6-29-2015

Back to the Bedside: Developing a Bedside Aid for Concussion and Brain Injury Decisions in the Emergency Department

Edward R. Melnick
Yale School of Medicine, edward.melnick@yale.edu

Kevin Lopez
University of Connecticut, kevin.lopez.91@gmail.com

Erik P. Hess
Mayo Clinic, Hess.Erik@mayo.edu

Fuad Abujarad
Yale School of Medicine, fuad.abujarad@yale.edu

See next pages for additional authors

Follow this and additional works at: http://repository.edm-forum.org/egems

Part of the Emergency Medicine Commons, Health Information Technology Commons, and the Trauma Commons

Recommended Citation
DOI: http://dx.doi.org/10.13063/2327-9214.1136
Available at: http://repository.edm-forum.org/egems/vol3/iss2/6

This Informatics Case Study is brought to you for free and open access by the the Publish at EDM Forum Community. It has been peer-reviewed and accepted for publication in eGEMs (Generating Evidence & Methods to improve patient outcomes).

The Electronic Data Methods (EDM) Forum is supported by the Agency for Healthcare Research and Quality (AHRQ), Grant 1U18HS022789-01. eGEMs publications do not reflect the official views of AHRQ or the United States Department of Health and Human Services.
Back to the Bedside: Developing a Bedside Aid for Concussion and Brain Injury Decisions in the Emergency Department

Abstract

Context: Current information-rich electronic health record (EHR) interfaces require large, high-resolution screens running on desktop computers. This interface compromises the provider’s already limited time at the bedside by physically separating the patient from the doctor. The case study presented here describes a patient-centered clinical decision support (CDS) design process that aims to bring the physician back to the bedside by integrating a patient decision aid with CDS for shared use by the patient and provider on a touchscreen tablet computer for deciding whether or not to obtain a CT scan for minor head injury in the emergency department, a clinical scenario that could benefit from CDS but has failed previous implementation attempts.

Case Description: This case study follows the user-centered design (UCD) approach to build a bedside aid that is useful and usable, and that promotes shared decision-making between patients and their providers using a tablet computer at the bedside. The patient-centered decision support design process focuses on the prototype build using agile software development, but also describes the following: (1) the requirement gathering phase including triangulated qualitative research (focus groups and cognitive task analysis) to understand current challenges, (2) features for patient education, the physician, and shared decision-making, (3) system architecture and technical requirements, and (4) future plans for formative usability testing and field testing.

Lessons Learned: We share specific lessons learned and general recommendations from critical insights gained in the patient-centered decision support design process about early stakeholder engagement, EHR integration, external expert feedback, challenges to two users on a single device, project management, and accessibility.

Conclusions: Successful implementation of this tool will require seamless integration into the provider’s workflow. This protocol can create an effective interface for shared decision-making and safe resource reduction at the bedside in the austere and dynamic clinical environment of the ED and is generalizable for these purposes in other clinical environments as well.

Acknowledgements

We would like to thank the Mayo Clinic Knowledge & Evaluation Research Unit, North Shore/LIJ Department of Internal Medicine, University of Maryland, Human-Computer Interaction Lab, Oregon Health & Science University-Human Computer Interaction in Biomedicine course instructors, Pediatric Emergency Care Applied Research Network investigators, and Epic’s Research & Development team for taking the time to view demonstrations of our prototype and provide feedback to improve our final product.

Keywords

Methods, Informatics, Quality Improvement

Disciplines

Emergency Medicine | Health Information Technology | Trauma

This case study is available at EDM Forum Community: http://repository.edm-forum.org/egems/vol3/iss2/6
Back to the Bedside: Developing a Bedside Aid for Concussion and Brain Injury Decisions in the Emergency Department

Edward R. Melnick, MD; Kevin Lopez, MS; Erik P. Hess, MD; Fuad Abujarad, PhD; MSc; Cynthia A. Brandt, MD, MPH; Richard N. Shiffman, MD, MCIS; Lori A. Post, PhD

Abstract

Context: Current information-rich electronic health record (EHR) interfaces require large, high-resolution screens running on desktop computers. This interface compromises the provider’s already limited time at the bedside by physically separating the patient from the doctor. The case study presented here describes a patient-centered clinical decision support (CDS) design process that aims to bring the physician back to the bedside by integrating a patient decision aid with CDS for shared use by the patient and provider on a touchscreen tablet computer for deciding whether or not to obtain a CT scan has failed previous implementation attempts.

