Chapter 7
Using EHR to Conduct Outcome and Health Services Research

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Take Home Messages

- Electronic Health Records have become an essential tool in clinical research, both as a supplement to existing methods, but also in the growing domains of outcomes research and analytics.
- While EHR data is extensive and analytics are powerful, it is essential to fully understand the biases and limitations introduced when used in health services research.

7.1 Introduction

Data from electronic health records (EHR) can be a powerful tool for research. However, researchers must be aware of the fallibility of data collected for clinical purposes and of biases inherent to using EHR data to conduct sound health outcomes and health services research. Innovative methods are currently being developed to improve the quality of data and thus our ability to draw conclusions from studies that use EHR data.

The United States devotes a large share of the Gross Domestic Product (17.6% in 2009) to health care [1]. With such a huge financial and social investment in healthcare, important questions are fundamental to evaluating this investment:

- How do we know what treatment works and for which patients?
- How much should health care cost? When is too much to pay? In what type of care should we invest more or less resources?
- How does the health system work and how could it function better?

Health services research is a field of research that lives at the intersection of health care policy, management, and clinical care delivery and seeks to answer
these questions. Fundamentally, health services research places the health system under the microscope as the organism of study.

To begin to address these questions, researchers need large volumes of data across multiple patients, across different types of health delivery structures, and across time. The simultaneous growth of this field of research in the past 15 years has coincided with the development of the electronic health record and the increasing number of providers who make use of them in their workspace [2]. The EHR provides large quantities of raw data to fuel this research, both at the granular level of the patient and provider and at the aggregated level of the hospital, state, or nation.

Conducting research with EHR data has many challenges. EHR data are riddled with biases, collected for purposes other than research, inputted by a variety of users for the same patient, and difficult to integrate across health systems [See previous chapter “Confounding by Indication”]. This chapter will focus on the attempts to capitalize on the promise of the EHR for health services research with careful consideration of the challenges researchers must address to derive meaningful and valid conclusions.

### 7.2 The Rise of EHRs in Health Services Research

#### 7.2.1 The EHR in Outcomes and Observational Studies

Observational studies, either retrospective or prospective, attempt to draw inferences about the effects of different exposures. Within health services research, these exposures include both different types of clinical exposures (e.g., does hormone replacement therapy help or hurt patients?) and health care delivery exposures (e.g., does admission to a large hospital for cardiac revascularization improve survival from myocardial infarction over admission to a small hospital). The availability of the extensive health data in electronic health records has fueled this type of research, as data extraction and transcription from paper records has ceased to be a barrier to research. These studies capitalize on the demographic and clinical elements that are routinely recorded as part of an encounter with the health system (e.g., age, sex, race, procedures performed, length of stay, critical care resources used).

We have highlighted a number of examples of this type of research below. Each one is an example of research that has made use of electronic health data, either at the national or hospital level, to draw inferences about health care delivery and care.

Does health care delivery vary? The researchers who compile and examine the Dartmouth Atlas have demonstrated substantial geographic variation in care. In their original article in *Science*, Wennberg and Gittlesohn noted wide variations in the use of health services in Vermont [3]. These authors employed data derived from the use of different types of medical services—home health services, inpatient discharges, etc.—to draw these inferences. Subsequent investigations into national variation in care have been able to capitalize on the availability of such data electronically [4].
Do hospitals with more experience in a particular area perform better? Birkmeyer and colleagues studied the intersection of hospital volume and surgical outcomes with absolute differences in adjusted mortality rates between low volume hospitals and high volume hospitals ranging from 12% for pancreatic resection to 0.2% for carotid endarterectomy [5]. Kahn et al. also used data available in over 20,000 patients to demonstrate that mortality associated with mechanical ventilation was 37% lower in high volume hospitals compared with low volume hospitals [6]. Both of these research groups made use of large volumes of clinical and claims data—Medicare claims data in the case of Birkmeyer and colleagues and the APACHE database from Cerner for Kahn et al.—to ask important questions about where patients should seek different types of care.

