Objective: Online social networking sites are web services in which users create public or semipublic profiles and connect to build online communities, finding likeminded people through self-labeled personal attributes including ethnicity, leisure interests, political beliefs, and, increasingly, health status. Thirty-nine percent of patients in the United States identified themselves as users of social networks in a recent survey. “Tags,” user-generated descriptors functioning as labels for user-generated content, are increasingly important to social networking, and the language used by patients is thus becoming important for knowledge representation in these systems. However, patient language poses considerable challenges for health communication and networking. How have information systems traditionally incorporated these languages in their controlled vocabularies and thesauri? How do system builders know what consumers and patients say?

Methods: This comprehensive review of the literature of health care (PubMed MEDLINE, CINAHL), library science, and information science (Library and Information Science and Technology Abstracts, Library and Information Science Abstracts, and Library Literature) examines the research domains in which consumer and patient language has been explored.

Results: Consumer contributions to controlled vocabulary appear to be seriously under-researched inside and outside of health care.

Conclusion: The author reflects on the implications of these findings for online social networks devoted to patients and the patient experience.

INTRODUCTION

Online social networks, or social networking sites (SNS), have been a feature of the web since 1997, with the founding of SixDegrees.com. SNS are websites that allow users to “(1) construct a public or semi-public profile within a bounded system, (2) articulate a list of other users with whom they share a connection, and (3) view and traverse their list of connections and those made by others within the system” [1]. More and more SNS target people who define themselves by communities according to geographic location, sexual orientation, belief systems, ethnicity, education, and countless other social attributes [2]. Online information, advocacy, and support organizations oriented to specific medical diagnoses were among the first communities of Internet users [3], and SNS for patients are now also a part of the web landscape—making “community” the “killer app in health care” [4]. In fact, the Pew Internet and American Life Project identifies 39% of US “e-patients” as users of social networks, particularly users aged 18–29 [5], which implies a long potential lifespan for this trend. For example, PatientsLikeMe [6] hosts patient communities in 16 varied diagnostic categories, including approximately 5% of all amyotrophic lateral sclerosis (ALS) and primary lateral sclerosis patients in the United States [7, 8]. This SNS incorporates not only a bulletin board, but also clinical tools. Community members report symptoms to find other “patients like them”; “tagging” of symptoms becomes useful data “emergent from shared information” [9].
bidirectional: Not only do many consumers have difficulty with medical terminology, but information systems constructed to understand medical terminologies have difficulty with consumer language. For this reason, consumer health vocabulary research and development is identified in this white paper as a strategy for improved, consumer-centered health communication [10].

What do we know about the language of laypersons, inside and outside the formal health care setting? How do we discover it? And how has this knowledge been used to contribute to controlled vocabularies and thesauri for health information representation and provision?

The discussion that follows reviews (1) research in library and information studies (LIS) and in medicine and nursing, in which the language of users is related to controlled vocabularies and thesauri for information systems, and (2) studies in these domains that focus on the direct language, including health language, of laypeople. “Laypeople” are frequently “consumers” in both large bodies of literature, a group that includes but is not entirely synonymous with “patients.” The article concludes with a reflection on the implications of these findings for social networks devoted to patients and the patient experience.

METHODS

Searches were conducted during the month of May 2010, and databases were searched from the beginning of each file, as noted below.

Consumer language

To locate studies on the relationship of consumers to controlled vocabularies and thesauri, the library and information studies databases used were Library and Information Science Abstracts (LISA; CSA, file begins 1969–), Library and Information Science and Technology Abstracts (LISTA; EBSCO, file begins 1961–), and Library Literature and Information Science/Library Literature and Information Science Retro (HW Wilson, current file begins 1984, Retro file begins 1905). No document types were excluded in these searches. The only limitations were to English-language articles (note that LISTA does not permit limiting by language of publication, so filtering had to be done manually). The LISA search strategy was:

\[
\text{([DE=}(\text{"controlled vocabulary" or "thesauri"}) \text{ AND (KW= user OR end-user OR customer OR client)})]
\]

