Diagnosing Diagnosis Errors: Lessons from a Multi-institutional Collaborative Project


Abstract

Background: Diagnosis errors are frequent and important, but represent an underemphasized and understudied area of patient safety. Diagnosis errors are challenging to detect and dissect. It is often difficult to agree whether an error has occurred, and even harder to determine with certainty its causes and consequence. The authors applied four safety paradigms: (1) diagnosis as part of a system, (2) less reliance on human memory, (3) need to open “breathing space” to reflect and discuss, (4) multidisciplinary perspectives and collaboration. Methods: The authors reviewed literature on diagnosis errors and developed a taxonomy delineating stages in the diagnostic process: (1) access and presentation, (2) history taking/collection, (3) the physical exam, (4) testing, (5) assessment, (6) referral, and (7) followup. The taxonomy identifies where in the diagnostic process the failures occur. The authors used this approach to analyze diagnosis errors collected over a 3-year period of weekly case conferences and by a survey of physicians. Results: The authors summarize challenges encountered from their review of diagnosis error cases, presenting lessons learned using four prototypical cases. A recurring issue is the sorting-out of relationships among errors in the diagnostic process, delay and misdiagnosis, and adverse patient outcomes. To help understand these relationships, the authors present a model that identifies four key challenges in assessing potential diagnosis error cases: (1) uncertainties about diagnosis and findings, (2) the relationship between diagnosis failure and adverse outcomes, (3) challenges in reconstructing clinician assessment of the patient and clinician actions, and (4) global assessment of improvement opportunities. Conclusions and recommendations: Finally the authors catalogue a series of ideas for change. These include: reengineering followup of abnormal test results; standardizing protocols for reading x-rays/lab tests, particularly in training programs and after hours; identifying “red flag” and “don’t miss” diagnoses and situations and use of manual and automated check-lists; engaging patients on multiple levels to become “coproducers” of safer medical diagnosis practices; and weaving “safety nets” to mitigate harm from uncertainties and errors in diagnosis. These change ideas need to be tested and implemented for more timely and error-free diagnoses.
**Diagnosing Diagnosis Errors: Lessons From A Multi-Institutional Collaborative Project**

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Introduction

Diagnosis errors are frequent and important, but represent an underemphasized and understudied area of patient-safety.\(^1-8\) This belief led us to embark on a 3-year project, funded by the Agency for Healthcare Research and Quality (AHRQ), to better understand where and how diagnosis fails and explore ways to target interventions that might prevent such failures. It is known that diagnosis errors are common and underemphasized, but they are also challenging to detect and dissect. It is often difficult even to agree whether or not a diagnosis error has occurred.

In this article we describe how we have applied patient safety paradigms (blame-free reporting/reviewing/learning, attention to process and systems, an emphasis on communication and information technology) to better understand diagnosis error.\(^2, 7-9\)

We review evidence about the types and importance of diagnosis errors and summarize challenges we have encountered in our review of more than 300 cases of diagnosis error. In the second half of the article, we present lessons learned through analysis of four prototypical cases. We conclude with suggested “change ideas”—interventions for improvement, testing, and future research.

Although much of the patient safety spotlight has focused on medication errors, two recent studies of malpractice claims revealed that diagnosis errors far outnumber medication errors as a cause of claims lodged (26 percent versus 12 percent in one study;\(^10\) 32 percent versus 8 percent in another study\(^11\)). A Harris poll commissioned by the National Patient Safety Foundation found that one in six people had personally experienced a medical error related to misdiagnosis.\(^12\)

Most medical error studies find that 10–30 percent (range = 0.6–56.8 percent) of errors are errors in diagnosis (Table 1).\(^1-3, 5, 11, 13-21\) A recent review of 53 autopsy studies found an average rate of 23.5 percent major missed diagnoses (range = 4.1–49.8 percent). Selected disease-specific studies (Table 2),\(^6, 22-32\) also show that substantial percentages of patients (range = 2.1 – 61 percent) experienced missed or delayed diagnoses. Thus, while these studies view the problem from varying vantage points using heterogeneous methodologies (some nonsystematic and lacking in standardized definitions), what emerges is compelling evidence for the frequency and impact of diagnosis error and delay.