Case Description: This case study follows the user-centered design (UCD) approach to build a bedside aid that is useful and usable, and that promotes shared decision-making between patients and their providers using a tablet computer at the bedside. The patient-centered decision support design process focuses on the prototype build using agile software development, but also describes the following: (1) the requirement gathering phase including triangulated qualitative research (focus groups and cognitive task analysis) to understand current challenges, (2) features for patient education, the physician, and shared decision-making, (3) system architecture and technical requirements, and (4) future plans for formative usability testing and field testing.

Lessons Learned: We share specific lessons learned and general recommendations from critical insights gained in the patient-centered decision support design process about early stakeholder engagement, EHR integration, external expert feedback, challenges to two users on a single device, project management, and accessibility.

Yale School of Medicine, University of Connecticut, Mayo Clinic
**Introduction**

Counterintuitively, in an information-rich world advances in technology can increase, not decrease, cognitive demands on users. In the rush to adopt electronic health records (EHRs) to qualify for federal incentive payments, clinicians now find themselves working with products with poor usability that are neither integrated nor interoperable into the clinical workflow. Since computerized clinical decision support (CDS) is most effective when integrated as part of the physician’s normal workflow at the time and location of decision-making, the potential patient safety and outcome benefits of CDS have not yet been fully realized. Furthermore, current information-rich EHR interfaces require large, high-resolution screens running on desktop computers. This interface compromises the physician’s already limited time at the bedside by physically separating the patient and the doctor.

We developed a possible solution to these interface challenges in a clinical scenario that could benefit from CDS but had failed previous implementation attempts. Diagnostic imaging is the fastest growing segment of health care spending in the United States, increasing twice as fast as total health care costs. In the emergency department (ED), use of advanced diagnostic imaging in injured patients has increased dramatically—leading to increased health care costs, exposure to ionizing radiation, and length of stay without objective metrics of improved patient outcomes. In particular, despite implementation of validated, highly sensitive clinical guidelines designed to safely reduce the use of computed tomography (CT) in minor head injury, CT is frequently obtained in low-risk, minor head injury patients in whom it is not clinically indicated.

This case study describes a patient-centered CDS design process that aims to bring the physician back to the patient’s bedside by integrating a patient decision aid with CDS for shared use by the patient and provider using a touchscreen tablet computer (Figure 1). Furthermore, newer generation tablet computers are flat, portable, and potentially less likely to cause hospital-acquired infections than desktop computers as they can be more easily sanitized since they do not house internal fans. The objective of this case study was to do the necessary foundational work to uncover and disentangle the human and environmental factors, as well as the chaotic clinical workflow, and address them in the design process such that the eventual CDS interface can more effectively support the physician at the point of care. The tool described in this case study has been prototyped and will subsequently undergo usability and field-...
testing, will be interfaced with our institution’s EHR, and will be studied in an implementation trial to demonstrate that technology is better than no technology for patient engagement and safe reduction of diagnostic imaging in the ED—an austere and dynamic clinical environment, where the physician is faced with a substantial volume of high acuity patients, time pressures, and interruptions.31-34

**Case Description**

**Background**

More than 1.3 million patients are treated annually in United States EDs for traumatic brain injury.35 Most of these injuries are mild, but in a small proportion of patients with mild injury, clinical deterioration occurs.36 In patients with clinically important traumatic brain injury, CT imaging yields a quick and accurate diagnosis such that neurological intervention can prevent deleterious outcomes from intracranial hematoma. Although CT has greatly improved our diagnostic ability, it exposes patients to significant amounts of ionizing radiation.37 In addition, over 90 percent of CT scans for minor head injury are negative for clinically important brain injury.38

