

Patient Experiences Using an Inpatient Personal Health Record

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Summary

Objective: To investigate patients' experience using an inpatient personal health record (PHR) on a tablet computer to increase engagement in their hospital care.

Methods: We performed observations and conducted semi-structured interviews with 14 post-operative cardiac surgical patients and their family members who received an inpatient PHR. Themes were identified using an inductive coding scheme.

Results: All participants responded favorably to having access to view their clinical information. A majority (85.7%) of participants used the application following an initial training session. Patients reported high satisfaction with being able to view their hospital medications and access educational materials related to their medical conditions. Patients reported a desire to view daily progress reports about their hospital stay and have access to educational information about their post-acute recovery. In addition, patients expressed a common desire to view their diagnoses, laboratory test results, radiology reports, and procedure notes in language that is patient-friendly.

Conclusion: Patients have unmet information needs in the hospital setting. Our findings suggest that for some inpatients and their family members, providing personalized health information through a tablet computer may improve satisfaction, decrease anxiety, increase understanding of their health conditions, and improve safety and quality of care.

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1. Background

Being a patient in the hospital is an anxiety-inducing experience. Poor communication and lack of access to information can have detrimental effects on a patient's psychological wellbeing and coping abilities [1–5]. Policymakers [6, 7] and patient advocates [8, 9] are increasingly focusing on this problem. This study investigates how an inpatient personal health record (PHR) may help address the information needs of patients and improve their engagement and experience in the hospital.

Provision of medical information and education about care can allay a patient's uncertainty and stress [10–15]. Further, sharing information has been shown to reduce patient requests for clinicians' time, foster a sense of control and empowerment while promoting self-efficacy and ownership of illness [16–18], and lead to increased participation as an informed partner in making health decisions [19–22]. Previous research suggests that patients want to use PHRs and believe that they will be valuable [23]. Moreover, patients believe that their adherence to treatment regimens will improve if they have the opportunity to read and understand their health information [24, 25]. However, patients are not typically given the opportunity to review their health information in the hospital. For example, in a survey of hospitalized patients conducted by Cumbler and colleagues, 90% of respondents said they wanted to review their hospital medication list for accuracy, but only 28% said they were given the opportunity to do so [26]. Without access to a documented care plan, patients are uncertain as to the care they have received and what to expect. For example, studies show that patients remember less than half of what physicians explain to them in the hospital [27, 28]. These findings suggest that there is potential to expand data-sharing practices with hospitalized patients in order to improve patient experience.

In previous work, we conducted a small pilot study where we provided a prototype of an inpatient PHR to five hospitalized patients in a supervised manner to assess their initial impressions and interest [38]. Using tablet computers, patients viewed photos of their care teams, medications they received, and their reconciled home medications and allergies. The response of patients and health-care providers was positive [38, 39], demonstrating that the inpatient PHR could be a useful platform for addressing patients' information needs in the hospital. Based on the feedback we received from this earlier study, we enhanced the inpatient PHR system to address a number of limitations. The present study describes a larger deployment of the system and assesses patients' perceptions and information needs after having access to the technology without supervision for at least 24 hours during their hospital stay.

2. Objectives

The three aims of our study were to assess (1) the factors surrounding patients' use of an inpatient PHR, (2) patients' experience with the application, and (3) patients' information needs that might be partially met using an inpatient PHR.

3. Methods

3.1. Setting and Technology

In 2009, NewYork-Presbyterian Hospital (NYP) / Columbia University Medical Center (CUMC), a large urban academic medical center in New York City, implemented an outpatient portal called myNYP.org. In addition to functions such as the ability to search for NYP physicians, schedule appointments, and pay bills, the system allowed patients to access their laboratory test results, radiology reports, operative notes, and discharge summaries after leaving the hospital. In 2011, we developed a version of myNYP specifically for hospitalized patients. The initial inpatient PHR allowed patients and their families to view their care team (names, photos, and biographies), confirm allergies and home medications, and track hospital medications.