Total articles found were 109. The LISTA search strategy was:

\[
\text{(DE=Subject headings) AND (user OR end-user OR customer OR client)}
\]

“User OR end-user OR customer OR client” was searched as “All Text.” Total articles found were 162. The Library Literature Search strategy was:

\[
\text{((user <in> Keyword OR end-user <in> Keyword OR customer OR consumer OR client) <in> Keyword) AND ("Terminology" OR "Authority control" OR "Indexing vocabularies") OR ("Indexing vocabularies" OR "Thesauri")}
\]

Total articles were 109 in the current file and 5 in the retrospective file. Terms not identified as “keyword” were searched as “Subject.”

Patient language

To locate studies on patient language, the databases searched were PubMed MEDLINE (file begins 1965) and CINAHLPlus (EBSCO, file begins 1937). Document types were chosen to focus on research about language, as opposed to research on patients in which language was one of multiple facets being investigated. For PubMed MEDLINE, included document types were: meta-analysis, review, address, bibliography, biography, classical article, comment, comparative study, congress, corrected or republished, duplicate publication, English abstract, evaluation study, fest-schrift, government publication, historical article, interview, journal article, introductory journal article, lecture, published erratum, retracted publication, technical report, and validation study. No document types were excluded in the CINAHL search. Controlled vocabulary terms in Medical Subject Headings (MeSH) and CINAHL Headings were all major. All articles were limited to the English language. The PubMed MEDLINE search strategy was:

\[
\text{[\text{"patient language" [searched in All Fields]] OR [Patients AND (Terminology as Topic OR Vocabulary OR Communication Barriers OR Language)]}}
\]

Total articles found were 170. The CINAHL search strategy was:

\[
\text{[Patients AND (Language OR Vocabulary, Controlled)] OR ["patient language" [searched as keyword in All Text]}}
\]

Total articles found were 52.

Eliminated from the literature review that follows were articles on bi- or multilingual patients when the focus was the native language of the patient (for example, preparation of patient education materials in Spanish or translation of a survey or test instrument into another language) and articles about email communication between physicians and patients, unless terminology was the major point of the article.

RESULTS

The library and information science databases had an overlap of 29 articles (8%, 5 due to internal database duplication, leaving 24 or 6%). Only 5 citations (1%) were shared by LISA, LISTA, and Library Literature. Fourteen articles were found in LISA, 16 in LISTA, and 9 in Library Literature’s combined files that dealt with user contributions to controlled vocabulary and thesauri (Table 1). In the health-related databases, CINAHL and MEDLINE had an overlap of 5 articles.
(2%). One hundred forty-one relevant articles were identified in MEDLINE and 7 in CINAHL relating to patient language (Table 1).

**Consumer language: library and information science literature**

Knowledge representation by users, not builders, is a key element of Web 2.0, because user-generated keywords, or tags, are a critical feature of SNS. Tagging by web users enables retrieval of information according to community attributes of interest. The community orientation of tagging is so strong that a person who customizes narrowly—with tags perceived as esoteric, idiosyncratic, and thus socially unuseful—is called a “selfish tagger”; the individual is compiling a “personomy” [11]. Tufekci [12] argues that SNS are a feature of “expressive” Internet use, and an SNS user’s profile has been called “a representation of the self” [13].

An SNS for a specific population can be seen as a community information system, depending on “a tight interplay between the organization of knowledge and communicative processes within communities of practice” [14]. There must exist a complete and accurate understanding of the community, the information it exchanges, and the recipients of that information. Information systems can represent the institutions that create them; text can “represent the institution as an authoritative source in information provision and decision-making procedures affecting the patient” [15]. However, system and user need to understand each other, or “We may be alienating a user community by not speaking their language” [16].

When an SNS is focused on health status, knowledge representation for information exchange presents a particular challenge. In the Web 2.0 era, builders and users may be two discrete but overlapping groups of people. User-created content may drive the information provided by the system. Arguably, folksonomies become collections of user-created descriptors for that user-created content.

