Designing Visual Communication of Everyday Illness Experiences in Complex Pediatric Care

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ABSTRACT
In complex chronic care, patients’ ongoing awareness of their health status and ability to articulate health needs are vital to active participation in care, yet they face various challenges that could thwart their potential to engage in such participation. My research explores how design methods in HCI can evolve to meet these challenges by engaging both adolescents and family caregivers throughout the process of tracking the patients’ illness experiences and co-designing rich representations that are expected to support adolescents’ communication of these experiences in care. This thesis will contribute 1) a critical understanding of the ways in which human-centered design can address primary challenges that adolescents face when engaging in care, 2) a novel method for conducting co-design research with chronically ill patient families, and a 3) family-centered mobile health technology that demonstrates the feasibility of engaging pediatric patient families.

KEYWORDS
mobile health; chronic illness; patient-generated health data; co-design; adolescent, family.
1 **Complex Chronic Conditions**: Complex chronic conditions (CCC), in its working definition in clinical studies, include a range of diseases that "can be reasonably expected to last at least 12 months and to involve either several different organ systems or one system severely enough to require specialty pediatric care and probably some period of hospitalization in a tertiary care center" [3]. By this definition, CCCs are different from more prevalent forms of pediatric chronic illnesses such as obesity and diabetes. Pediatric rheumatism, renal failure, cancer, and blood disorders, which are the focus of my work, fall under this definition based on Feudtner et al.’s characterization.

2 **Patient-Generated Health Data**: Patient-Generated Health Data (PGHD) are health-related data that are created or inferred by patients or care partners to help address a health concern. Two types of PGHD inform health care: clinician-, and patient-defined PGHD [1]. Clinician-defined PGHD are assessments that professionals find important, but that must be gathered by the person experiencing daily life to contribute to those assessments. On the other hand, patient-defined PGHD reflect concepts that are uniquely defined and seen as important by the patient.

3 **Observations of Daily Living**: Observations of Daily Living (ODLs) is a conceptual framework and a specific type of patient-defined PGHD that describes PGHD reflecting the patients’ perspective [1]. ODLs reflect feelings, thoughts, behaviors, exposures and actions that serve as cues to healthy action and indicators of personal health states.

Sidebar 1: Key Concepts and Background

**CONTEXT AND MOTIVATION**

Design research in HCI has played a fundamental role in advancing the design of health technologies for chronically ill patients, yet research involving pediatric patients with special health care needs has been limited to date. In care of these patients, decisions about treatment and supportive care, are made based on a comprehensive understanding of the patient’s health data, including physiological data, physician’s global assessment, and caregiver- and patient-reported observations of the patient’s health status [2]. It is important that patients articulate their physical, psychological, social, and emotional well-being to the care team, but adolescent (ages 10-19) patients with CCC’s, in particular, face difficulty expressing these illness needs as their transition into adulthood marks an important yet challenging period of biopsychosocial development. While they have the firsthand experience, there is little consideration for the patients’ own assessments of their health in clinical decision making due to their dependence on family and clinical caregivers, conflicting interpretations about their health status, poor recall, and lack of means to understand and express their health needs.

To communicate these illness needs, patients will first need to build on meaningful representations of their everyday illness experience to support ongoing awareness of their health status [7]. Technology has the vast potential to help adolescents construct meaningful representations of their lived experience, yet no tools exist to support their engagement and communication of illness needs that build upon these representations. My answer to this problem and broader research goal is to re-envision design methods to create collaborative health technologies that scaffold patients’ gradual participation in care as designers—by means of engaging family members as co-design partners.

For my dissertation research, I am working with oncology, hematology, nephrology and rheumatology clinical collaborators at Children’s Healthcare of Atlanta (CHOA) to understand how design methods can evolve to better engage adolescents and family caregivers throughout the process of tracking and co-designing rich representations of the patients’ daily illness experiences that have personal relevance to their management of treatment effects.

**COMPLETED AND ONGOING WORK**

For the past four years, I have been leading various research studies involving chronically ill adolescent patients in both clinical and non-clinical settings that span participant observations, interviews, portal usage log analysis, surveys, co-design, and a probe study. Thus far, I have completed two projects and started collecting data for the final project. Three research questions (Sidebar 2) guide my work.

1. **Understanding Patient Participation in Complex Care**: My early formative work provides an overview of the problem and design space for adolescents’ participation in complex care, which address RQ 1. Through individual interviews (in private) with patients and their parents, I learned that adolescents’ faced challenges participating in clinical conversations, communicating emotionally
RQ 1: How do adolescents with CCC’s and family caregivers participate in care? [5, 6]
- Empirical analysis of patient-parent perspectives toward their participation in complex care prior to, and after being exposed to patient portal technology.
- Characterization of challenges and design space to support adolescents’ participation in complex care.

RQ 2: How can we design tools to elicit information about adolescents’ everyday illness experiences? [4, 8]
- Creation and expansion of co-design artifacts (Visual ODLs) and methodological innovation (scaffolding technique).
- Formative usability evaluation of UIs integrating Visual ODLs for clinical use.

RQ 3: How can we engage adolescent patients undergoing complex treatment in the design process, to create illness representations that are meaningful to these patients and their family members?
- Empirical analyses of interviews and diary entries, participant-generated artifacts from co-design, and quantitative EMA and survey measures.
- Characterization of technology affordances and family-tracking practices in capturing ODL data.
- Design and deployment of novel family-oriented mHealth application (CO-OP).