How can we identify harm to patients despite usual care? Herzig and colleagues made use of the granular EHR at a single institution and found that the widely-prescribed medications that suppress acid production were associated with an increased risk of pneumonia [7]. Other authors have similarly looked at the EHR found that these types of medications are often continued on discharge from the hospital [8, 9].

To facilitate appropriate modeling and identification of confounders in observational studies, researchers have had to devise methods to extract markers of diagnoses, severity of illness, and patient comorbidities using only the electronic fingerprint. Post et al. [10] developed an algorithm to search for patients who had diuretic-refractory hypertension by querying for patients who had a diagnosis of hypertension despite 6 months treatment with a diuretic. Previously validated methods for reliably measuring the severity of a patient’s illness, such as APACHE or SAPS scores [11, 12], have data elements that are not easily extracted in the absence of manual inputting of data. To meet these challenges, researchers such as Escobar and Elixhauser have proposed alternative, electronically derived methods for both severity of illness measures [13, 14] and identification of comorbidities [14]. Escobar’s work, with a severity of illness measure with an area under the curve of 0.88, makes use of highly granular electronic data including laboratory values; Elixhauser’s comorbidity measure is publically available through the Agency for Healthcare Research and Quality and solely requires billing data [15].

Finally, researchers must develop and employ appropriate mathematical models that can accommodate the short-comings of electronic health data or else they risk drawing inaccurate conclusions. Examples of such modeling techniques are extensive have included propensity scores, causal methods such as marginal structural models and inverse probability weights, and designs from other fields such as instrumental variable analysis [16–19]. The details of these methods are discussed elsewhere in this text.

### 7.2.2 The EHR as Tool to Facilitate Patient Enrollment in Prospective Trials

Despite the power of the EHR to conduct health services and outcomes research retrospectively, the gold standard in research remains prospective and randomized trials. The EHR has functioned as a valuable tool to screen patients at a large scale...
for eligibility. In this instance, research staff uses the data available through the electronic record as a high-volume screening technique to target recruitment efforts to the most appropriate patients. Clinical trials that develop electronic strategies for patient identification and recruitment are at an even greater advantage, although such robust methods have been described as sensitive but not specific, and frequently require coupling screening efforts with manual review of individual records [20]. Embi et al. [21] have proposed using the EHR to simultaneously generate Clinical Trial Alerts, particularly in commercial EHRs such as Epic to leverage the EHR in a point of care strategy. This strategy could expedite enrollment although it must be weighed against the risk of losing patient confidentiality, an ongoing tension between patient care and clinical trial enrollment [22].

7.2.3 The EHR as Tool to Study and Improve Patient Outcomes

Quality can also be tracked and reported through EHRs, either for internal quality improvement or for national benchmarking; the Veterans’ Affairs’ (VA) healthcare system highlights this. Byrne et al. [23] reported that in the 1990s, the VA spent more money on information technology infrastructure and achieved higher rates of adoption compared to the private sector. Their home-grown EHR, which is called VistA, provided a way to track preventative care processes such as cancer and diabetes screening through electronic pop up messages. Between 2004 and 2007, they found that the VA system achieved better glucose and lipid control for diabetics compared to a Medicare HMO benchmark [23]. While much capital investment was needed during the initial implementation of VistA, it is estimated that adopting this infrastructure saved the VA system $3.09 billion in the long term. It also continues to be a source of quality improvement as quality metrics evolve over time [23].