Based on the feedback from an earlier pilot study, we enhanced the inpatient PHR to increase usability and provide feature enhancements. These enhancements included user interface and navi-

gation improvements, electronic self-reporting of pain level, and a feature to record notes, comments and questions for health providers that were made available to the care team within the hospital's electronic health record. Additionally, we made available links to medication information on MedlinePlus.gov, as well as links to curated educational content on myNYP.org, such as educational videos and explanations of medical conditions. A sample screen of the patient comment interface is shown in ► Figure 1.

3.2. Study Design

Patients and their family members from a post-operative cardiac surgery unit at NewYork-Presbyterian/Columbia University Medical Center were invited to participate in the study from August to December 2012. English-speaking participants were recruited based on consultation with the attending physician managing post-operative patients on the floor. Exclusion criteria included patients with severe mental illness, substance abuse, language barriers or severe physical discomfort. This study was approved by the Columbia University Medical Center Institutional Review Board.

After providing informed consent, participants were given an Apple iPad tablet computer with access to the inpatient PHR application. Initial use of the application was observed for 15 minutes to assess usability, answer questions, provide guidance on navigation of application, and collect initial impressions. The following day, participants participated in a semi-structured interview that took on average 20–30 minutes. The interview questions consisted of 23 questions based on core concepts from the Technology Acceptance Model [40]. To ensure accuracy of interpretations of interview data, we reflected back to participants our understanding of what we heard them reporting during the interviews [41]. Questions covered patients' demographics, technological adeptness, health information seeking behavior, knowledge of medical conditions, previous use of PHRs, information needs while in the hospital, and experience and use of the inpatient PHR application (see Appendix). Interviews were audio-recorded and transcribed by members of the research team, who met weekly to review general findings and themes of collected data. Participant recruitment continued until we reached thematic data saturation, i.e., when no new general themes were observed [42].

3.3 Data Analysis

Using a qualitative content analysis approach, two members of the research team (JW and AS) independently reviewed all the data and through the process of consensus building generated the coding scheme. They independently coded the data and discussed to clarify meaning and boundary of codes to finalize themes [43]. Participant demographics and self-reported usage of application were analyzed using descriptive statistics.

4. Results

Twenty-four patients were approached, twenty patients consented to participate in the study, and fourteen completed the semi-structured interviews (► Table 1). The mean age of the participants was 59.0 (SD=16.0), with an equal split between male and female. The majority of participants (57.1%) had at least some college education. Almost all participants (92.9%) reported using the Internet on a daily basis and 21.4% had previous experience using a tablet computer. Three of the fourteen patients (21.4%) used the inpatient PHR along with a family member.

Half of the participants (50.0%) stated that they kept a physical file of their medical records at home. Nearly three-quarters of participants (71.4%) reported searching online for health information from consumer health websites. Two of the fourteen patients (14.3%) reported having access to their personal health information via a portal maintained by their provider. One patient reported having a digital copy of his records provided by his physician on a USB drive.

4.1 Application Usage

Most participants (85.7%) reported using the application during the period after the initial observation session (► Table 2). The majority of participants (57.1%) reported not using the tablet computer for other purposes (e.g., checking email, social media, browsing the Internet), even though they were

invited to do so. The main factor patients reported that influenced their use of the application was their well-being. Patients reported feeling less inclined to use the application if they were experiencing pain or nausea, or had general malaise. In addition, patients who underwent multiple tests and/or procedures reported less use. In several cases where the patient was too ill or otherwise disinclined to participate, a family member reported using the application (with the patient's permission).

In the small sample of participants, there were no apparent socio-demographic variables that impacted tablet usage. For example, there was no significant difference in application usage between female and male patients, and neither education level nor age played a role. However, we observed that patients who owned and used a tablet device previously had an easier time using the tablet and navigating through the application. Although most patients reported that the application was easy-to-use, during initial observations, it was noted that about half had difficulty operating the tablet computer. After about 15 minutes of instruction, each of these participants reported feeling comfortable navigating the application.

4.2 Patients' Experience with the Inpatient PHR

Ten themes emerged from analysis of interviews: 1) desire to understand and engage, 2) desire for status report, 3) report test results, 4) ownership of data, 5) desire for guidance, 6) inpatient PHR serves as information and memory aid, 7) improves safety, 8) well-being affects use, 9) messaging providers: mixed response, 10) self-reporting pain: mostly unfavorable. Definitions of these themes and example quotes are further detailed in ► Table 3.