Of the 93 safety projects funded by AHRQ, only 1 is focused on diagnosis error, and none of the 20 evidence-based AHRQ Patient Safety Indicators directly measures failure to diagnose.\(^33\) Nonetheless, for each of AHRQ’s 26 “sentinel complications” (e.g., decubitus ulcer, iatrogenic pneumothorax, postoperative sepsis, accidental puncture/laceration), timely diagnosis can be decisive in determining whether patients experience major adverse outcomes. Hence, while diagnosis error remains more in the shadows than in the spotlight of patient safety, this aspect of clinical medicine is clearly vulnerable to well-documented failures and warrants an examination through the lens of modern patient safety and quality improvement principles.
<table>
<thead>
<tr>
<th>Author, Year</th>
<th>Context/Design</th>
<th>Diagnosis errors/Total errors (%)</th>
<th>Comments/Types of Diagnostic Errors</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flannery FT (1991)</td>
<td>Physicians and surgeons update, St. Paul, MN. Malpractice claims data reviewed</td>
<td>1126/7233 (27.4%) failure to diagnose</td>
<td>Top 5 diagnoses (cancer, circulatory/thrombosis, fracture/dislocation, lack of attendance, infection)</td>
</tr>
<tr>
<td>Leape LL (1991)</td>
<td>AE in hospitalized patients</td>
<td>168/1276 (13.8%)</td>
<td>Failure to use indicated tests, act on test, appropriate test, delay in diagnosis, practicing outside area of expertise</td>
</tr>
<tr>
<td>Bogner M (1994)</td>
<td>Medical practice study, 1984, AE in NY hospitals (Leape)</td>
<td>11731/68645 (17.1%)</td>
<td>17.1% preventable errors due to diagnostic errors. 71% of them were negligent.</td>
</tr>
<tr>
<td>Bhasale AL (1998)</td>
<td>Austrian GP, Diagnostic incidents</td>
<td>275/805 (34.2%)</td>
<td>Calculated rates (N = 275): missed diagnosis (40 per 100), delayed (34), misdiagnosed (23), and diagnostic procedural (14)</td>
</tr>
<tr>
<td>Wilson RM (1999)</td>
<td>The Quality in Australian Health Care Study</td>
<td>267/470 (56.8%)</td>
<td>1922/2351 AEs (81.8%) associated w/ human errors, 470 (20%) delays in diagnosis or treatment, 460 (19.6%) treatment, and 252 (10.7%) investigation. 1922 AEs w/ 2940 causes identified. Causes of human errors: 465/2940 (15.8%) failure to synthesis/decide/act on information, 346 (11.8%) failure to request, arrange an investigation.</td>
</tr>
<tr>
<td>Weingart S (2000)</td>
<td>Potential adverse events</td>
<td>29/110 (26.4%) process of care problems associated w/ diagnosis</td>
<td>18/110 (16.4%) inadequate evaluation, 7 (6.4%) diagnostic error, 4 (3.6%) delayed consultation.</td>
</tr>
<tr>
<td>Neale G (2001)</td>
<td>AE in England hospitals</td>
<td>5/118 (0.6%)</td>
<td>118/840 (14%) had AE. 57% of all AE, cognitive. 5/118 (0.6%) of admissions were associated with incorrect diagnoses. 2 missed heart failures, 2 incorrect assessment of abdominal pain, and 1 missed fracture.</td>
</tr>
<tr>
<td>Baker GR (2002)</td>
<td>Mailed survey questionnaires regarding patient safety issues, Canadian health care facilities, and colleges and associations</td>
<td>1/25 (4%) of health care errors in health care facilities, 9/21 (39%) in colleges/associations</td>
<td>Human factors, 3 (12%), 9 (39%); competency, 5 (20%), 2 (9%)</td>
</tr>
</tbody>
</table>
Table 1. General medical error studies that reported errors in diagnosis, cont.

<table>
<thead>
<tr>
<th>Author, Year</th>
<th>Context/Design</th>
<th>Diagnosis errors/Total errors (%)</th>
<th>Comments/Types of Diagnostic Errors</th>
</tr>
</thead>
<tbody>
<tr>
<td>JCAHO Sentinel event advisory group (2002)</td>
<td>ED sentinel event</td>
<td>55/23 (42%)</td>
<td>55 delays in treatment, due to misdiagnosis (42%), test results availability (13%), delayed initial assessment (7%). Most frequent missed diagnosis: meningitis 7/23 (30%), cardiac disease, PE, trauma, asthma, neurologic disorder</td>
</tr>
<tr>
<td>Makeham MA (2002)</td>
<td>GP in 6 countries</td>
<td>17/104 (13%) in Austria, 55/236 (19%) in other countries</td>
<td>Process errors 104/134 (78%) in Austria, 235/301 (78%), Of the process errors, investigation errors (lab errors, diagnostic imaging errors, and others) were 13%, 19%</td>
</tr>
<tr>
<td>Medical malpractice lawyers and attorneys online (2002)</td>
<td>Medical malpractice</td>
<td>40%</td>
<td>Failure-to-diagnose, failed to diagnosis in timely fashion</td>
</tr>
<tr>
<td>Chaudhry S (2003)</td>
<td>Error detection by attending physicians, general medicine inpatients</td>
<td>12/63 (19.1%) of all errors, 8/39 (20.5%) of near misses, and 4/24 (16.7%) of AEs were related with diagnostic errors</td>
<td>55/528 (10.4%) patients admitted to the hospitalists had at least 1 error</td>
</tr>
<tr>
<td>Kravitz RL (2003)</td>
<td>Malpractice claims data, 4 specialties</td>
<td>3–8%</td>
<td>1371 claims, Failure to appropriate diagnostic testing/monitoring</td>
</tr>
<tr>
<td>Phillips R (2004)</td>
<td>Malpractice claims in primary care</td>
<td>26126 claims peer reviewed, 5921/26126 (23%) related with negligent</td>
<td>2003/5921 (34%) associated with diagnosis error, 16% failure to monitor case, 4% failure/delay in referral, 2% failure to recognize a complication</td>
</tr>
</tbody>
</table>

AE = adverse events; GP = general practitioners; JCAHO = Joint Commission for Accreditation of Healthcare Organizations; ED = emergency department
Table 2. Illustrative disease-specific studies of diagnosis errors