The Canadian CT Head Rule (CCHR) is a *clinical decision rule* that was developed using a rigorous, evidence-based derivation and validation process to identify appropriate use of CT to differentiate mild traumatic brain injury from clinically important brain injury.38 In both Canada and the United States, the CCHR has been validated to be 100 percent sensitive and more specific than other guidelines and decision rules.38,40-43 A prospective cluster-randomized trial to implement a similar prediction rule—the Canadian C-spine Rule—led to a significant decrease in imaging.36 When the CCHR was implemented at the same centers with
many of the same patients, however, CT imaging rates did not decrease. In fact, imaging rates were 74–76 percent with the implementation, compared to 63–68 percent without it. These rates were more than double compared to 12 years earlier in the same region. The authors suspected that “CT imaging has become the local standard of care for patients with minor head injury...[and has] led to expedient over-testing.” The CCHR failed to reduce testing due to implementation failures not rule performance. Specifically, compliance with the CCHR could safely decrease the number of CT scans performed in minor head injury by 35 percent. If the CCHR were successfully implemented in the United States, a significant number of radiation-induced cancer deaths could be averted with a cost savings of up to $394 million annually. Until workflow barriers, patients’ values and preferences, and how they affect decision-making regarding use of CT are addressed in this clinical scenario, these patients will continue to be exposed to undue radiation risk and cost.

**Patient-Centered Decision Support Design Process**

We followed a user-centered design (UCD) approach—an iterative, multistage user interface design and evaluation process—to build a bedside aid that is useful and usable, and that promotes shared decision-making between patients and providers at the patient’s bedside on a tablet computer. First, a triangulated qualitative study was performed to identify the factors that either promote or inhibit the appropriate use of CT in patients presenting to the ED with minor head injury.

The findings of this qualitative research have been reported elsewhere. Briefly, seven focus groups were performed—three with exclusively providers and four with exclusively patients. Understanding that users may “have very limited insight into their own performance, and even more limited ability to articulate what might improve it,” we triangulated the focus group findings with direct field observation in the form of a cognitive task analysis. The analysis included more than 150 hours of direct observation in the ED of peer-nominated, senior emergency-physician subject matter experts (SMEs) in safely minimizing testing in the ED via patient engagement and subsequent critical decision method interviews with the SMEs. The qualitative research findings included nonclinical human factors in six primary domains of establishing trust, bedside manner, anxiety (of both the patient and the provider), constraints (e.g., time), the influence of others (e.g., other providers, patient family members, Internet), and patient expectations. These results informed the conclusion that identifying and disseminating approaches and designing systems that help clinicians establish trust and manage uncertainty within the ED context could optimize CT use in minor head injury. To our knowledge, such patient-centered themes have not previously been considered up front in the CDS design process.

Qualitative factors identified during the focus groups and cognitive task analysis were integrated in subsequent design and development of a prototype tool to formulate the initial CDS concept. Although there are many potential options for the specific details of a tool’s interface such that it may eventually be seamlessly integrated into provider workflow, the initial interface must function within the constraints of what typically occurs at the bedside in a patient encounter—e.g., in the ED, patient arrival, waiting for provider, provider evaluation, diagnostic work-up if necessary, and patient-provider discussion regarding patient disposition (i.e., discharged home, transferred to another facility, admitted to the hospital). The tool must facilitate a patient-focused workflow for the provider. Therefore, the prototype was designed to include the following: (1) information for the patient to review while waiting for the provider and awaiting...
diagnostic work-up and discharge (akin to a patient decision aid) and (2) decision support with content for the provider to complete at the bedside in discussion with the patient.

A rapid prototyping model was used following the International Patient Decision Aid Standards and the agile software development approach, which provides the flexibility to make changes from feedback on usability in addition to rapid prototyping. Given the constant evolution of available technology, the prototype was programmed to be device agnostic: capable of running on any modern device regardless of its operating system or form factor. The prototype’s interface evolved through many development cycles. To develop the initial prototype, a multidisciplinary team (including several clinical informaticists with a variety of clinical backgrounds, a systems architect, a computer programmer, key stakeholders from the health system’s information technology (IT) leadership, and potential users) reviewed the factors identified in the requirements-gathering phase, resulting in a “rough draft” prototype. Next, the flexible development process was enhanced by eliciting feedback from end users at every stage. The initial prototype was developed for demonstration and feedback from multiple audiences (including patient decision aid designers, internal- and emergency-medicine physicians, a human computer interaction class for physicians with a variety of clinical backgrounds and experiences, and key stakeholders from our institution’s EHR vendor) on its content, format, and usability as well as its potential to increase patient knowledge and address patient concerns, values, and preferences. This feedback was used to iteratively modify the prototype.