4.3 Hospitalized Patients' Information Needs

Thirteen of fourteen patients in the study (92.9%) wanted more information about their hospital care, and the same number stated that the tablet-based inpatient PHR was helpful in addressing their information needs and valuable in increasing their understanding of their hospital stay. Some patients said they believed they should be given access to review any part of their medical record during their hospital stay, even if they cannot fully understand it. Though most participants pushed for greater access, 28.6% expressed hesitation about seeing sensitive material (e.g., a terminal prognosis) before their doctor could share it with them at an appropriate time. They felt safeguards should be put in place for sensitive information. Nearly half of participants (42.8%) believed that in some instances, the application could replace face-to-face conversations with their providers.

4.3.1 Information patients found most useful in the inpatient PHR

Of the information provided in the inpatient PHR, the information reported as most useful was the list of hospital medications administered and links to educational materials. Specifically, 85.7% of patients found the medication names and links to MedlinePlus.gov helpful for participating in and understanding their hospital care. For example, a potential medical error was prevented when a family member recognized a dosing error while using the application and notified the provider. Most participants also indicated that they liked seeing photos and short bios of their care team members as well.

4.3.2 Additional information patients wanted

In addition to what was accessible in the inpatient PHR, participants expressed interest in seeing their progress notes (85.7%), operative reports (71.4%), documented medical conditions (50%), laboratory test results (35.7%), and radiology reports (35.7%). Participants were particularly interested in test results that were noted by their physicians to be important for discharge. Most participants desired more tailored information in the inpatient PHR, such as the following: a personalized care plan, a daily progress report, intended discharge date, key items providers needed to approve discharge, and a daily schedule (e.g., tests, procedures, and therapy sessions). Patients reported that this would help give them a sense of control over their day and help them plan when their guests should visit. Most participants requested educational information about their medical conditions and surgical procedure, along with self-care instructions and what to expect after discharge. In addition, a few

patients wanted to be able to view information about their hospital bill and follow-up doctors appointments.

All participants reported wanting access to their records upon leaving the hospital. Some participants requested the ability to email their information to themselves or to their primary care provider. Several patients requested that information not typically part of a medical record be accessible outside the hospital or during their next hospital stay, such as viewing previous care team members. One patient wanted the ability to review ratings and provide reviews of exemplary staff members.

5. Discussion

This study explored hospital patients' usage, experience, and information needs using an inpatient PHR during their post-operative hospital stay. Participants uniformly responded favorably to having access to their clinical data and believed that the inpatient PHR was useful. The results of our study suggest that an inpatient PHR can be helpful to improve understanding, engagement, and safety for some patients in the hospital. These findings are consistent with the results of previous research investigating the effects of sharing medical information with patients through PHRs and patient portals in the outpatient setting [17, 19, 44–49]. Many patients want to play an active role in their care [8, 17, 44, 50], but patients cannot express informed preferences unless they are given sufficient and appropriate information about their medical care.

Our study adds to the limited research that has examined the benefits of providing hospitalized patients with access to their medical information and educational information.[29–36] There is still only a rudimentary understanding of how PHRs can contribute to improving inpatient experience and increasing patient engagement in the hospital setting.[37]

5.1 Implications

5.1.1 Challenges

Fulfilling patients' desires for personalized information about their hospital stay presents numerous challenges. Some of these challenges include determining what information is most beneficial to present to patients and choosing the format and context in which such information should be presented. Although patients wanted access to "everything," presenting the whole medical record (or even the many items requested by the study participants), may overwhelm rather than empower and truly inform. To avoid "information overload," considerable effort will be required to identify the most informative pieces of medical data that are meaningful to patients with various conditions and across various health services.