<table>
<thead>
<tr>
<th>Author</th>
<th>Disease/Design</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Steere AC (1993)</td>
<td>Lyme disease</td>
<td>Overdiagnosis of Lyme disease: patients given the diagnosis but on review did not meet criteria for diagnosis. - 452/788 (57%)</td>
</tr>
<tr>
<td>Cravan ER (1994)</td>
<td>Glaucoma</td>
<td>Diagnosis error in glaucoma claims/lawsuits - 42/194 (21.7%)</td>
</tr>
<tr>
<td>Lederle FA (1994)</td>
<td>Ruptured abdominal aortic aneurysm</td>
<td>Ruptured aneurysm diagnosis initially missed - 14/23 (61%)</td>
</tr>
<tr>
<td>Mayer PL (1996)</td>
<td>Symptomatic cerebral aneurysm</td>
<td>Patients initially misdiagnosed - 54/217 (25%, ranging from 13% to 35% at 4 sites); Of these, 26/54 (48%) deteriorated before definite treatment; erroneous working diagnoses in 54 pts; 8 (15%) viral meningitis, 7 (13%) migraine, 7 (13%) headache of uncertain etiology</td>
</tr>
<tr>
<td>Williams V (1997)</td>
<td>Brain and spinal cord biopsies</td>
<td>Errors based on &quot;second opinion&quot; reviews - 214/500 (43%); 44/214 (20%) w/ serious complications, 96 (45%) w/ substantial, 50 (23%) w/ minor errors</td>
</tr>
<tr>
<td>Clark S (1998)</td>
<td>Pyrogenic spinal infection</td>
<td>Delay in diagnosis - 41/69 (59.4%); more than 1 month of back/neck pain before specialist referral</td>
</tr>
<tr>
<td>Arbiser ZK (2000)</td>
<td>Soft tissue pathology</td>
<td>Second option reviews: minor discrepancy - 20/266 (7.5%); major discrepancy - 65 (25%)</td>
</tr>
<tr>
<td>Pope JH (2000)</td>
<td>Acute cardiac ischemia in ED</td>
<td>Mistakenly discharged from ED; MI - 894/889 (2.1%);</td>
</tr>
<tr>
<td>Edelman D (2002)</td>
<td>Diabetes in an outpatient clinic</td>
<td>Blood glucoxs meeting criteria for DM with no diagnosis of diabetes in records - 258/1426 (18%)</td>
</tr>
<tr>
<td>Goodson WH (2002)</td>
<td>Breast Cancer</td>
<td>Inappropriately reassured to have benign lesions - 21/435 (5%); 14 (3%) misread mammogram, 4 (1%) misread pathologic finding, 5 (1%) missed by poor fine-needle biopsy</td>
</tr>
<tr>
<td>Kawalski RG (2004)</td>
<td>Aneurysmal subarachnoid hemorrhage (SAH)</td>
<td>Patients w/ SAH initially misdiagnosed - 56/482 (12%); migraine/tension headache most common incorrect dx – 20/56 (36%); failure to obtain a CT was most common error - 41/56 (73%)</td>
</tr>
</tbody>
</table>

### Traditional and innovative approaches to learning from diagnosis error

Traditionally, studying missed diagnoses or incorrect diagnoses had a central role in medical education, research, and quality assurance in the form of autopsies.34–36 Other traditional methods of learning about misdiagnosed cases include malpractice litigation, morbidity and mortality (M&M) conferences, unsystematic feedback from patients, other providers, or simply from patients’ illnesses as they evolved over time.3, 14, 15, 32 Beyond the negative aspects of being grounded in patients’ deaths or malpractice accusations, there are other limitations of these historical approaches, including
• Lack of systematic approaches to surveillance, reporting, and learning from errors, with nonrandom sample of cases subjected to such review.\textsuperscript{37, 39}

• Lack of timeliness, with cases often reviewed months or years after the event.\textsuperscript{38}

• Examinations that rarely dig to the root of problems: not focused on the “Five Whys.”\textsuperscript{40}

• Postmortems that seldom go beyond the case-at-hand, with minimal linkages to formal quality improvement activities.\textsuperscript{41}

• Atrophy of the value of even these suboptimal approaches, with autopsy rates in the single digits (in many hospitals, zero), many malpractice experiences sealed by nondisclosure agreements, and shorter hospitalizations limiting opportunities for followup to ultimate diagnosis.\textsuperscript{34, 41, 42}

What is needed to overcome these limitations is not only a more systematic method for examining cases of diagnosis failure, but also a fresh approach. Therefore, our team approached diagnosis error with the following perspectives:

**Diagnosis as part of a system.** Diagnostic accuracy should be viewed as a system property rather than simply what happens between the doctor’s two ears.\textsuperscript{2, 43–45} While cognitive issues figure heavily in the diagnostic process, a quite from Don Berwick summarizes\textsuperscript{46} a much lacking and needed perspective: “Genius diagnosticians make great stories, but they don’t make great health care. The idea is to make accuracy reliable, not heroic.”

**Less reliance on human memory.** Relying on clinicians’ memory—to trigger consideration of a particular diagnosis, recall a disease’s signs/symptoms/pattern from a textbook or experience—or simply to remember to check on a patient’s lab result—is an invitation to variations and failures. This lesson from other error research resonates powerfully with clinicians, who are losing the battle to keep up to date.\textsuperscript{9, 45, 47}

**Need for “space” to allow open reflection and discussion.** Transforming an adversarial atmosphere into one conducive to honest reflection is an essential first step.\textsuperscript{48, 49} However, an equally important and difficult challenge is creating venues that allow clinicians (and patients) to discuss concerns in an efficient and productive manner.\textsuperscript{37} Cases need to be reviewed in sufficient detail to make them “real.” Firsthand clinical information often radically changes our understanding from what the more superficial “first story” suggested. As complex clinical circumstances are better understood, new light is often shed on what at first appeared to be indefensible diagnostic decisions and actions. Unsuspected additional errors also emerge. Equally important is not to get mired in details or making judgments (whether to label a case as a diagnosis error). Instead, it is more valuable to focus on generalizable lessons of how to ensure better treatment of similar future patients.\textsuperscript{16}
Adopting multidisciplinary perspectives and collaboration. A broad range of skills and vantage points are valuable in understanding the complex diagnostic problems that we encountered. We considered input from specialists and primary care physicians to be essential. In addition, specialists in emergency medicine (where many patients first present) offered a vital perspective, both for their diagnostic expertise and their pivotal interface with system constraints (resource limits mean that not every patient with a confusing diagnosis can be hospitalized). Even more valuable has been the role of non-MDs, including nursing quality specialists, information scientists, and social scientists (cognitive psychologist, decision theory specialist) in forging a team to broadly examine diagnosis errors.

Innovative screening approaches. Developing new ways to uncover errors is a priority. We cannot afford to wait for a death, lawsuit, or manual review. Approaches we have been exploring include electronic screening that links pharmacy and lab data (e.g., to screen for abnormal results, such as elevated thyroid stimulating hormone [TSH], unaddressed by thyroxin therapy), trajectory studies (retrospectively probing delays in a series of cases with a particular diagnosis), and screening for discrepancies between admitting and discharge diagnoses. A related approach is to survey specialists (who are poised to see diagnoses missed in referred patients), primary care physicians (about their own missed diagnoses), or patients themselves (who frequently have stories to share about incorrect diagnoses), in addition to various ad hoc queries and self-reports.