The estimated 20% of web users who browse via links are best served by lists of readable, understandable, meaningful labels as guides to navigation [17]. Tags that are ranked by popularity can be labels and constitute a “self-rewarding positive loop” [18]. Immediate community feedback in SNS assists retrieval: “A user can clarify and focus on her own image or concept of her need. The structure of the index language serves as a catalyst” [19]. Furthermore, the knowledge represented in user-generated tags is graphically displayed simultaneously as the medium and the object of collaboration [20]. Individual members of purposeful communities thus increase the quality of knowledge representation geometrically, because their representations engender knowledge work by other users in turn.

The field of information science is where the automation of controlled vocabularies began. An early question asked by information scientists was: Who controls the controlled vocabulary? One author found user-generated terms not useful: “[If no [controlled] index vocabulary is available, do not attempt to generate your own” [21]. However, others argued for a “committee approach” to thesaurus construction, suggesting that users could be a part of this approach: A “thesaurus for sociology or political science should not only be useful for the specialists, but also for the common man” [22].

One early research study on personal indexing systems suggested that personal index terms themselves could populate a classification system. One advantage was that the user would be choosing keywords out of the full text of the document: “They are his own selection and he knows them….As the user, he should in any case have a strong voice in the selection” [23]. Strong and Drott argued almost twenty years later for “a mechanism…for users to suggest their own facets when they do not find the thesaurus facets adequate” [24].

In information science, however, these were minority voices. LIS studies of users and terms have not relied on user-submitted terms. Instead, researchers have investigated users’ choices from existing lists to expand queries [25] or to test the “fit” of captured terms. Today’s tagging researchers do the same thing with captured tags: Library of Congress Subject Headings cataloging terms (Adler [26]), Connotea user tags, and MeSH terms (Lin et al. [27]), while Daly and Ballantyne solicited tags from a known image community [28]. However, in all of these “extent-of-match” studies, users are not, themselves, presented with a thesaurus. Researchers who ask users for terms typically do so to increase the relevance of returned results. Palmquist and Balakrishnan wanted to understand more about users’ unexpressed ideas [29], while Wacholder and Liu [30] investigated searches’ preference for human-constructed versus computer-generated term sets. Again, neither study related

| Database                                      | Total citations retrieved | Citations about user contributions to thesauri or patient language |
|-----------------------------------------------|---------------------------|-------------------------------------------------------------------|
| Library and Information Science Abstracts     | 109                       | 14                                                                |
| Library and Information Science and Technology Abstracts | 162                       | 16                                                                |
| Library Literature (current and Retro file)   | 114                       | 9                                                                 |
| PubMed MEDLINE                                | 170                       | 141                                                               |
| CINAHL                                        | 52                        | 7                                                                 |

*Table 1: Citations reviewed*
users’ terms to either a controlled vocabulary or thesaurus.

In information science research, then, user queries become proxies for the larger, implicitly expressed subtextual information need. The researcher/scientist mediates between user and thesaurus, as librarians have historically mediated between user and system. As in indexing research, “Few studies mention user participation” [31]. Transaction log analysis and similar studies are called exemplars of “user-centered” research [32]. However, users are typically represented in this literature primarily as end users: people existing in relation to systems, not thought about unless they are sitting at a terminal, and certainly not people explicitly involved in controlled vocabularies and thesaurus building. Three notable exceptions in the LIS literature are found, all involving dictionary building: actual musicians contributing to a “musician’s word list” [33], heroin addicts compiling a glossary of drug addiction [34], and indigenous language–speaking community members helping to develop a dictionary of medical terms [35].

Thesaurus builders, unlike dictionary builders, tend to present an existing list of thesaurus terms to users and then ask for their opinions, rather than solicit terms themselves. Commercial and government systems developers typically solicit user feedback as a continuous quality improvement process, as described for the development of the CINAHL thesaurus [36]. User interaction with the existing controlled vocabulary is occasionally mentioned in the literature as a maintenance issue [37]. But in general, the user as contributor has been absent from the literature of thesaurus development since the 1980s. Is the situation any different in the realm of health care?

