after intensity modulated radiotherapy (IMRT) for nasopharyngeal cancer ⁴¹; prognosis with glioblastoma ⁴²; prostate cancer response to IMRT ^{43,44}; and skin dose in low-kV intraoperative radiotherapy. ⁴⁵ In medical oncology, ML has been used to predict which patients will acquire a resistance to EGFR inhibitors ⁴⁶; to predict breast cancer bone metastasis ⁴⁷; and to predict chemotherapy-induced peripheral neuropathy. ⁴⁸ In radiomics, ML has tried to improve pulmonary nodule screening for lung cancer, ⁴⁹ to predict mutation status of renal cell carcinoma, ⁵⁰ and to detect prostate cancer in prostatectomy specimens. ⁵¹ In pathology, ML methods have attempted to identify papillary thyroid carcinoma based on the appearance of fine needle aspiration cytology. ⁵²

Artificial intelligence can be described broadly as the application of computers to mimic the thought processes of human brains; it includes both NLP, ML, and other methods. Artificial intelligence methods can be employed sequentially, using NLP to mine unstructured texts, and ML for creating predictions using the structured output. This sequential strategy has been used to automate extraction of detailed prostate cancer data from clinical notes⁵³; to identify local recurrences of breast cancer⁵⁴; and to extract clinical information from discharge summaries.⁵⁵

Regulatory

The US Food and Drug Administration (FDA) has recently expressed increasing interest in using real-world data, primarily derived from EHRs, to inform regulatory decisions, including postmarketing surveillance and decisions about label expansions. In late 2018, FDA commissioner Gottlieb announced the formation of a new office at the FDA, the Framework for the Real-World Evidence Program, dedicated to utilizing this data.⁵⁶ The FDA and the pharmaceutical industry have a particular interest in the concept of "synthetic controls" - populations derived from heavily annotated EHR data that can serve as a proxy to a prospective control arm in a randomized controlled trial. Taking this idea further, some have advocated for synthetic controls which are amalgamated across EHRs — the ultimate "average patient." These approaches have generated healthy skepticism but are very likely to inform at least a portion of future regulatory decisions.

Education

As the use of EHR data for research continues to increase in both frequency and complexity, the clinical, informatic, and educator communities will need to collaboratively establish and implement core informatics competencies in healthcare.

A newly offered subspecialty medical certification in Clinical Informatics (https://www.theabpm.org)⁵⁷ and a proliferation of high-quality, accessible online education programs, such as the AMIA 10×10 initiative (https://www.amia.org/educa tion), represent important training programs in informatics. However, these resources may not reach the average healthcare provider. To combat this, the NIH Big Data to Knowledge (BD2K) invested \$200 million to address data science challenges, including educational program development, and HITComp (http://hitcomp.org) has formed as an international effort to standardize health information technology competencies. Ultimately though, the fundamentals of EHR-based research remain unfamiliar to the majority of healthcare professionals, leaving them unprepared to assess the validity and applicability of EHR research to real-world healthcare questions. Multidisciplinary efforts to establish and implement standardized informatics education at all levels of the healthcare training pipeline, such as through the AMIA Informatics Educators Forum (https://www.amia.org/ief2019), are critical to closing the informatics knowledge gap.

In the future, a "Complete EHR"?

In order to anticipate what the future may bring to the EHR, it is useful to look to the past. In 1995, the Computer-based Patient Record Institute Work Group defined the EHR as "a virtual compilation of nonredundant health data about a person across a lifetime, including facts, observations, interpretations, plans, actions, and outcomes." While this definition is laudable and provides one with a useful framework, no current EHR technology meets this extremely broad definition, nor is this a goal of current EHR technology. In fact, EHR data is highly redundant and nonstandardized. ⁵⁸ Some facts, observations, and actions are captured in a structured fashion, whereas the majority of interpretations, plans, and outcomes are represented in an unstructured fashion, when they are present at all in the EHR.

Another useful framework is the concept of the Designated Record Set. 59 Per the American Health Information Management Association, "there is no one-size-fits-all definition for the designated record set. The healthcare organization must explicitly define both in a multidisciplinary team approach. Medical staff, for example, should provide guidance to ensure that patient care needs will be met for immediate, long-term, and research uses." Unfortunately, even if it were well defined, the Designated Record Set is more geared to describing the current, as opposed to the possible. Many clinical data elements, some of which may be critically important to the diagnosis, treatment, and continuing care of patients, are simply not present in the EHR with any predictability. Much of what has been coined the "health tapestry" by Weber et al. is completely absent or sparsely recorded within clinical notes. 60

Using the Computer-based Patient Record Institute model, we have approximated the current state (as of 2019) for the average "comprehensive" EHR, ⁷ in **Table 1**. For each of the categories of facts, observations, interpretations, plans,

actions, and outcomes, we have determined: (1) which data elements are likely to be found in structured data; (2) which are likely to be found in free text (some of which will be amenable to extraction through NLP); and (3) which are unlikely to be found in the EHR. As we anticipate which data streams EHR will begin to capture in the future, it is possible that we may also witness revolutions in clinical documentation, such as the introduction of the SOAP note in the 1960s⁶¹ and our recent proposal to "wikify" the medical record.⁶²

Conclusions

In recent decades, EHRs have been adopted in nearly every healthcare system in the US. Due to the financial incentives, EHR have been traditionally designed to aid administration (registration, scheduling, billing) and basic clinical care, rather than research. With the inflation-adjusted decline in research funding, rise in costs of traditional gold-standard prospective randomized clinical trials, and rise in computer science applications for healthcare data, it seems obvious to try to harness this growing amount of data to improve clinical outcomes and efficiency. EHR are gradually starting to be re-designed to be more useful for research purposes, and we are finding new techniques for overcoming limitations. The use of EHR for research is an exciting field with dynamic advances to constantly evolving data and has amazing potential.

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