Where does the diagnostic process fail?

One of the most powerful heuristics in medication safety has been delineation of the steps in the medication-use process (prescribing, transcribing, dispensing, administering, and monitoring) to help localize where an error has occurred. Diagnosis, while more difficult to neatly classify (because compared to medications, stages are more concurrent, recurrent, and complex), nonetheless can be divided into seven stages: (1) access/presentation, (2) history taking/collection, (3) the physical exam, (4) testing, (5) assessment, (6) referral, and (7) followup. We have found this framework helpful for organizing discussions, aggregating cases, and targeting areas for improvement and research. It identifies what went wrong, and situates where in the diagnostic process the failure occurred (Table 3). We have used it for a preliminary analysis of several hundred diagnosis error cases we collected by surveying physicians.

This taxonomy for categorizing diagnostic “assessment” draws on work of Kassirer and others, highlighting the two key steps of (a) hypothesis generation, and (b) differential diagnosis or hypothesis weighing/prioritization. We add another aspect of diagnostic assessment, one that connects to other medical and iatrogenic error work—the need to recognize the urgency of diagnoses and complications. This addition underscores the fact that failure to make the exact diagnosis is often less important than correctly assessing the urgency of the patient’s illness. We divide the “testing” stage into three components—ordering, performing, and clinician processing (similar but not identical to the laboratory literature classification of the phases of lab testing as preanalytic, analytic, and
Table 3. Taxonomy of where and what errors occurred

<table>
<thead>
<tr>
<th>Where in Diagnostic Process (~Anatomic localization)</th>
<th>What Went Wrong (~Lesion)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Access/presentation</td>
<td>Denied care</td>
</tr>
<tr>
<td></td>
<td>Delayed presentation</td>
</tr>
<tr>
<td>2. History</td>
<td>Failure/delay in eliciting critical piece of history data</td>
</tr>
<tr>
<td></td>
<td>Inaccurate/misinterpretation &quot;</td>
</tr>
<tr>
<td></td>
<td>Suboptimal weighing &quot;</td>
</tr>
<tr>
<td></td>
<td>Failure/delay to followup &quot;</td>
</tr>
<tr>
<td>3. Physical exam</td>
<td>Failure/delay in eliciting critical physical exam finding</td>
</tr>
<tr>
<td></td>
<td>Inaccurate/misinterpreted &quot;</td>
</tr>
<tr>
<td></td>
<td>Suboptimal weighing &quot;</td>
</tr>
<tr>
<td></td>
<td>Failure/delay to followup &quot;</td>
</tr>
<tr>
<td>4. Tests (lab/radiology)</td>
<td>Failure/delay in ordering needed test(s)</td>
</tr>
<tr>
<td></td>
<td>Failure/delay in performing ordered test(s)</td>
</tr>
<tr>
<td></td>
<td>Suboptimal test sequencing</td>
</tr>
<tr>
<td></td>
<td>Ordering of wrong test(s)</td>
</tr>
<tr>
<td></td>
<td><strong>Performance</strong></td>
</tr>
<tr>
<td></td>
<td>Sample mix-up/mislabeled (e.g., wrong patient)</td>
</tr>
<tr>
<td></td>
<td>Technical errors/poor processing of specimen/test</td>
</tr>
<tr>
<td></td>
<td>Erroneous lab/radiol reading of test</td>
</tr>
<tr>
<td></td>
<td>Failed/delayed transmission of result to clinician</td>
</tr>
<tr>
<td></td>
<td><strong>Clinician processing</strong></td>
</tr>
<tr>
<td></td>
<td>Failed/delayed followup action on test result</td>
</tr>
<tr>
<td></td>
<td>Erroneous clinician interpretation of test</td>
</tr>
<tr>
<td>5. Assessment</td>
<td>Failure/delay in considering the correct diagnosis</td>
</tr>
<tr>
<td></td>
<td><strong>Suboptimal weighing/prioritizing</strong></td>
</tr>
<tr>
<td></td>
<td>Too much weight to low(er) probability/priority dx</td>
</tr>
<tr>
<td></td>
<td>Too little consideration of high(er) probability/priority dx</td>
</tr>
<tr>
<td></td>
<td>Too much weight on competing diagnosis</td>
</tr>
<tr>
<td></td>
<td><strong>Recognizing urgency/complications</strong></td>
</tr>
<tr>
<td></td>
<td>Failure to appreciate urgency/acuity of illness</td>
</tr>
<tr>
<td></td>
<td>Failure/delay in recognizing complication(s)</td>
</tr>
<tr>
<td>6. Referral/consultation</td>
<td>Failure/delay in ordering needed referral</td>
</tr>
<tr>
<td></td>
<td>Inappropriate/unneeded referral</td>
</tr>
<tr>
<td></td>
<td>Suboptimal consultation diagnostic performance</td>
</tr>
<tr>
<td></td>
<td>Failed/delayed communication/followup of consultation</td>
</tr>
<tr>
<td>7. Followup</td>
<td>Failure to refer to setting for close monitoring</td>
</tr>
<tr>
<td></td>
<td>Failure/delay in timely followup/rechecking of patient</td>
</tr>
</tbody>
</table>
Diagnosing Diagnosis Errors

For each broad category, we specified the types of problems we observed.

A recurring theme running through our reviews of potential diagnosis error cases pertains to the relationship between errors in the diagnostic process, delay and misdiagnosis, and adverse patient outcomes. Bates has promulgated a useful model for depicting the relationships between medication errors and outcomes. Similarly, we find that most errors in the diagnostic process do not adversely impact patient outcomes. And, many adverse outcomes associated with misdiagnosis or delay do not necessarily result from any error in the diagnostic process—the cancer may simply be undiagnosable at that stage, the illness presentation too atypical, rare, or unlikely even for the best of clinicians to diagnose early. These situations are often referred to as “no fault” or “forgivable” errors—terms best avoided because they imply fault or blame for preventable errors (Figure 1).