This process is ongoing. At the time of this writing, the next step planned is for the prototype to undergo formative usability testing to maximize its efficiency, effectiveness, and user satisfaction. Usability evaluations will be conducted with representative patients and providers (end users) to assess the degree to which the prototype tool matches their needs for shared decision-making and workflow. To optimize the adoption of this tool in the complex, high-pressure environment of the ED warrants special attention. In this complex sociotechnical system, maximizing the human-computer interaction is not sufficient. The human-environment interaction, or ecology, must also be taken into account. It is crucial that CDS makes work easier for the provider—otherwise providers will not use the tool, will experience information overload and alert fatigue, and will try to ignore or circumnavigate the CDS in order to get other tasks done. Field-testing will rely on the principles of ecological interface design to provide the right information at the right time and in the right way while considering the demands of the ED work environment.

Features

The application was designed to allow users (both patient and provider) smooth, user-friendly navigation through the screens while completing the tasks of patient education, risk communication, and shared decision-making about whether or not to perform a CT. The application is equipped with features to educate and empower patients with knowledge that facilitates expressing concerns. When the provider arrives at the bedside, the patient can make an informed decision, and the provider can efficiently address the patient’s concerns.

The patients fill out two forms when they first receive the tablet. The first is an eligibility form. If the patient is eligible, the patient then continues on to a questionnaire form. The answers to the questionnaire autopopulate the subjective components of the clinical decision rule later on in the provider workflow.
to streamline the provider’s time at the bedside, thus maximizing the opportunity for a conversation regarding the patient’s specific concerns. Following the forms, the tool is divided into three sections: (1) a patient education section, (2) a physician’s section, and (3) a shared decision-making section. These sections all follow a visual metaphor using a design reminiscent of a decision aid on paper cards.

The tool’s patient education section contains information for the patient about concussion, more severe brain injuries, and CT scans. The information provided on each card is focused and simplified, given the wide range of education backgrounds of potential users as well as the presence of a recent head injury. If the patient wants to learn more, there is the option to get more detailed information on specific questions on each topic. With the goal of maintaining the patient’s attention, the information section was designed so the patients can read in any order they choose. In this section, the patients are also given the opportunity to flag their specific concerns for future discussion with their providers (Figure 2).

On entering the patient’s room, the provider logs in and is prompted to go through the CCHR clinical decision rule with the patient. The inputs to the rule prompt a risk communication conversation by providing patient-specific risk estimates of any brain injury on CT, clinically important brain injury, the need for neurosurgical intervention, and risk of cancer from a head CT. The “What is best for YOU” screen in the provider section (Figure 3) engages the patient by addressing issues that we identified in our qualitative research—such as identifying and

Figure 2. Card Allowing Patient to Communicate Their Concerns with the Provider
addressing patient concerns, establishing trust, and managing patient uncertainty. This section also has a card that appears only if the patient is 65 years or older and addresses issues specific to older adults.

The decision area is where the patient and the provider come together to make a shared decision. Three cards provide the details of three choices: (1) get a CT, (2) to go home now with active surveillance for new or concerning symptoms, or (3) stay in the ED for observation (available only to patients under age 65 due to the increased risk of a slow-bleeding subdural hemorrhage). A doctor’s note includes the patient-specific risk estimates and documents the shared decision-making conversation. For example, the following shared decision-making note is generated for a low risk patient who does not undergo CT imaging:

I have used a decision aid to share decision-making with the patient about whether or not to get a CT scan for a minor head injury. This patient’s injury is low risk based on the Canadian CT Head Rule. We estimated the patient’s risks as follows: (1) need for neurosurgical intervention to be 0.0%, (2) clinically important brain injury to be 1.1%, (3) any brain injury by CT to be 2.7%, and (4) lifetime risk of cancer from a CT scan today to be 0.007%. After considering the patient’s unique circumstances and the pros and cons of the alternatives, we decided the patient should go home now without a CT.

This note will push to the patient’s chart in the EHR along with the ability for the provider to push an order for a CT or discharge instructions depending on the ultimate results of the shared decision.

**Figure 3. The Provider Sees the Preidentified Patient Concerns and Discusses Them with the Patient**
System Database Architecture

The prototype phase also included creation of a database to collect, edit, store, and retrieve data generated by the tool. The database grew throughout the rapid prototyping process as the application’s needs expanded. Originally the model view controller framework provided some tables that handle user authentication. As the application grew, new tables were added for additional functionality. At the end of the prototyping phase, the database had six tables (Figure 4).