Sidebar 2: Research Questions and Summary of Contributions (bullets).

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To address RQ 3, I apply the ECD method to adolescent patients with CCC’s, as a case study, to show that the method is useful, valid, and reliable for its intended purpose [9]. To do this, I employ mobile ecological momentary assessment (EMA) methods that are geared towards achieving high sensitive information, and managing physical and emotional responses. In particular, while patients downplayed their symptoms not to worry their parents, symptom reporting was often assumed by parental caregivers, even if the reported data may not adequately represent the patient’s true felt experience [6]. In another study, I conducted log analysis, survey and interviews to understand adolescent and parent perspectives after they have navigated the patient’s health information through a tethered personal health records system, yet found that current patient portal technologies had limited means to engage these patients in care [5]. These studies created contours of a design space for accommodating patient and parent perspectives in symptom reporting by supporting their gradually evolving partnerships through the design of a patient-friendly and engaging technology.

2. Characterizing Patient-Defined Patient-Generated Data: Informed by research on the positive impact of images on symptom communication [7] and the concept of Observations of Daily Living (ODLs), my recent work, addressing RQ 2, explored the use of co-design methods to characterize patient-defined PGHD for communicating adolescents’ lived experience [4]. In this work I adapted the ODL framework and solicited domain knowledge from clinicians to generate design artifacts, or Visual ODLs (Fig. 1), to elicit everyday illness experiences through co-design sessions with adolescent families. Through co-design, I found the need for technology to scaffold the process of encoding and articulating symptomatic experiences through representation of how symptoms affect patients’ ability to engage in daily activities, and to support distinct roles which family caregivers can serve in tracking the patient experience (e.g., timing and frequency). In another study, I examined the potential value of Visual ODLs in clinical conversations that uncovered important use cases demonstrating clinicians’ preference to promote patient-initiated first-person illness narratives during face-to-face encounters [8]. These insights informed the design of the current CO-OP system (Fig. 2) and research goal to understand how the family could work together as co-designers while tracking the patients’ health.

3. Supporting Situated Design Work in Everyday Life: I am currently drawing on design methods to create an interactive mHealth system as an endpoint of the novel design process that investigates how technology can support patients and family caregivers’ collaborative effort to track and co-create personally meaningful representations of everyday illness experiences. Over a 2–3 week period, the system will elicit and probe patients’ and family caregivers’ observations of symptomatic experiences in relation to their everyday activities, and their design input, through a suit of media technology readily available on their mobile device. One goal of this work is to create and evaluate a new method, ‘Episode-Based Co-Design (ECD)’, which is motivated by the lack of methods for engaging patients and family members in the co-design process when the design needs to be close in time to phenomena of interest (e.g., in this case, an episode of the illness experience).

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ecological validity by placing the data collection activities in the hands of patients and their parental caregivers, in their natural setting. To better understand patient and parents’ conceptions of the illness experience, I am conducting co-design activities that are informed by a rich set of data collaboratively collected on a daily basis by the patient family. To measure the relevance of illness representations, co-designed artifacts will be assessed by a subgroup of patient populations belonging to similar disease categories. Through this research, I aim to analyze many factors with respect to patients’ engagement in care, including compliance, quantity and quality of design outcomes, distribution of tracking tasks among family members, and the unique privacy needs arising from family-based tracking.

My work is expected to inform the design of tools that support patient and family partners’ engagement with their everyday lived experiences by scaffolding their co-creation of personally meaningful representations. These co-designed representations could support other vulnerable populations who rely on a network of caregivers to better articulate their experiences.

REFERENCES

APPENDIX 1. EXPECTED BENEFITS OF PARTICIPATION

I am a PhD Candidate in the Human-Centered Computing program at Georgia Institute of Technology. I work in the Health Experience and Applications Lab with my advisor Dr. Lauren Wilcox. I proposed my dissertation research in May 2018 and plan to defend next year, so the CHI 2019 doctoral consortium comes in a timely manner (to receive feedback) as I finish my dissertation studies. I am particularly interested in discussing my plans for analyzing granular data generated from family-based mobile deployment studies that involve design at the end point of the design process. Additionally, I hope to receive insights regarding how I can better situate my empirical and methodological contributions within Human-Computer Interaction (HCI), from the perspective of the CHI community. With the rapid expansion of health tracking tools, health has quickly evolved as a theme at CHI. To this effect, I believe this doctoral consortium is the perfect setting for me to present my dissertation work and engage with scholars from various academic disciplines who share similar visions of human-centered computing for sociotechnical systems in healthcare. Having been trained in Health Informatics (HI) and HCI, I have done substantial work to frame my research by synthesizing work from both fields. I expect that doctoral consortium participants will gain a critical understanding of ways to merge or adapt concepts from disparate fields to contribute to HCI research. For example, I dissect scholarship around Patient-Generated Health Data to reappraise and adapt the concept of Observations of Daily Living (ODLs) in my research, an important framework suited for understanding PGHD reflecting patient perspectives. I also merge in-situ design methods with participatory or co-design approaches and propose methodological innovations to better engage vulnerable populations in the human-centered design process. Overall, I highly anticipate my participation in the CHI 2019 doctoral consortium to be a valuable and intellectually stimulating experience that will push me to think about the implications of my work.