While deceptively simple, the model raises a series of extremely challenging questions—questions we found ourselves repeatedly returning to in our weekly discussions. We hope these questions can provide insights into recurring themes and challenges we faced, and perhaps even serve as a checklist for others to structure their own patient care reviews. While humbled by our own inability to provide more conclusive answers to these questions, we believe researchers and practitioners will be forced to grapple with them before we can make significant progress.

Questions for consideration by diagnosis error evaluation and research (DEER) investigators in assessing cases

Uncertainties about diagnosis and findings

1. What is the correct diagnosis? How much certainty do we have, even now, about what the correct diagnosis is?

2. What were the findings at the various points in time when the patient was being seen; how much certainty do we have that a particular finding and diagnosis was actually present at the time(s) we are positing an error?

Relationship between diagnosis failure and adverse outcomes

3. What is the probability that the error resulted in the adverse outcome? How treatable is the condition, and how critical is timely diagnosis and treatment for impacting on the outcome—both in general and in this case?

4. How did the error in the diagnostic process contribute to making the wrong diagnosis and wrong treatment?
Figure 1. Relationships between diagnostic process errors, misdiagnosis, and adverse events

*Delayed, missed, or misdiagnosis

Caption

Group A = Errors in diagnostic process (blood sample switched between two patients, MD doesn’t do a physical exam for patient with abdominal pain)

Group B = Diagnostic process error with resulting misdiagnosis (patient given wrong diagnosis because blood samples switched)

Group C = Adverse outcome resulting from error-related misdiagnosis (Patient is given toxic treatment and has adverse effect as result of switched samples. Fail to diagnose appendicitis because of failure to examine abdomen, and it ruptures and patient dies)

Group D = Harm from error in diagnostic process (colon perforation from colonoscopy done on wrong patient)

Group E = Misdiagnosis, delayed diagnosis or missed diagnosis, but no error in care or harm (incidental prostate cancer found on autopsy)

Group F = Adverse event due to misdiagnosis but no identifiable process error (death from acute MI but no chest pain or other symptoms that were missed)

Group G = Adverse events but not related to misdiagnosis, delay, or error in diagnostic process, e.g., death from correctly diagnosed disease complication, or nonpreventable drug reaction (PCN anaphylaxis in patient never previously exposed)
Clinician assessment and actions

5. What was the physician’s diagnostic assessment? How much consideration was given to the correct diagnosis? (This is usually difficult to reconstruct because differential diagnosis often is not well documented.)

6. How good or bad was the diagnostic assessment based on evidence clinicians had on hand at that time (should have been obvious from available data vs. no way anyone could have suspected)?

7. How erroneous was the diagnostic assessment, based on the difficulty in making the diagnosis at this point? (Was there a difficult “signal-to-noise” situation, a rare low-probability diagnosis, or an atypical presentation?)

8. How justifiable was the failure to obtain additional information (i.e., history, tests) at a particular point in time? How can this be analyzed absolutely, as well as relative to the difficulties and constraints in obtaining this missing data? (Did the patient withhold or refuse to give accurate/additional history; were there backlogs and delays that made it impossible to obtain the desired test?)

9. Was there a problem in diagnostic assessment of the severity of the illness, with resulting failure to observe or follow up the patient more closely? (Again, both absolutely and relative to constraints.)

Global assessment of improvement opportunities

10. To what extent did the clinicians’ actions deviate from the standard-of-care (i.e., was there negligent care with failure to follow accepted diagnostic guidelines and expected practices, or to pursue abnormal finding that should never be ignored)?

11. How preventable was the error? How ameliorable or amenable to change are the factors/problems that contributed to the error? How much would such changes, designed to prevent this error in the future, cost?

12. What should we do better the next time we encounter a similar patient or situation? Is there a general rule, or are there measures that can be implemented to ensure this is reliably done each time?

Diagnosis error case vignettes

Case 1

A 25-year-old woman presents with crampy abdominal pain, vaginal bleeding, and amenorrhea for 6 weeks. Her serum human choriogonadotropin (HCG) level is markedly elevated. A pelvic ultrasound is read by the on-call radiology chief resident and obstetrics (OB) attending physician as showing an empty uterus,
suggesting ectopic pregnancy. The patient is informed of the findings and treated with methotrexate. The following morning the radiology attending reviews the ultrasound and amends the report, officially reading it as “normal intrauterine pregnancy.”

**Case 2**

A 49-year-old, previously healthy man presents to the emergency department (ED) with nonproductive cough, “chest congestion,” and dyspnea lasting 2 weeks; he has a history of smoking. The patient is afebrile, with pulse = 105, respiration rate (RR) = 22, and white blood count (WBC) = 6.4. Chest x-ray shows “marked cardiomegaly, diffuse interstitial and reticulonodular densities with blunting of the right costophrenic angle; impression—congestive heart failure (CHF)/pneumonia. Rule out (R/O) cardiomyopathy, valve disease or pericardial effusion.” The patient is sent home with the diagnosis of pneumonia, with an oral antibiotic.

The patient returns 1 week later with worsening symptoms. He is found to have pulsus paradoxicus, and an emergency echocardiogram shows massive pericardial effusion. Pericardiocentesis obtains 350 cc fluid with cytology positive for adenocarcinoma. Computed tomography of the chest suggests “lymphangitic carcinomatosis.”

**Case 3**

A 50-year-old woman with frequent ED visits for asthma (four visits in the preceding month) presents to the ED with a chief complaint of dyspnea and new back pain. She is treated for asthma exacerbation and discharged with nonsteroidal anti-inflammatory drugs (NSAID) for back pain.

She returns 2 days later with acutely worsening back pain, which started when reaching for something in her cupboard. A chest x-ray shows a “tortuous and slightly ectatic aorta,” and the radiologist’s impression concludes, “If aortic dissection is suspected, further evaluation with chest CT with intravenous (IV) contrast is recommended.” The ED resident proceeds to order a chest CT, which concludes “no evidence of aneurysm or dissection.” The patient is discharged.