7.3 How to Avoid Common Pitfalls When Using EHR to Do Health Services Research

We would propose the following hypothetical research study as a case study to highlight common challenges to conducting health services research with electronic health data:

Proposed research study: Antipsychotic medications (e.g. haloperidol) are prescribed frequently in the intensive care unit to treat patients with active delirium. However, these medications have been associated with their own potential risk of harm [24] that is separate from the overall risk of harm from delirium. The researchers are interested in whether treatment with antipsychotics increases the risk of in-hospital death and increases the cost of care and use of resources in the hospital.
7.3.1 **Step 1: Recognize the Fallibility of the EHR**

The EHR is rarely complete or correct. Hogan et al. [25] tried to estimate how complete and accurate data are in studies that are conducted on an EHR, finding significant variability in both. Completeness ranged from 31 to 100 % and correctness ranged from 67 to 100 % [25]. Table 7.1 highlights examples of different diagnoses and possible sources of data, which may or may not be present for all patients.

Proposed research study: The researchers will need to extract which patients were exposed to antipsychotics and which were not. However, there is unlikely to be one single place where this information is stored. Should they use pharmacy dispensing data? Nursing administration data? Should they look at which patients were charged for the medications? What if they need these data from multiple hospitals with different electronic health records?

Additionally, even with a robust data extraction strategy, the fidelity of different types of data is variable [26–33]. For example, many EHR systems have the option of entering free text for a medical condition, which may be spelled wrong or be worded unconventionally. As another example, the relative reimbursement of a particular billing code may influence the incidence of that code in the electronic health record so billing may not reflect the true incidence and prevalence of the disease [34, 35].

7.3.2 **Step 2: Understand Confounding, Bias, and Missing Data When Using the EHR for Research**

We would highlight the following methodological issues inherent in conducting research with electronic health records: selection bias, confounding, and missing data. These are explored in greater depth in other chapters of this text.

| Disease state                  | Data source       | Example                                                                 |
|-------------------------------|------------------|-------------------------------------------------------------------------|
| Ischemic heart disease        | Billing data     | ICD-9 code 410 [48]                                                     |
|                               | Laboratory data  | Positive troponin during admission                                       |
|                               | Physician        | In the discharge summary: “the patient was noted to have ST elevations on ECG and was taken to the cath lab”… |
|                               | documentation    |                                                                           |
| Acute lung injury             | Billing data     | ICD-9 code 518.5 and 518.82 with the procedural codes 96.70, 96.71 and 96.72 for mechanical ventilation [49] |
|                               | Radiology data   | “Bilateral” and “infiltrates” on chest x-ray reads [50]                  |
|                               | Laboratory data  | PaO2/FiO2 < 300 mmHg                                                      |
Selection bias, or the failure of the population of study to represent the generalizable population, can occur if all the patients, including controls, are already seeking medical care within an EHR-based system. For example, in EHR-based studies comparing medical versus surgical approaches to the same condition may not be comparing equivalent patients in each group; patients seeking a surgical correction may fundamentally differ from those seeking a more conservative approach. Hripcsak et al. [36] used a large clinical data set from a tertiary center in 2007 to compare mortality from pneumonia to a hand-collected data set that had been published previously; the different search criteria altered the patient population and the subsequent risk of death. While it is not eliminated entirely, selection bias is reduced when prospective randomization takes place [37].

Confounding bias represents the failure to appropriately account for an additional variable that influences both the dependent and independent variable. In research with electronic health records, confounding represents a particular challenge, as identification of all possible confounding variables is nearly impossible.

Proposed research study: The researchers in this study are interested in the patient-level outcomes of what happens to those patients exposed to antipsychotics during their stay. But patients who are actively delirious while in the ICU are likely to be sicker than those who are not actively delirious and sicker patients require more hospital resources. As a result, antipsychotics will appear to be associated with a higher risk of in-hospital mortality and use of hospital resources not due to the independent effect of the drug but rather as a result of confounding by indication.