Technical System Requirements. The application was developed and tested on a machine that was running Internet Information Services 8 (running on Microsoft Windows 6.2). This machine had 16 GB of memory and a 4.5 GHz octa-core processor. Currently, the Web service is hosted on a machine that has Windows 2012 R2 Datacenter edition with two processors. Our server used 4GB of RAM and...
a 250GB hard disk. While this machine uses fewer resources, these specifications allow the application to run smoothly. We used similar specifications for the database server. With these modest specifications the application runs smoothly and yet does not utilize all of the server’s resources.

**Lessons Learned**

Specific lessons gained from the patient-centered decision support design process described above are provided in Table 1 along with generalized, actionable recommendations such that future design and implementation efforts might benefit from the critical insights gained from this process. Given the literature available on similar topics, this list of lessons is by no means exhaustive; rather, it is intended to provide key pitfalls and successes based on our experiences that may not yet appear in the literature or would most benefit an early stage developer.

Early stakeholder involvement is a key tenet in implementation science and change management. A common pitfall in implementation science is when stakeholders resist change due to lack of engagement early on in the change process. Stakeholders for this project include groups such as ED patients with minor head injury, ED providers locally (and beyond should the implementation succeed locally), departmental leadership, health system IT leadership and informatics work force, clinical informatics research community, and the EHR vendor. Engagement of health system IT leadership and patients are both challenging and crucial for success. Therefore, we placed special emphasis on recommendations to engage these stakeholders early and throughout the development process.

Provider stakeholders have made it clear in the requirement-gathering phase that they will use a tool if it can streamline their workflow. Therefore, we are working to integrate the tool within our institution’s EHR such that it can facilitate CT order entry for the provider and generate discharge instructions for the patient (patient handout on concussion). We are working with our health system’s IT leadership and our institution’s EHR leadership to optimize this interoperability. Lessons learned thus far in EHR integration are both refreshing and frustrating. Evolution of the EHR interface is driven by market pressure for functionality over usability. Despite the barriers to receiving vendor-specific training coursework for research applied to their interface, there are large initiatives on the vendor’s part to continue to improve EHR usability with a patient-centered focus.

During the prototype development process, we demonstrated the prototype to outside expert groups (in human-computer interaction, health IT usability research, and decision aids) to seek feedback. These outside groups were particularly helpful to set goals in innovative areas of inquiry within the patient-centered decision support design process. For example, traditional usability evaluations test a single user on a single device. Usability evaluation of our tool presents unique challenges to traditional usability evaluation techniques since our tool will be used by two users on a single device. To address this challenge, we have developed a standardized protocol for usability testing that evaluates one user type and simulates the other user with a standardized script. For example, real physician users will test the tool with standardized patients in a simulated clinical environment. The decision aid expert group helped us to appreciate that the tool’s primary goal for both evaluation and implementation will be its ability to facilitate high quality conversation between the patient and provider.
Table 1. Lessons Learned and Recommendations to Facilitate CDS Development and Interoperability

<table>
<thead>
<tr>
<th>LESSONS LEARNED</th>
<th>RECOMMENDATIONS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Early stakeholder engagement</strong></td>
<td>Early engagement of health system IT leadership allowed introduction of project’s goals, networking with leaders locally, and access to EHR training coursework that will support integration of tool with EHR.</td>
</tr>
<tr>
<td>Prototype pilot testing addressed gaps in instructions provided to new users.</td>
<td>Pilot test prototype with new users prior to usability testing to identify gaps that developers may overlook due to familiarity with their tool.</td>
</tr>
<tr>
<td>Patient input is challenging. A patient advisory committee was recruited from focus group participants for future input on tool development.</td>
<td>Involve patient end users at every stage of development. There is a growing community of patient representatives, advocates, and volunteers who could help provide this input.</td>
</tr>
<tr>
<td><strong>EHR integration</strong></td>
<td>Despite physician user frustration with EHR usability, vendors are developing patient-centered interfaces, and they know who else is doing similar work and how best to integrate new tools with their software.</td>
</tr>
<tr>
<td>Other researchers have worked on similar integration challenges. One group integrated a Web service into provider decision support workflow with our EHR vendor. As a result, the vendor has incorporated the ability to access a Web service from within the EHR as a standard for the latest version of their software.</td>
<td>1. Network with EHR vendors to find people whose goals are aligned with yours. 2. Epic’s App Exchange and Developer Guide are a good starting point.</td>
</tr>
<tr>
<td>EHR vendors create barriers to research applied to their interface. There are many logistical challenges to completing vendor-specific EHR training to learn skills to do this type of research.</td>
<td>1. Build off of experiences of others in the field. 2. Programming a Web service that can be accessed from within the EHR can maintain EHR functionality and allow the possibility of being device- and platform agnostic.</td>
</tr>
<tr>
<td></td>
<td>Begin EHR vendor training coursework early, and anticipate delays and challenges to coursework completion.</td>
</tr>
</tbody>
</table>
Table 1. Lessons Learned and Recommendations to Facilitate CDS Development and Interoperability (Cont’d)

