Patients Are Knowledge Workers in the Clinical Information Space

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Abstract

Background Limited research exists on patient knowledge/cognition or “getting inside patients’ heads.” Because patients possess unique and privileged knowledge, clinicians need this information to make patient-centered and coordinated treatment planning decisions. To achieve patient-centered care, we characterize patient knowledge and contributions to the clinical information space.

Methods and Objectives In a theoretical overview, we explore the relevance of patient knowledge to care provision, apply historical perspectives of knowledge acquisition to patient knowledge, propose a representation of patient knowledge types across the continuum of care, and include illustrative vignettes about Mr. Jones. We highlight how the field of human factors (a core competency of health informatics) provides a perspective and methods for eliciting and characterizing patient knowledge.

Conclusion Patients play a vital role in the clinical information space by possessing and sharing unique knowledge relevant to the clinical picture. Without a patient’s contributions, the clinical picture of the patient is incomplete. A human factors perspective informs patient-centered care and health information technology solutions to support clinical information sharing.

Keywords ► patient knowledge  ► human factors  ► health informatics  ► patient-centered care  ► knowledge  ► cognition  ► patient care

Background and Significance

There has recently been more research on characterizing patient behavior within and outside of clinical settings, mainly in the context of chronic or long-term illness, to inform the development of human-centered technologies. Patient ergonomics, a subdiscipline of human factors, studies the health-related work of patients, caregivers, and community members.1 Limited research has been concerned with “getting inside patients’ heads” or patient cognition. Cognition refers to the processes (associated with knowledge, thinking, memory, attention) underlying outward behaviors. Patients possess unique and privileged knowledge particularly about historical and daily experiences with their illness. Although clinicians need this information from patients to provide patient-centered care, patient knowledge may not be systematically elicited or documented by clinicians. In this article, we propose a theoretical characterization of patient knowledge and explore its relevance to care provision by applying traditional cognitive science and human factors (a core competency of health informatics2) perspectives to define patients as knowledge workers across the continuum of care. To demonstrate that patients possess and share knowledge relevant to the clinical picture, we present vignettes about Mr. Jones.

• Background—A patient has experience and comprehension of illness, medication side effects, and self-care management strategies: Mr. Jones, a 50-year-old male with a
master’s degree, was diagnosed with multiple myeloma 4 years ago. Mr. Jones also suffers from hypertension, for which he takes blood pressure medication and exercises regularly. Mr. Jones’ cancer has relapsed multiple times but it has been controlled with chemotherapy and stem cell replacement. He regularly sees his oncologist and his primary care provider. Since his diagnosis, Mr. Jones has had several emergency room visits and hospitalizations at both his local hospital and the academic medical center where he gets his cancer care. With the help of his wife, Mr. Jones manages the logistics of his doctors’ visits and chemotherapy infusions, as well as medications and self-care activities. He keeps his test results in a binder and adjusts lifestyle and activity levels according to his oncologist’s guidance and personal strategies. Recently, Mr. Jones has begun to suffer from debilitating neuropathy caused by one of his cancer medications. He also had a stairlift installed in his home to prevent falls due to fatigue and lightheadedness. Mr. Jones has an informed understanding of his condition and the mechanisms of action of his medications. He has also developed strategies for managing his life, given medication side effects.

- **Patient and caregiver communication of patient status may impact care safety and quality:** While neutropenic due to chemotherapy, Mr. Jones had an acute event that lands him in the local hospital’s emergency room, rather than the academic medical center where he receives his cancer care. The electronic health record (EHR) systems are not integrated. The emergency room clinicians do not follow guidelines for neutropenic precautions and prepare to conduct a series of unnecessary tests. Mr. Jones highlights his immunocompromised status to the clinicians and asks his wife to provide blood test results from the previous day. Clinicians put in place neutropenic precautions.

- **Strategically withholding patient knowledge may impact decision making regarding treatment options:** Mr. Jones is eligible for a clinical trial for a promising new medication. Mr. Jones sees this clinical trial as his last hope and strategically withholds from his care team the poor quality of life that he is experiencing at home for fear of not being included.