Missing data or unevenly sampled data collected as part of the EHR creates its own complex set of challenges for health services research. For example, restricting the analysis to patients with only a complete set of data may yield very different (and poorly generalizable) inferences. The multidimensionality of this problem often goes unexamined and underestimated. Nearly all conventional analytic software presumes completeness of the matrix of data, leading many researchers to fail to fully address these issues. For example, data can be misaligned due to lack of sampling, missing data, or simple misalignment. In other words, the data could not be measured during a period of time for an intentional reason (e.g., a patient was extubated and therefore no values for mechanical ventilation were documented) and should not be imputed or the data was measured but was unintentionally not recorded and therefore can be imputed. Rusanov et al. studied 10,000 outpatients at a tertiary center who underwent general anesthesia for elective procedures. Patients with a higher risk of adverse outcome going into surgery had more data points including laboratory values, medication orders and possibly admission orders compared to less sick patients [38], making the missing data for less sick patients intentional. Methods for handling missing data have included omitting cases are not complete, pairwise deletion, mean substitution, regression substitution, or using modeling techniques for maximum likelihood and multiple imputation [39].
7.4 Future Directions for the EHR and Health Services Research

7.4.1 Ensuring Adequate Patient Privacy Protection

It is controversial whether using EHR for research goes against our national privacy standard. In large cohorts, many patients may be present with the same health information, thereby rendering the data sufficiently deidentified. Further, Ingelfinger et al. acknowledge that countries with healthcare registries such as Scandinavia have a distinct research advantage [40]. However, health information is a protected class of information under the Health Insurance Portability and Accountability Act, so there is significant awareness among U.S. healthcare professionals and researchers about its proper storage and dissemination. Some argue that patients should be consented (versus just notified) that their information could be used for research purposes in the future. Ingelfinger et al. [40] recommends IRB approval of registries and a rigorous deidentification process.

Public perception on the secondary use of EHR may not be as prohibitive as policymakers may have believed. In a survey of 3300 people, they were more willing to have their information used for research by university hospitals, compared to public health departments or for quality improvement purposes [41]. They were much less willing to contribute to marketing efforts or have the information used by pharmaceutical companies [41].

With the growing amount of information being entered into EHRs across the country, the American Medical Informatics Association convened a panel to make recommendations for how best to use EHR securely for purposes other than direct patient care. In 2006, the panel called for a national standard to deal with the issue of privacy. They described complex situations where there were security breaches due to problems with deidentification or data was being sold by physicians for profit [42]. While the panel demanded that the national framework be transparent, comprehensive and publicly accepted, they did not propose a particular standard at that time [42]. Other groups such as the Patient-Centered Outcomes Research Institute have since addressed the same conflict in a national forum in 2012. Similarly, while visions were discussed, no explicit recommendation was set forth [PCORI]. Controversy continues in this area.

7.5 Multidimensional Collaborations

Going forward, the true power of integrated data can only be harnessed by forming more collaborations, both within institutions and between them. Research on a national scale in the U.S. has been shown to be feasible. The FDA implemented a pilot program in 2009 called the Mini-Sentinel program. It brought together 31 academic and private organizations to monitor for safety events related to
medications and devices currently on the market [43]. Admittedly, merging databases may require significant financial resources, especially if the datasets need to be coded and/or validated, but researchers like Bradley et al. [44] believe this is a cost-effective use of grant money because of the vast potential to make advances in the way we deliver care. Fundamental to the feasibility of multidimensional collaborations is the ability to ensure accuracy of large-scale data and integrate it across multiple health record technologies and platforms. Efforts to ensure data quality and accessibility must be promoted alongside patient privacy.

7.6 Conclusion

Researchers continue to ask fundamental questions of our health system, making use of the deluge of data generated by EHRs. Unfortunately, that deluge is messy and problematic. As the field of health services research with EHRs continues to evolve, we must hold researchers to rigorous standards [45] and encourage more investment in research-friendly clinical databases as well as cross-institutional collaborations. Only then will the discoveries in health outcomes and health services research be one click away [46, 47]. It is time for healthcare to reap the same reward from a rich data source that is already in existence.

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