In order to fully explore the potential of inpatient participation and real-time sharing of medical records with inpatients, additional research is needed to determine how best to present medical record information to diverse patients. This is not a small challenge, as there are many factors that may influence patient interest and engagement with their health information and care [51]. For example, Weed proselytized the importance of format to profoundly help or hinder error discovery in and understanding of the medical record [52]. Key to maximizing understanding and engagement will be designing effective user interfaces [53]. Presenting medical record information to inpatients creates numerous new challenges: accommodating a patient's native language, health literacy level [54–56], information disclosure preferences [57–60], lack of familiarity with new technology, feelings of isolation and compromised constitution.

Providing daily "patient-friendly" progress reports as requested by participants may increase demands on clinicians to create and keep these reports up-to-date [61]. While clinicians typically write daily progress notes of patients for other clinicians, the content of these notes tends to be highly technical, is often in short form and is thick with abbreviations, reducing its informative value to patients. It is possible that existing technologies, such as natural language processing, could be leveraged to help interpret these notes to make them more accessible to patients and their families [62,63].

Ensuring privacy is a significant challenge for an inpatient PHR. To protect electronic health information, security measures must be instituted to prevent access to unauthorized persons. How-

ever, such measures (e.g., passwords) may hinder accessibility and convenience and, thus, use of an inpatient PHR. Passwords are easily forgotten, especially by impaired hospital patients. Additionally, patients have expressed a desire to have granular control over of their medical record.[20] New policies, logistics and technical challenges must be resolved in order to meet such desires without hampering usability, data sharing, or accessibility.

5.1.2 Benefits

From the interviews, we discovered how an inpatient PHR can bring value to hospitalized patients in five ways:

1. *Facilitating data ownership*: Patients view medical data as their property. Regardless of whether it is actionable or understandable, patients expressed it was their right to own and have ready access to their medical record. An inpatient PHR facilitates patients' feelings of ownership of their data.
2. *Reducing uncertainty and anxiety*: Study participants reported that receiving information about their care helped alleviate their uncertainty and stress by providing a sense of control.
3. *Providing understandable information*: Study participants appreciated the application's patient-friendly language, crediting it for helping them understand and take greater interest in their medical conditions. Additional work is needed to address issues of health literacy and numeracy, including providing information in a patient's preferred language[64];
4. *Improving accuracy of data in the health record*: Medical records can be inaccurate, incomplete, and biased.[65] With increased patient demand for access to medical records [8] and increased incentives to provide that access in recent federal legislation [7], more patients may review their medical records and help correct mistakes they discover [66]. Furthermore, this practice may motivate clinicians to improve data collection and documentation. Improving data quality and accuracy could in turn improve reuse of the data for secondary purposes such as quality improvement activities[67];
5. *Improving patient safety*: Providing patients with their medical information is beneficial for safety, as exemplified by a study participant in our study who intercepted a medication dosing error. An inpatient PHR may enable patients and family members help reduce medical errors, which have been identified as among America's leading causes of death [68].

5.2 Suggestions for Future Research

There remains a critical gap in the rigorous evaluation of inpatient PHRs to enhance patient engagement. Encouragingly, more health systems and an electronic medical records vendor have recently spearheaded studies of inpatient PHRs [34–36, 69, 70]. Further research is needed to clarify the appropriate design and implementation of these systems. Though usage in our study of the application was relatively high (86% of patients), it was a very high-touch intervention, with a research assistant spending considerable time educating patients about the technology and addressing concerns. Future research should help clarify what usage might be in a typical hospital ward.

The right balance must be struck between sharing all of a patient's record versus sharing information that is deemed useful, to avoid overwhelming patients with large volumes of data. Consideration must be given to policies of sharing sensitive test results or diagnoses before physicians can have conversations with their patients. Progress must also be made on creating user-interfaces and device ergonomics that take into consideration that "a hospital patient should be treated as a situationally-impaired user" [71], and their physical and mental state is likely to fluctuate over the course of a hospital stay. As part of this process, family members should also be considered as users, particularly for frail and elderly patients, and attention should be given to appropriate security and privacy controls when granting them access [26, 60].

Future research should quantify the benefits and negative consequences of patient-facing technologies at the bedside. Specifically, a „dose-response“ paradigm could be employed to determine whether the level of technology use is associated with changes in patient outcomes such as satisfaction, activation, health system utilization, readmission rates, and mortality. Randomized clinical trials should be conducted on a large and diverse inpatient population spanning a variety of health-care settings (e.g., community hospital, long-term acute care, and academic medical center) to ensure results are widely applicable.