- **Experiences of medication side effects may impact medication options:** Mr. Jones is interested in exploring a new hypertension medication. When asked by his primary care physician about bothersome side effects of his cancer medications, Mr. Jones forgets to mention neuropathy. Mr. Jones is thus prescribed a medication that exacerbates his neuropathy, thus impacting his quality of life as well as his functionality.

Despite the impetus toward patient–clinician teaming in health care and policy for systematic sharing of patient-generated health data, research to characterize patient cognition as underlying their contributions to the clinical information space is limited. This is a challenge for both technology development and synchronous and asynchronous communication. The content of patient knowledge informs thinking and underlies behaviors of managing wellness, health, and interfacing with the health care system. Qualitative researchers like nurse and sociologist, Corbin and Strauss have a history of capturing patient experiences with chronic illness and characterizing activities of self-care management as “work.” However, rather than answering research and applied questions of how to support the work, their research objective was the discovery of patient experiences of work. Designing human-centered health information technologies (HITs) to support capturing, sharing, and integrating of patient-held knowledge requires the understanding of not only what patients do, but when, why, and how they do it.

Human factors approaches have been used to characterize decision making of professional experts in real-world domains of aviation, nuclear power, military, and health care. Findings have informed models of knowledge acquisition and decision making. Only recently have patients become recognized as active participants and decision makers in and outside of the health care system. Despite the relevance of cognitive science and human factors literature to this applied problem space, most theory and research on patient cognition have stopped short of applying it.

### Why Is Patient Knowledge Relevant?

Patient knowledge is relevant because of its relationship with clinical outcomes. For instance, research by Khan et al in pediatric hospital settings highlights that families play a role in patient safety through vigilance and reporting of potential errors. Cognitive processes associated with decision making underlie vigilance and reporting. Work by Weiner and Schwartz spanning the last decade stresses the importance of avoiding contextual mistakes in medical decision making by physicians asking the “right” questions of patients. By gathering contextual information from patients, particularly about life and social constraints, physicians can better inform patient-centered treatment plans, highlighting the value of patient contributions. This work is an example of the importance of clinician-driven knowledge elicitation from patients.

Further, lower health literacy is associated with lower rates of preventative care, medication management challenges, more hospitalizations, more emergency care use, poorer health, and higher mortality rates. On the other hand, patients and caregivers with higher health literacy may actively influence their care plans through engagement and shared decision making. In a BMJ Comment, Kennedy highlights the idea that patients are experts in their “experience, feelings, fears, hopes, and desires.” Thus, even in the absence of illness-specific knowledge, patients may be experts in their own right. The term “expert patient” refers to patients who are not just health care consumers but producers of health through deep knowledge and comprehension.

### Cognitive Science Perspectives on Knowledge Acquisition

Traditionally, cognitive scientists have examined the acquisition of knowledge in service of education. In 1980, Dreyfus and Dreyfus proposed a stage model of knowledge...
acquisition where a learner progresses from a rule-bound novice to an intuitive reasoner making decisions based on tacit knowledge. Generalized to nursing practice, the Dreyfus model continues to be heavily cited in medicine today. In addressing designing medical education to support knowledge acquisition, Ericsson’s deliberate practice theory is widely accepted. This theory’s idea is that expert performance is a function of intentional practice designed by educators. This practice incorporates opportunities for problem solving, immediate feedback, evaluation, and repeat performance. Ericsson’s theory has been generalized to other professional domains such as transportation, entrepreneurial skills, writing, teaching, and even music, chess, drawing, math, and software design.

In complement, human factors scientists have been seeking to understand knowledge through theoretical perspectives such as naturalistic decision making (NDM) that accounts for expert performance in complex real-world domains. NDM provides both a theoretical perspective and a set of research tools to examine how professionals (primarily subject matter experts or SMEs) solve problems, make decisions, and perform cognitively and perceptually complex work. This work has traditionally been conducted in domains marked by uncertainty, time limitations, and high consequences for erroneous actions. Because it is not feasible to bring real-world complexity into the laboratory, methods include interviews to elicit challenging lived incidents from SMEs and identify and unpack decision points. Again, much of this research has been focused on professional domains to capture expertise and deliver it to novices as part of training, education, and decision support solutions. In health care, substantial literature exists using the NDM perspective to characterize clinicians’ knowledge and decision making. However, again, the topic of patient cognition has received little attention even in human factors.