She returns to the ED 3 days later, again complaining of worsening asthma and back pain. While waiting to be seen, she collapses in the waiting room and is unable to be resuscitated. Autopsy shows a ruptured aneurysm of the ascending aorta.

**Case 4**

A 50-year-old woman with a past history diabetes and alcohol and IV drug abuse, presents with symptoms of abdominal pain and vomiting and is diagnosed as having “acute chronic pancreatitis.” Her amylase and lipase levels are normal. She is admitted and treated with IV fluids and analgesics. On hospital day 2 she begins having spiking fevers and antibiotics are administered. The next day, blood cultures are growing gram negative organisms.
At this point, the service is clueless about the patient’s correct diagnosis. It only becomes evident the following day when (a) review of laboratory data over the past year shows that patient had four prior blood cultures, each positive with different gram negative organisms; (b) a nurse reports patient was “behaving suspiciously,” rummaging through the supply room where syringes were kept; and (c) a medical student looks up posthospital outpatient records from 4 months earlier and finds several notes stating that “the patient has probable Munchausen syndrome rather than pancreatitis.” Upon discovering these findings, the patient’s IVs are discontinued and sensitive, appropriate followup primary and psychiatric care are arranged.

A postscript to this admission: 3 months later, the patient was again readmitted to the same hospital for “pancreatitis” and an unusual “massive leg abscess.” The physicians caring for her were unaware of her past diagnoses and never suspected or discovered the likely etiology of her abscess (self-induced from unsterile injections).

Lessons and issues raised by the diagnosis error cases

Difficulties in sorting out “don’t miss” diagnoses

Before starting our project, we compiled a list of “don’t miss” diagnoses (available from the authors). These are diagnoses that are considered critical, but often difficult to make—critical because timely diagnosis and treatment can have major impact (for the patient or the public’s health, or both), and difficult because they either are rare or pose diagnostic challenges. Diagnoses such as spinal epidural abscess (where paraplegia can result from delayed diagnosis), or active pulmonary tuberculosis (TB) (where preventing spread of infection is critical) are examples of “don’t miss” diagnoses. While there is a scant evidence base to definitively compile and prioritize such a list, three of our cases—ectopic pregnancy, dissecting aortic aneurysm, and pericardial effusion with tamponade—are diagnoses that would unquestionably be considered as life-threatening diagnoses that ought not be delayed or missed.25

Although numerous issues concerning diagnosis error are raised by these cases, they also illustrate problems relating to uncertainties, lack of gold standards (for both testing and standard of care), and difficulties reaching consensus about best ways to prevent future errors and harmful delays. Below we briefly discuss some of these issues and controversies.

Diagnostic criteria and strategies for diagnosing ectopic pregnancy are controversial, and our patient’s findings were particularly confusing. Even after careful review of all aspects of the case, we were still not certain who was “right”—the physicians who read the initial images and interpreted them as consistent with ectopic pregnancy, or the attending physician rereading the films the next day as normal. The literature is unclear about criteria for establishing this
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In addition, there is a lack of standards in performance and interpretation of ultrasound exams, plus controversies about timing of interventions. Thus this “obvious” error is obviously more complex, highlighting a problem-prone clinical situation.

The patient who was found to have a malignant pericardial effusion illustrates problems in determining the appropriate course of action for patients with unexplained cardiomegaly, which the ED physicians failed to address on his first presentation: Was he hemodynamically stable at this time? If so, did he require an urgent echocardiogram? What criteria should have mandated this be done immediately? How did his cardiomegaly get “lost” as his diagnosis prematurely “closed” on pneumonia when endorsed from the day to the night ED shift? How should one assess the empiric treatment for pneumonia given his abnormal chest x-ray? Was this patient with metastatic lung cancer harmed by the 1 week delay?

The diagnosis of aneurysms (e.g., aortic, intracranial) arises repeatedly in discussions of misdiagnosis. Every physician seems to recall a case of a missed aneurysm with catastrophic outcomes where, in retrospect, warnings may have been overlooked. A Wall Street Journal article recently won a Pulitzer Prize for publicizing such aneurysm cases. Our patient’s back pain was initially dismissed. Because of frequent visits, she had been labeled as a “frequent flyer”—and back pain is an extremely common and nonspecific symptom. A review of literature on the frequency of dissecting aortic aneurysm reveals that it is surprisingly rare, perhaps less than 1 out of 50,000 ED visits for chest pain, and likely an equally rare cause for back pain. She did ultimately undergo a recommended imaging study after a suspicious plain chest x-ray, however it was deemed “negative.”

Thus, in each case, seemingly egregious and unequivocal errors were found to be more complex and uncertain.

**Issues related to limitations of diagnostic testing**

Even during our 3 years of diagnosis case reviews, clinicians have been confronted with rapid changes in diagnostic testing. New imaging modalities, lab tests, and testing recommendations have been introduced, often leaving clinicians confused about which tests to order and how to interpret their confusing and, at times, contradictory (from one radiologist to the next) results.

If diagnosis errors are to be avoided, clinicians must be aware of the limitations of the diagnostic tests they are using. It is well known that a normal mammogram in a woman with a breast lump does not rule out the diagnosis of breast cancer, because the sensitivity of test is only 70 to 85 percent. A recurring theme in our cases is failure to appreciate pitfalls in weighing test results in the context of the patient’s pretest disease probabilities. Local factors, such as the variation in quality of test performance and readings, combined with communication failures between radiology/laboratory and ordering physicians (either no direct communication or interactions where complex interpretations get
reduced to “positive” or “negative,” overlooking subtleties and limitations) provide further sources of error.66

The woman with the suspected ectopic pregnancy, whose emergency ultrasound was initially interpreted as being “positive,” illustrates the pitfalls of taking irreversible therapeutic actions without carefully weighing test reading limitations. Perhaps an impending rupture of an ectopic pregnancy warrants urgent action. However, it is also imperative for decisions of this sort that institutions have fail-safe protocols that anticipate such emergencies and associated test limitations.