**Patient language: medical and nursing literature**

*Human medicine is the only scientific field in which the subject literally tells the scientist what the problem is.* [38]

When the ideas of “patient” and “language” are discussed in the literature of medicine and nursing, the intention is seldom to discuss “language of patients.” Instead, patients are typically represented either as “receivers” of language or as members of a group that health care providers are talking or writing about. For example, the patient is a member of health care’s target audience for patient education (Lambert et al. investigated causes of patient confusion about drugs [39]; Stapleton et al. studied the effect of word usage in midwife consultations [40]). Other researchers study what we call patients in health policy [41], mental health [42], and particularly obstetrics [43]. Finally, health care professional language is itself a subject of much study, inasmuch as it relates to patient comprehension of the language and its impact on the patient receiving it (Nordby provides an excellent summary [44]). Researchers consider semantic gaps in the meaning of “asthma” [45], “life expectancy” [46], “gift” (in the context of organ donation) [47], “back pain” [48], “black” and “white” (in the context of moral values) [49], and “euthanasia” and “assisted suicide” [50]. The following discussion excludes these themes, each of which has a copious literature of its own, but instead considers a smaller body of work: studies concerning language used by patients.

The importance of data contributed by the patient—and of asking the patient the right questions—was realized as early as 1000 AD by Rufus of Ephesus, who wrote, “It is important to ask questions of patients because with the help of those questions one will know more exactly some of the things that concern disease, and one will treat the disease better” [51]. The cognitive anthropologist Charles Frake stressed the importance of questions and answers in linguistic discovery of representations of illness: “For every response, the set of inquiries which appropriately evoke the response should also be discovered” [52].

Literature about patients’ own language does suggest some general conclusions about the usefulness of patient-generated verbal descriptors for understanding patients’ health status. It also identifies some factors affecting these verbal descriptors and their susceptibility to processing by information systems. The oldest and most well-documented form of patient language must be the chief complaint, “the patient’s primary reason for seeking medical care” expressed during medical history-taking [53]. The stress on eliciting the patient’s own words whenever possible was made by William Osler, who wrote, “In taking histories…ask no leading questions….Give the patient’s own words in the complaint” [54]. Later medical educators extended this logically to suggest that physicians literally, verbally, echo the language of the patient during the interview, “[A]voiding, of course, the use of four letter words or obscenities.”

Orthographic rules exist for history-taking precisely in order to distinguish patient language from clinician-mediated language: “Responses should be recorded as nearly as possible verbatim which should be indicated by quotation marks” [53], a phenomenon still observed in natural language processing research. Medical informatics research has considered the chief complaint as a data source. No standard coding terminology exists for data of this kind, which exhibits characteristics “idiosyncratic to a specific area or hospital” [55]. This, in turn, presents all kinds of problems for natural language processing, among them, all the communications hurdles encountered in verbal expression, for example, synonyms and paraphrasing.

Patient language appears in the literature of health care in several other domains. First is the work by developers of pain assessment instruments. Pain language is an experience so personal, so individual, and so subjective that no “gold standard” exists for describing clinical pain [56]; thus, “The only way to successfully assess pain is to believe the patient” [57].

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Reliance on patient language can cause difficulties in knowledge representation, however, when words fail the patient: “No pathognomonic sensory descriptor exists for neuropathic pain” [58]. Thus, language in pain assessment instruments was first used to help quantify how much pain the patient was in and, using “the verbal judgment of the patient,” to objectively represent that subjective experience [59]. This high subjectivity of pain descriptors means, too, that they have little predictive power in discriminating between kinds of pain or kinds of patients with pain. Putzke et al. found of pain instruments that “12 of 15 words...[were] selected by >20% of subjects across a variety of pain populations” [60], for example, acute versus chronic pain sufferers [61], but Mauro et al. found no statistically significant differences [62]. In pain research, patients serve not only as recipients, but also as judges of pain instruments (of the Verbal Rating Scale [59, 63], the Visual Analogue Scale [56, 59, 63, 64], the Verbal Descriptor Checklist [64], and the Tursky Pain Perception Profile [61], among others). The ubiquitous McGill Pain Questionnaire [65] has been translated into other languages, a process itself documented in the research literature [56, 62, 66].