Finally, in pursuing precision medicine's goal to take into account individuals' differences, future studies could characterize patient demographic traits associated with inpatient PHR usage patterns. Variability exists in patients' ability to handle the uncertainties and stress of a hospitalization. Accordingly, we anticipate an intervention like ours will be less compelling to patients who are relatively comfortable with uncertainty and have fewer information needs. Notwithstanding, family members may benefit, even if patients themselves do not wish to use the PHR system.

5.3 Limitations

This study had several limitations. First, it was a small qualitative study conducted in a single setting at a large academic medical center. All of the participants were post-operative cardiac surgical patients in the post-ICU step-down unit. These patients may not be representative of all hospitalized patients. Second, there is the potential for volunteer bias, as our study was a convenience sample of English-speaking patients who were mentally and physically stable. Twenty-four patients were approached, and fourteen patients completed interviews. This could lead us to overestimate the number of patients or family members that would use an inpatient PHR. Third, there is the potential for response bias since the study was not blinded and participants may have chosen to answer questions in a way they thought was favorable to the researchers. A subset of answers, such as to questions about participants' daily Internet use and the time participants spent interacting with the application, could have been subject to recall bias.

The impact of taking part in a formal study on inpatient PHRs may have had an effect on the amount that the tablet and inpatient PHR were used, as mentioned previously. With 14 patients, we did not have sufficient power to detect differences in usage based on sociodemographic variables. In addition, interviewer bias could have resulted in a more positive interpretation of participant responses than was shown in the data. We attempted to mitigate this last issue, by conducting two independent content analyses of the transcribed interviews and comparing results. Finally, inpatient PHRs may vary in content, usability, and function and thus the experience of participants in this study with the application may not generalize to experience with other inpatient PHR applications.

6. Conclusion

This study uncovered a high desire by inpatients to have access to information about their hospital care. Providing hospital patients access to an inpatient PHR may potentially help improve patient satisfaction, understanding of care, and safety. Our results highlight the need for further investigation of the optimal ways to share medical information with inpatients and the efficacy of inpatient PHRs in improving patient experience and health outcomes.

Clinical Relevance Statement

Patients have unmet information needs in the hospital setting. Providing hospitalized patients with personalized health information through a tablet computer may yield considerable benefits--improved data quality and patient safety, increased patient understanding of health conditions and hospital care processes, increased patient awareness of the recovery process, decreased patient stress, and increased patient satisfaction.

Conflicts of Interest

The authors have no competing interests to declare.

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Human Subject Research Approval

This study was approved by the Columbia University Medical Center Institutional Review Board.