The patient with a dissecting aortic aneurysm clearly had a missed diagnosis, as confirmed by autopsy. This diagnosis was suspected premortem, but considered to be “ruled out” by a CT scan that did not show a dissection. Studies of the role of chest CT, particularly when earlier CT scanning technology was used, show sensitivity for dissecting aneurysm of only 83 percent.67 When we reexamined the old films for this patient, several radiologists questioned the adequacy of the study (quality of the infusion plus question of motion artifact). Newer, faster scanners reportedly are less prone to these errors, but the experience is variable. We identified another patient where a “known” artifact on a spiral CT nearly led to an unnecessary aneurysm surgery; it was only prevented by the fact that she was a Jehovah’s Witness and, because her religious beliefs precluded transfusions, surgery was considered too risky.62

The role of information transfer and the communication of critical laboratory information

Failure of diagnosis because of missing information is another theme in our weekly case reviews and the medical literature. Critical information can be missed because of failures in history-taking, lack of access to medical records, failures in the transmission of diagnostic test results, or faulty records organization (either paper or electronic) that created problems for quickly reviewing or finding needed information.

For the patient with the self-induced illness, all of the “missing” information was available online. Ironically, although many patients with a diagnosis of Munchausen’s often go to great lengths to conceal information (i.e., giving false names, using multiple hospitals), in our case, there was so much data in the computer from previous admissions and outpatient visits that the condition was “lost” in a sea of information overload—a problem certain to grow as more and more clinical information is stored online. While this patient is an unusual example of the general problems related to information transfer, this case illustrates important principles related to the need for conscientious review, the synthesizing of information, and continuity (both of physicians and information) to avoid errors.

Simply creating and maintaining a patient problem list can help prevent diagnosis errors. It can ensure that each active problem is being addressed, helping all caregivers to be aware of diagnoses, allergies, and unexplained
findings. Had our patient with “unexplained cardiomegaly” been discharged listing this as one of his problems, instead of only “pneumonia,” perhaps this problem would not have been overlooked. However, making this seemingly simple documentation tool operational has been unsuccessful in most institutions, even ones with advanced electronic information systems, and thus represents a challenge as much as a panacea.

One area of information transfer, the followup of abnormal laboratory test results, represents an important example of this information transfer paradigm in diagnostic patient safety.\textsuperscript{68–73} We identified failure rates of more than 1 in 50 for both followup abnormal thyroid tests (where the diagnosis of hypothyroidism was missed in 23 out of 982, or 2.3 percent, of patients with markedly elevated TSH results), and our earlier study on failure to act on elevated potassium levels (674 out of 32,563, or 2.0 percent, of potassium prescriptions were written for hyperkalemic patients).\textsuperscript{74} Issues of communication, teamwork, systems design, and information technology stand as areas for improvement.\textsuperscript{75–77} Recognizing this, the Massachusetts Safety Coalition has launched a statewide initiative on Communicating Critical Test Results.\textsuperscript{78}

**Physician time, test availability, and other system constraints**

Our project was based in two busy urban hospitals, including a public hospital with serious constraints on bed availability and access to certain diagnostic tests. An important recurring theme in our case discussions (and in health care generally) is the interaction between diagnostic imperatives and these resource limitations.

To what extent is failure to obtain an echocardiogram, or even a more thorough history or physical exam, understandable and justified by the circumstances under which physicians find themselves practicing? Certainly our patient with the massive cardiomegaly needed an echocardiogram at some time. Was it reasonable for the ED to defer the test (meaning a wait of perhaps several months in the clinic), or would a more “just-in-time” approach be more efficient, as well as safer in minimizing diagnosis error and delay?\textsuperscript{79} Since, by definition, we expect ED physicians to triage and treat emergencies, not thoroughly work up every problem patients have, we find complex trade-offs operating at multiple levels.

Similar trade-offs impact whether a physician had time to review all of the past records of our factitious-illness patient (only the medical student did), or how much radiology expertise is available around-the-clock to read ultrasound or CT exams, to diagnose ectopic pregnancy or aortic dissection. This is perhaps the most profound and poorly explored aspect of diagnosis error and delay, but one that will increasingly be front-and-center in health care.
Cognitive issues in diagnosis error

We briefly conclude where most diagnosis error discussions begin, with cognitive errors.45, 53, 80–84

Hindsight bias and the difficulty of weighing prior probabilities of the possible diagnoses bedeviled our efforts to assess decisions and actions retrospectively. Many “don’t miss” diagnoses are rare; it would be an error to pursue each one for every patient. We struggled to delineate guidelines that would accurately identify high-risk patients and to design strategies to prevent missing these diagnoses.

Our case reviews and firsthand interviews often found that each physician had his or her own individual way of approaching patients and their problems. Such differences made for lively conference discussions, but have disturbing implications for developing more standardized approaches to diagnosis.

The putative dichotomy between “cognitive” and “process” errors is in many ways an artificial distinction.7, 8 If a physician is interrupted while talking to the patient or thinking about a diagnosis and forgets to ask a critical question or consider a critical diagnosis, is this a process or cognitive error?

Conclusion

Because of their complexity, there are no quick fixes for diagnosis errors. As we review what we learned from a variety of approaches and cases, certain areas stood out as ripe for improvement—both small-scale improvements that can be tested locally, as well as larger improvements that need more rigorous, formal research. Table 4 summarizes these change ideas, which harvest the lessons of our 3-year project.