<table>
<thead>
<tr>
<th>LESSONS LEARNED</th>
<th>RECOMMENDATIONS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Feedback from prototype demonstrations to outside expert groups</strong></td>
<td>Tool will achieve its goals only if it can promote conversation between patient and provider.</td>
</tr>
<tr>
<td></td>
<td>It is more difficult to make changes to a computerized prototype if programming is started too early in the development process.</td>
</tr>
<tr>
<td><strong>Challenge of having two users</strong></td>
<td>Best practices for authentication of two users for a single tool on a single device are not established. Double user authentication is particularly difficult given HIPAA regulations.</td>
</tr>
<tr>
<td></td>
<td>Usability testing methods for two users on a single device is not established—limiting data collection options for both users using conventional usability evaluation methods and software.</td>
</tr>
<tr>
<td><strong>Project management</strong></td>
<td>A high volume of programming specifications can be difficult to track and prioritize.</td>
</tr>
<tr>
<td><strong>Accessibility</strong></td>
<td>Section 508 of the Rehabilitation Act of 1973 in the United States requires all federally funded information technology (IT) to be accessible to people with disabilities.</td>
</tr>
</tbody>
</table>
Conclusions

This case study describes a novel patient-centered decision support design process that aims to bring the physician back to the patient’s bedside by integrating a patient decision aid with CDS for shared use by the patient and provider on a touchscreen tablet computer for deciding whether or not to obtain a CT scan for minor head injury in the ED, a clinical scenario that could benefit from CDS but has failed previous implementation attempts. The study focuses on the prototype building process in the context of requirement gathering, usability- and field-testing, and EHR integration, and a future implementation trial to demonstrate that some technological support is better than no technology for patient engagement and safe reduction of diagnostic imaging in the ED. Once the cycle of development and testing is complete, we aim to prospectively test the effect of using the tool in ED patients with minor head trauma. The ultimate objective of developing this patient-centered decision support tool is that it will engage patients in their care and safely reduce CT use in minor head injury patients in the ED. The tool will facilitate the patient and provider in engaging in transparent, informed, shared decision-making regarding risk communication and CT use in minor head injury. It will be seamlessly integrated into the provider’s EHR workflow including automatically generating a note for the patient’s chart on the shared decision-making conversation, thereby simultaneously streamlining the provider’s workflow and respecting the patient’s preferences.

The current challenges to EHR usability reflect the vendor market’s pressure for functionality over usability.45,69 It has been argued that current EHRs contain vast amounts of data and information hidden in unreadable interfaces and that “no system of data management will ever replace...good medicine.”71 This conclusion fails to recognize the trajectory of innovation that EHR usability challenges represent. In Ackoff’s seminal paper, “From Data to Wisdom,” he differentiates data and information from knowledge.2 Data is raw material; it exists in isolation without significance. Information is data that has been given meaning but may or may not be useful; whereas knowledge is the collection of information with the intent to be useful. We believe that medicine is only in the dawn of its information age; the smart phone’s pervasiveness is evidence that as a society we are well into a knowledge age. The complexity of the changing health care system locally, regionally, and nationally and the rapid growth of knowledge in complex fields all delay EHR usability. As medicine catches up to available technologies, future electronic systems must not only be usable, but must also support knowledge and promote conversation between patients and their doctors at the bedside.

Acknowledgements

We would like to thank the Mayo Clinic Knowledge & Evaluation Research Unit, North Shore/LIJ Department of Internal Medicine, University of Maryland, Human-Computer Interaction Lab, Oregon Health & Science University-Human Computer Interaction in Biomedicine course instructors, Pediatric Emergency Care Applied Research Network investigators, and Epic’s Research & Development team for taking the time to view demonstrations of our prototype and provide feedback to improve our final product.

References