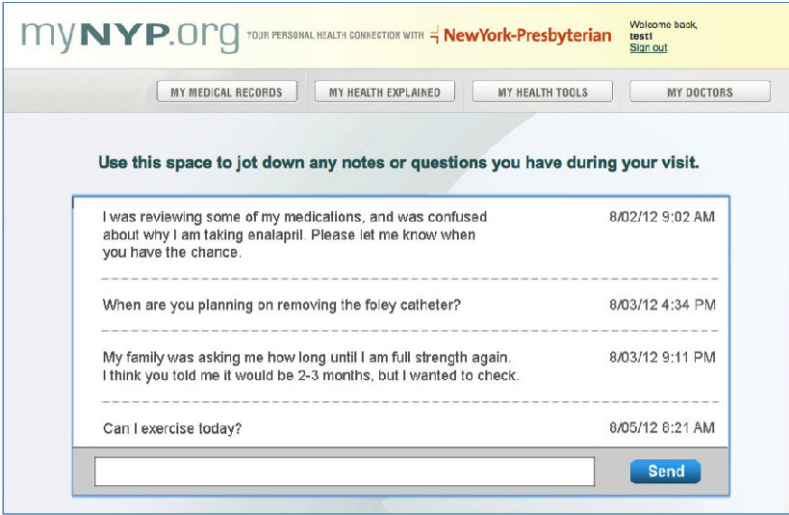


Fig. 1
Patient-entered ques-
tions and comments for
the care team

		Overall (%)
Gender	Female	7 (50%)
Ages	26 – 79	Mean = 59
		SD = 16
Education	High school or less	2 (14.3%)
	Some college	4 (28.6%)
	College graduate	8 (57.1%)
Prior Computer Experience	Yes	13 (92.9%)
	No	1 (7.1%)
Prior Tablet Experience	Yes	3 (21.4%)
	No	11 (78.6%)
Average Daily Internet Use	Never	1 (7.1%)
	1–30 min/day	3 (21.4%)
	31–90 min/day	4 (28.6%)
	>90 min/day	6 (43.9%)

Table 1

Participant Demographics and Technology Experience (n=14)

		Overall (%)
Reported Inpatient PHR Use		
Did not use after observation period		2 (14.3%)
0–15 mins		3 (21.4%)
16–30 mins		7 (50%)
1–2 hrs		2 (14.3%)
Reported Tablet Use for Other Purposes		
No		8 (57.1%)
Yes		6 (43.9%)

Table 2

Participant Use of Inpatient PHR (n=14)

Table 3 Themes Related to Patients' Experience with the Inpatient PHR

Themes	Description	Examples Quotes from Patients and Family Members
Desire to Understand and Engage	An inpatient PHR can help patients and their families learn about and engage with their care.	"I looked up the medicines to see what it does. I would have never known what [the medicine] was." "I can't believe I am taking all these medications." "They started all kinds of new medications, and it was very easy for me to go on the medication [page] and see the side effects. Instead of thinking there is something else wrong with me I was able to say, 'Gee, well maybe it's a side effect from this.'" "I read about the whole procedure that I had, the cardiac bypass surgery, and it's amazing exactly what happened and what they were able to do. And I didn't really realize that because they use all these big terms, and when you break it down to someone who is not medical, it's very interesting." "Access to their medical information gives patients and their caregivers perspective and insight into their hospital care and empowers them with knowledge about [what is going on], which reduces anxiety."

Table 3 Continued

Themes	Description	Examples Quotes from Patients and Family Members
Desire for Status Report	Inpatients want information concerning their health status and recovery progress.	"When am I getting out of here?...What progress am I making?" "Like right now, [I would like my care team to tell me] what's my plan? 'All x-rays are good. We are now waiting for your INRs to reach the level of 3. It is now at a level of 2.5. So hopefully we can increase the heparin dosage and by tomorrow it should reach it, and you should be discharged.'"
Report Test Results	Inpatients want to learn about their diagnostic tests and see their results.	"If [the clinician] is performing a procedure on me, I don't know what the hell it is called." "He's had x-rays and blood draws and I would like to see the results." "I know they were giving him phosphorous, so I would like to see what the levels were."
Ownership of data	Inpatients consider their medical record as something they have a right to own.	"I think if it pertains to them, they should have anything that they want. It's your record." "Everything."
Desire for Guidance	Inpatients want instructions about what to do and expect in and out of the hospital.	"I just had open heart surgery, something saying...the feelings you might experience, how long the feelings may be there for...things you have to do, how long you have to do it. The road to recovery." "You want to go home knowledgeable. If I could have a tablet...where I could see..., Dad had this, this, and this. This is what you should be looking for. That would be ideal. That's what I want."
Inpatient PHR Serves as Information and Memory Aid	Inpatients' questions can go unattended for long periods and can be missed or forgotten. An inpatient PHR could serve as an informative agent covering the time between provider visits.	"It's great cause I can look up all my questions and I won't forget anything." "I always am under the impression that doctors are busy people; they can't just stop and come to see me and explain things to me." "A lot of times doctors come in and tell you 20 things and out of the 20 things you'll only remember like 1 or 2. So you're like, 'What the hell did he say? I forgot what he said.'" "I had problems naming doctors because so many of them come to my room." "[The inpatient PHR is an] awesome idea because if you are sitting in bed and you have a question often times you forget it by the next day."
Improves Safety	Patients and family members can use the application to prevent errors.	"I like it because I discovered yesterday a medication he's getting that he is taking at home, and they're actually giving him half the dose he's getting at home." (A family member recognized a dosing error while using the application. She alerted a nurse and the medication was changed to the appropriate dose.)
Well-being Affects Use	Patients' lack of well-being affects use and interest in the application.	"When you first come in you might be too damn sick to really care about it or read the formalities. You have to be feeling better." "I was very nauseous and not feeling well so I really wasn't up to using it. Started this morning, trying to look at it a little bit because I was feeling better... if you're not feeling well you're not going to really be using it because you're not up to it... you don't have the energy."