Of particular relevance to informatics is the understanding that many types of patient knowledge, especially in the cases of chronic and long-term illness, are not acquired in a deliberate manner through formal education and training, but are emergent as a function of experience and continuity across time and space of the care continuum. Some types of patient knowledge may be acquired through implicit learning or learning that occurs without conscious effort, and may potentially yield deep understanding. Although authors suggest that self-management of chronic illness like diabetes “draws on the same cognitive skills found in experts from diverse professional domains,” few studies have focused on examining the acquisition and content of such knowledge. For instance, Lippa et al suggest that patient’s problem detection skills and strategies acquired through practice in diabetes self-management are associated with better adherence and greater glycemic control, which may indicate proficient performance. Recently, Holden et al used the NDM approach of incident-based interviews to characterize decision making personas to inform the design of solutions to support patients with heart failure. Their study highlights the need to understand cognitive processes, which are associated with decision making to inform human-centered design.

Traditionally, there are two types of knowledge, explicit and tacit. Explicit (e.g., declarative, expressive) knowledge can be easily articulated, codified, and transferred to others. For example, explicit knowledge may be acquired through book learning. In contrast, tacit (e.g., implicit), proposed by Polanyi, is acquired through practical, lived experience, and often cannot be readily articulated. Knowledge associated with carrying out skills or tasks (e.g., riding a bike, tying shoelaces) is termed procedural knowledge and is considered implicit. Although there are some slight variations in the literature regarding terminology and distinctions, we emphasize that a substantial literature exists in cognitive science and propose the need to extend and adapt accordingly to examine and characterize patient knowledge. Further, patients possess and manage knowledge that falls under the theoretically accepted categories of knowledge of explicit, tacit, and procedural and additional categories of privileged knowledge or “knowledge in the head,” such as preferences and history. Privileged patient knowledge that is unelicited due to barriers (e.g., time constraints, individual differences, bias) will remain so.

**Patient Is a Knowledge Worker**

Holden et al positioned the patient, alongside the clinicians, at the center of the health care complex sociotechnical system in SEIPS 2.0. The authors highlight that the patient conducts “work” ranging from logistical planning and execution to medication to self-care management, even in cases where there is no disease present. The SEIPS 2.0 framework highlights an active patient role across patient-only activities as well as patient–clinician teams. A body of literature now exists examining patient work focused on understanding daily contexts and activities, culminating in the idea that investigating health care workflow is incomplete without capturing patients’ health-related activities in clinical and daily-living settings. As a function of these activities, patients possess, manage, apply, create, and share knowledge. A recent review of qualitative literature on heart failure proposes a characterization of patient knowledge based on content, development, application, communication, and experience, highlighting that patient knowledge is implicit, explicit, and dynamic. Through such work, patients are knowledge workers. A knowledge worker is a term that has been traditionally applied to professionals who conduct thinking and reasoning as part of their work.

**What Knowledge Do Patients Possess?**

Bodenheimer et al summarized one role of patient knowledge during the self-management of chronic diseases in primary care. They highlight the importance of patients knowing how to identify their problems from their perspective, take actions to address these problems, and adapt as circumstances change. However, only a few studies investigate how a patient acquires such knowledge in the context of a complex illness marked by a particular trajectory, and how that unfolds across time and settings. Recently introduced SEIPS 3.0 highlights the concept of the patient journey across space and time, with patient knowledge playing a role in processes and outcomes.
Given that no taxonomy of patient knowledge exists, we propose first to apply theoretically accepted categories of knowledge of explicit, tacit, and procedural to patient knowledge, as shown in ►Table 1. ►Table 1 contains cognitive science knowledge types and their descriptions, along with examples. Two observations emerge: (1) multiple knowledge examples belong in the tacit category, suggesting that patients may have trouble articulating them, and (2) there are additional knowledge examples that potentially do not fit in this framework.

Thus, we propose additional knowledge examples. We refer to these examples as patient factors (both static and dynamic), an all-encompassing term that refers to patients’ privileged knowledge. The examples of patient factors represented in ►Fig. 1 may be privileged to the patient—some (e.g., attributes) may be more systematically captured and potentially documented in the EHR. Still, most do not even have designated fields in the EHR. These patient factors both inform and are a function of the patient journey across the care continuum and may also evolve (e.g., preferences) as a function of time and experience. ►Fig. 1 represents the problem space of patient knowledge that is in particular need of research attention.