As outlined in the table, there needs to be a commitment to build learning organizations, in which feedback to earlier providers who may have failed to make a correct diagnosis becomes routine, so that institutions can learn from this aggregated feedback data. To better protect patients, we will need to conceptualize and construct safety nets to mitigate harm from uncertainties and errors in diagnosis. The followup of abnormal test results is a prime candidate for reengineering, to ensure low “defect” rates that are comparable to those achieved in other fields. More standardized and reliable protocols for reading x-rays and laboratory tests (such as pathology specimens), particularly in residency training programs and “after hours,” could minimize the errors we observed. In addition, we need to better delineate “red flag” and “don’t miss” diagnoses and situations, based on better understanding and data regarding pitfalls in diagnosis and ways to avoid them.
### Table 4. Change ideas for preventing and minimizing diagnostic error

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| **Upstream feedback** to earlier providers who have failed to make correct diagnosis | • Promotes culture of safety, accountability, continuous and blame-free learning, and communication  
• “Hard-wires” organizational and practitioner learning from diagnosis evolution and delays  
• Feedback from specialists poised to see missed diagnosis could be especially useful  
• Build in “feedback from the feedback” to capture reflective practitioner assessment of why errors may have occurred and considerations for future prevention  
• Permits aggregation for tracking, uncovering patterns, learning across cases, elucidating pitfalls, measuring improvements | • Logistical requirements for implementation and surveillance screening to identify errors  
• Avoiding “tampering,” from availability bias that neglects base rates (ordering aortogram on every chest pain patient to “rule out” dissection)  
• Protecting confidentiality, legal liabilities, blame-free atmosphere  
• Ways to extend to previous hospitals and physicians outside of own institution  
• To be highest leverage needs to be coupled with reporting, case conferences |
| **Safety nets to mitigate harm from diagnostic uncertainty and error** | • Well designed observation and followup venues and systems (e.g., admit for observation, followup calls or e-mail from MD in 48 hours, automated 2 wk phone followup to ensure tests obtained) for high-risk, uncertain diagnoses  
• Educating and empowering patients to have lower threshold for seeking followup care or advice, including better defining and specifying particular warning symptoms | • Logistics  
• Resource constraints (bed, test availability, clinician time)  
• Avoiding false positive errors, inappropriate use of scarce/costly resources  
• Not creating excessive patient worry/anxiety |
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| Timely and reliable **abnormal test result followup** systems | - Fail-safe, prospectively designed systems to identify which results are critical abnormalities (panic and routine), who to communicate result to, how to deliver  
- Uniform approach across various test types and disciplines (lab, pathology, radiology, cardiology)  
- Leverage information technologies to automate and increase reliability  
- Involve patients to ensure timely notification of results, and contacting provider when this fails  
- Modeled on Massachusetts Patient Safety Coalition program 78 | - Achieving consensus on test types and cut-off thresholds  
- On-call, cross-coverage issues for critical panic results  
- Defining responsibilities: e.g., for ED patients, for lab personnel  
- Practitioner efficiency issues: avoiding excess duplicate work; efficient documentation |
| **Fail-safe protocols for preliminary/resident and definitive readings of tests** | - Must be well-defined and organized system for supervision, creating and communicating final reports  
- Need for system for amending and alerting critical changes to clinicians  
- "After-hours" systems for readings, amending | - How to best recognize and convey variations in expertise of attendings who write final reports  
- Quality control poses major unmet challenges |
| **Prospectively defining red flag diagnoses and situations and instituting prospective readiness** | - Create "pull" systems for patients with particular medical problems to ensure standardized, expedited diagnostic evaluations (so don't have to "push" to quickly obtain)  
- Like AMI thrombolytic "clot box," in-place for ready activation the moment patient first presents  
- Embodies/requires coordinated multidisciplinary approach (e.g., pharmacy, radiology, specialists) | - Difficulties in evidence-based delineation of diagnoses, situations, patient selection, criteria, and standardized actions  
- Avoiding excessive work-up, diverting resources from other problems |
Table 4. Change ideas for preventing and minimizing diagnostic error, cont.

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| Automated screening check lists to avoid missing key history, physical, lab data | • Less reliance on human memory for more thorough questioning  
• Queries triggered by individual patient features  
• Could be customized based on presenting problem (i.e., work exposures for lung symptoms, travel history for fever) | • Evidence of efficacy to-date unconvincing; unclear value of unselective “review of systems”  
• Sorting out, avoiding false positive errors from data with poor signal:noise ratio  
• Effectively implementing with teamwork and integrated information technology |
| High-level patient education and engagement with diagnosis probabilities and uncertainties | • Since diagnosis often so complex and difficult, this needs to be shared with patients in ways to minimize disappointments and surprises  
• Support and enhance patients taking initiative to question diagnosis, particularly if not responding as expected | • Potentially more time-consuming for practitioners  
• Avoiding unnecessary testing  
• Doing in way that balances need for preserving patient confidence in their physicians and advice, with education and recognition of diagnosis fallabilities |
| Test and leverage information technology tools to avoid known cognitive and care process pitfalls | • Better design of ways to streamline documentation (including differential diagnosis) and access/display of historical data  
• Easing documentation time demands to give practitioners more time to talk to patients and think about their problems  
• Facilitating real-time access to medical knowledge sources  
• Sophisticated decision support tools that use complex rules and individualized patients data  
• Prompts to suggest consideration of medication effects in differential, based on linkages to patient’s medication profile, lab results | • Shortcomings of first generation of “artificial intelligence” diagnosis software  
• Challenges coupling knowledge bases with individual patient characteristics  
• Paucity of standardized, accepted, sharable clinical alerts/rules  
• New errors and distractions introduced by intrusion of computer into clinical encounter  
• Alleged atrophy of unaided cognitive skills |
To achieve many of these advances, automated (and manual) checklists and reminders will be needed to overcome current reliance on human memory. But information technology must also be deployed and reengineered to overcome growing problems associated with information overload. Finally, and most importantly, patients will have to be engaged on multiple levels to become “coproducers” in a safer practice of medical diagnosis. It is our hope that these change ideas can be tested and implemented to ensure safer treatment based on better diagnoses—diagnosis with fewer delays, mistakes, and process errors.

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