Table 3 Continued

Themes	Description	Examples Quotes from Patients and Family Members
Messaging Providers: Mixed Response	Inpatient-provider communication via electronic messaging may be beneficial for some and ignored by others.	"Me personally, I don't need to see you. I just want to text you." "I think it's a good thing—if they answer, some of them aren't online to answer. I think we should be able to email our doctors all the time. I don't care where they at. They always got computers, iPhones, whatever." "[Doctors] are not going to be available sitting down at the desk the whole entire day. They're going back and forth, so...they're just not going to have time to [answer questions]." "It would be nice to have the communication face-to-face, but for some questions like about meds [it] might be quicker to text the doctor." "I don't think it's a good idea. The doctor is supposed to see the patient, sometimes feel them." "I just think it'd be nice if I [sent a message] and then they said, 'I'll be with you in ten minutes,' or something."
Self-Reporting Pain: Mostly unfavorable	Although some participants thought entering pain scores in the application could replace or supplement clinician pain assessments, most participants did not think self-reporting pain scores would be advantageous.	"If I say I need pain medication and I enter it and tell you – and enter it as a 6 or a 7 – then I guess the nurse can determine how quick she has to get to me. So maybe it's a good thing." "I don't know the benefit that would do. Because if you're in pain all you do is ask the nurse, I'm in pain and they'll ask you 1 to 5, it's a 4, and they'll assign you some medication. It's very simple." "No. You know why? Because I want to see [that] the nurse knows."

Appendix Interview Questions

User Background

1. What is your age? Record gender as well.
2. What is your profession? Level of education?
3. Do you know your medical conditions? Do you know your medications and what they do? How knowledgeable would you say you are about your health?
4. Do you have a smartphone? How much time do you spend each day using the internet or apps on your smartphone on average?
5. Do you have a desktop or laptop computer? Do you have internet on your computer? How much time do you spend each day on your computer on average?
6. Do you have a tablet computer? How much time do you spend each day on your tablet on average?
7. Do you have a personal health record? Who do you use it for (self, children, parents)? How frequently do you use it?
8. Do you look up information about your health online? What websites do you use?

General Inpatient PHR Questions

1. What do you think about patients receiving an inpatient personal health record like the one you used?
2. What information do you think patients would like to see while they are in the hospital?
3. What information should patients not have access to?
4. What questions have you asked your doctors while in the hospital? What do you think about patients being able to ask questions or make comments to their care team in the application? Is this useful?
5. Should this be one-way or two-way communication? (ie should the patient send messages in which a member of the care team responds in person, or should patients and the care team communicate electronically through the application).
6. What do you think about patients entering their own pain scores? Could this replace the scores recorded by nursing?
7. Patients will still have access to their records when they leave, but how long should patients have access to the special features of the inpatient PHR, such as staff profiles or ability message with your team, after the hospital stay?

Questions about application (myNYP)

1. What do you think about the myNYP Inpatient application?
2. What was most useful about the application? What was least useful?
3. What do you think if anything is missing from the application? What do you wish we had included?
4. Was it easy to find what you needed? Did you find navigating through the application confusing?
5. Was the information accurate?
6. What did you think about x (care team, hospital medications, allergies, home medications, note-pad, pain)? What would you add or change to them? *(If they did not use it, show it to them and ask for their opinion)*
7. Was the text the right size? How was it entering notes on the iPad?
8. How much time would you say you spent looking at the application? What else did you use the iPad while you had it? (We want to get a sense of what patients might use iPad for apart from our application)

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