### Acquisition of Knowledge

A crucial mechanism that accounts for privileged patient knowledge is experience and continuity. Patients (and caregivers) are the only ones with continuity across the care continuum—time, space (home, clinical settings, contexts of daily living), and multiple clinicians. ►Fig. 2 represents the patient care continuum as a spaghetti junction—a term used to describe a complex traffic interchange. Not unlike the spaghetti junction, the patient journey (along with caregivers and primarily in the home) across the care continuum is characterized by a winding path with a variety of barriers (construction), highs and lows in physical and mental health (under- and overpasses), the potential for guidance from lay and clinical caregivers such as nurse navigators (a Global Positioning System), and interactions with the health care system requiring planning, logistics, and information exchange (exits off the highway). We note that health care interactions comprise a fraction of time and space within the full patient journey. We currently have no measures to account for patient continuity, and indeed, it is not explicitly represented in the EHR.

Patients may also acquire knowledge across the patient journey via formal or deliberate mechanisms. Through patient...
education, the health care system focuses on providing patients with tools to support empowerment (understanding the health care system and engaging in behaviors that influence situations and outcomes)\textsuperscript{48} and engagement.\textsuperscript{49} Numerous examples exist, including: American Cancer Society\textsuperscript{50} provides patient educational resources specific to each cancer type, as does American Heart Association,\textsuperscript{51} on hypertension. Individual organizations, offices, clinics, and clinicians may provide additional patient education. A growing body of literature exists on the acceptance of consumer HITs.\textsuperscript{52} Human-centered consumer HIT can provide opportunities for managing, storing, and sharing health information (e.g., heart rate, blood glucose monitoring, etc.) and learning about self-care management. Yet, nondeliberately or implicitly acquired knowledge continues to remain poorly understood. Brown and Duguid’s organizational knowledge acquisition\textsuperscript{53} theory, focusing on informal and fluid knowledge acquisition, calls for a focus on practice or application. It also highlights the value of communities in learning. Online patient communities that serve not just as spaces for support, but spaces for information exchange (assuming moderators filter unreliable information), come to mind.

**Implications: Applying Patient Knowledge to the Clinical Information Space and Research**

Patients are responsible for navigating an increasingly complex health care system with information distributed across numerous clinicians and staff, technologies, paper artifacts, and physical settings. Further, through privileged knowledge (phenomenology of symptoms, case history), patients may be the drivers of their care, as suggested by multiple sclerosis research.\textsuperscript{18} Some examples of ways in which patients drive their care include making decisions within and outside of their relationships with clinicians. Traditionally, when we talk about patient decision making, it is often focused on selecting treatment options as a function of patient–clinician interactions and the disease context. However, patients make other decisions that receive less research attention, which may or may not play a role in their outcomes. The following are just some examples of decisions that patients routinely engage in independently (or in concert with their nonclinical caregivers): self-detecting difference in health status,\textsuperscript{54} seeking medical care (when, how, from whom, etc.),\textsuperscript{18} managing medications,\textsuperscript{55} releasing privileged information strategically,\textsuperscript{18} self-medicating with over-the-counter medicines and self-treating,\textsuperscript{44,56} and requesting and receiving preventative care (seeking, etc.).\textsuperscript{57,58} No matter the level of knowledge, patients are decision makers.

In this article, we highlighted the clinical value of patient knowledge. We recognize the unprecedented complexity of studying and characterizing patient cognition and behavior, given they are “situated” within complex sociotechnical systems. Understanding patient cognition or the whys behind patient behaviors can support (1) the design of human-centered technologies for patient–clinician synchronous and asynchronous information sharing, (2) patient–clinician teaming in decision making, (3) informed and engaged patient navigation of the care continuum, and (4) patient assessment...
and adoption of information, resources, and tools for self-management of health and wellness.