Most importantly, patients themselves have spoken about words and pain. A majority in one study preferred verbal over quantitative scales, finding words “easier to understand” and saying they “felt more comfortable using words than numbers,” because verbal descriptors “allowed them better communication with their physician about their pain experience” [67]. Verbal descriptors may convey greater subtlety of meaning than numbers, because they use “natural language of the person...and [have] an inherent face validity” [61]. Clark, Gironda, and Young found a direct correlation between years of education and a preference for verbal, as opposed to numeric, scales [67]. Other cultural factors have been shown to play a role in how people verbally express pain [68], including “anxiety, fear and depression” and age, which has been found to affect subjects’ interpretation of verbal pain descriptors [69]. Particular cultures have “particular semiotics of pain expression” [70], with a terminological effect: “In some languages more than a dozen specific pain terms are in common use, each indicating a particular pain experience, while in other languages a single inclusive term is the norm” [62].

As was noted above, patient language has been investigated as a contributor to physician-patient communication dysfunction, but researchers interested in semantic gaps usually present patients with a list of words, as opposed to asking patients to generate words. For example, Vincent et al. [71] studied patient terms for worsening, and Levin [72] looked at Xhosa patient versus English-speaking physician labels for the same health concepts. Bernstein reminds, like Osler before him, that “patients alone know what they are feeling” [73], but the patient’s verbal expression of that feeling needs to be understood by the listener. Only Yoos et al. asked open-ended questions of young asthmatics and their parents and allowed them to “generate” terms for their own asthma symptoms [74].

Patient language has also been investigated from sociocultural perspectives. Röndahl, Innala, and Carlsson were concerned about sexual orientation—of both nurses and patients—as a factor implicated in “very cautious communication from both personnel and patients” [70]. Schouten and Meeuwesen’s three-year review of medicine and psychology literature found linguistic barriers to be a key predictor of cross-cultural communication problems [76]. The most common type of study found involved ethnic minority patients visiting white physicians, who were then compared with white patients visiting white physicians. Language concordance was found to have an effect in several of these studies [77]. Gender was investigated by pain researchers Vodopiutz et al., but few gender differences were found [78]. Bischoff, Hudelson, and Bovier asked what difference gender made to language discordance between physician and patient and found interpreters to function not just as mediators of language, but as mediators of culture [79].

Vocabulary development is one way in which information systems communicate with users. The information needs of health care professionals have, of course, been the driving force behind many medical informatics initiatives, and their origins are reflected in these systems’ vocabularies. For example, Stetson et al. attempted to develop an ontology modeling the contribution of clinical communication problems to clinical errors [80]. Health paraprofessionals like medical librarians have also served as human mediators of needs for the purposes of system design [81], particularly for ontology development [82]. The most extreme expression of user and librarian involvement found thus far has been the Faculty Research Interests Project (FRIP) at the University of Pittsburgh [83]. The University of Pittsburgh’s clinical faculty and researchers were profiled in a database that was pre-populated with mined MeSH terms attached to their publications. These authors were asked to supply additional author-generated keywords as they thought necessary. The keywords were then displayed online alongside the MeSH terms as part of the researcher’s profile. Searchers of FRIP could thus use these keywords side by side with a browsable thesaurus, as terminological assists and search augmentations [84]. Users were generating terms to describe themselves, or, at least, that portion of themselves revealed by their MEDLINE-indexed publications. This prefigured the practice of tagging in Web 2.0 by several years.