To make the above a reality, patients need to take part in the research process. There is no question that researchers have embraced the necessity of including patients to generate and evaluate patient-facing solutions. Some examples include patient evaluation of an application to support self-care during gastrointestinal cancer,\(^{59}\) language for medication instructions,\(^{60}\) and information web sites.\(^{61}\) A study to train patients to create a continuity of care document to share with clinicians across systems reduced duplication of laboratory tests.\(^{62}\) Further, Riggare\(^{63}\) writes about the need for and the value of persons with lived experiences with illness engaging in research. Twitter stakeholder communities, including patients and clinicians, such as #BTSM (brain tumor social media)\(^{64}\) and #BCSM (breast cancer social media),\(^{65}\) (among numerous other benefits) inform research ranging from bringing together ad hoc collaborations and developing research and operational questions, to disseminating research findings, not to mention serving as research mechanisms. Last but not least, using the term “patients marshaled epistemic authority,” a recent publication highlights how it is patients, through social media, that shed research attention on “long-haul Covid.”\(^{66}\) In addition, a 2020 panel at the Human Factors in Health Care Symposium highlighted the unique contributions of researchers bringing their patient perspectives to research.\(^{57}\) Ultimately, engaging patients as partners across the research, development, and implementation process is key to designing solutions that account for patient knowledge.

**Takeaways**

Patient knowledge is integral to inform patient–clinician teaming. Currently, a research gap exists on both sides of the information exchange equation between patients and clinicians. We presented a theoretical characterization of patient knowledge’s problem space as critical to informing patient-centered care. Future research needs to address the application and integration of patient knowledge as part of patient–clinician teaming and decision making.

In the absence of theory applied to informatics solutions, we may mistakenly assume that external behavior reflects an accurate understanding. However, there lies a potential disconnect. Studying cognition can help identify gaps and inaccuracies in understanding, thereby informing appropriately tailored informatics solutions. The study by Lippa and Klein on patient cognition of diabetes self-care is an example of such research.\(^{68}\) There is much work to be done—examining specific illnesses, settings, and patient life contexts. We leave the reader with the following takeaways:

- **Patients are knowledge workers.** Patients possess unique knowledge that clinicians need. By defining the patient as a knowledge worker, we can begin to develop a much-needed taxonomy to inform both research and practice.
- **Patient knowledge impacts clinical outcomes.** Patient-held knowledge is clinically relevant, stressing the importance of developing solutions to facilitate patient knowledge sharing and integration.

  - **The field of human factors offers a perspective and approaches for characterizing and eliciting the content of patient knowledge.** Based on decades of examining cognition and behavior, the field of human factors has approaches such as cognitive task analysis (CTA) applicable to characterizing patient cognition, as demonstrated by studies by Lippa et al.\(^{18,69}\) and Holden et al.\(^{34}\) Further, literature has highlighted the complementary nature of CTA and participatory design.\(^{42}\)
  - **Engaging patients as partners in research is needed.** The active participation of the general public (e.g., patients, citizens, etc.) as partners in the research process (a shift toward citizen science) is key to developing solutions that effectively account for patient knowledge.\(^{70}\)

**Clinical Relevance Statement**

The clinical picture is incomplete without privileged and unique knowledge from patients about their health, particularly for complex or chronic illness. Understanding patient knowledge is necessary to inform patient- and provider-centered HIT solutions to support information sharing and integration.

**Multiple Choice Questions**

1. Patient knowledge is clinically relevant because:
   - a. Patients may possess unique and privileged information.
   - b. Patients may have historical information regarding their illness.
   - c. Patients may be the only constant across the continuum of care.
   - d. All of the above.

   **Correct Answer:** The correct answer is option d.

2. Cognitive science can offer theoretical perspectives to studying patient knowledge because:
   - a. Cognitive scientists primarily conduct laboratory research.
   - b. Cognitive scientists have long studied knowledge acquisition.
   - c. It focuses on human behavior.
   - d. It has long focused on patient cognition in real-world settings.

   **Correct Answer:** The correct answer is option b.

**Protection of Human and Animal Subjects**

No human subjects were involved in this study.

**Funding**

This study was not funded.

**Conflict of Interest**

None declared.
Acknowledgments

We are grateful to Alexandria Cook, BFA, MS (Biomedical Visualization, Department of Biomedical and Health Information Sciences, University of Illinois at Chicago) for coming up with the term "spaghetti junction" as an analogy for the care continuum and for developing Fig. 2.

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Patients are Knowledge Workers in the Clinical Information Space


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