However, FRIP was concerned with profiling biomedical clinicians and researchers. What about systems profiling consumers and patients? How have lay people contributed their own language to knowledge representation in health care information systems?

In medical informatics, the professional-lay communication gap has been operationalized as “the consumer health vocabulary problem” [85]. Consum-
er health language research, and specifically vocabulary development targeting consumers, was identified by Keselman et al. as one important informatics strategy for addressing professional-consumer communication problems in health care [10]. Medical informatics researchers have explored the extent and severity of the consumer-professional gap [86, 87], its role in communications dysfunctions, and the implications for health literacy initiatives [88]. Exploration in this “consumer health vocabulary” subdomain, however, has typically been done to describe and demarcate what “consumer” territory exists, highlighting its difference from professional territory. It is not done explicitly in the interest of vocabulary development and control, although enhancing thesauri through addition of synonyms and entry terms is a desired corollary outcome.

Where does lay language come from in medical informatics? Various extant “consumer” texts, sources as diverse as the Dictionary of American Regional English [85] and emails [88], have been explored as sources of terms. An example is a large extent-of-match study, focusing on consumer utterances and using data from the Medical Library Association–funded project Ten Thousand Questions [89]. Consumers and patients are not typically “present” in this research: They are represented in the aggregate, and in absentia, by the trails they leave in the system. For example, patients are represented by query logs at ClinicalTrials.gov [90], MedlinePlus [91], or “Ask-A-Doctor” sites [92]. Other online consumer “speech” includes emails to a university health website [93], a cancer information service [94], or nurses caring for specific patients [95, 96]. Web-based bulletin board posts have also been studied for their characteristics and content [97, 98]. Typically, the data generated in these research explorations derive from text capture and are obtained from large groups of anonymous “speakers,” operationalized as “consumers” because any more specific identity is unknown to the researcher. If consumers, including patients, are consulted for their term preferences in an information systems context, it is typically via focus groups and interviews, such as those described by Slaughter, Ruland, and Rotega [99]. These focus groups are feedback groups: Like the previously discussed patients in pain studies, they rate existing lists but do not build new lists. One interesting near-exception was the study done by Zeng et al., using a mix of transaction log analysis and patient interviews. Queries logged at a “Find A Doctor” site were mined for content, and individual patients seen at the same hospital were interviewed in order to test their understanding of frequently used terms taken from the query logs [100]. Most common, unfortunately, is the kind of study in which the verbal expressions of patients are channeled by clinicians, so that the data are always secondhand. For example, plastic surgeons were interested in developing a set of lay synonyms for body parts to help in clinical consultations about liposuction and “made lists...we have heard from our patient populations over the years” [101].

Health communications researchers have written about the medical interview and its impact on patient-physician relations. Lindfors and Raevarla studied emergency department patients and discovered that patient language not only can be transformed by physicians, but it can apparently be conversationally transmissible. Patients were admitted to the emergency department with symptoms of chest pain and then observed during medical history taking. Health care providers perceived the patient’s expressions of pain as “vague,” which frustrated them. They then began to structure the interactions by verbalizing a “menu of potential responses,” naturally associated with their professional, clinical conceptions about the likely basis of the pain. The prompting of the emergency department physicians began to structure the patient’s own verbal expressions. The more trouble the patient had in communicating, the more suggestions were on the physician’s “menu.” After repeated interviews, the patient was “trained” to express symptoms in more focused—and more professional—language, increasingly citing physician expertise and increasingly quoting physician statements that incorporated more medical terminology, for example, “My doctor said I have angina.” The conclusion of these researchers could have been stated by any website designer or information architect who attempts to create a browsable web-based ontology: Patients “required to select from a finite number of descriptors...may not find a word that accurately reflects their experience” [103]. In fact, it may reflect somebody else’s experience: their doctors’.

Finally, pain researcher Nutkiewicz studied thirty-two children from ethnically diverse backgrounds seen for chronic pain. He compared the use of “oral testimony,” or “the children’s own words,” to find that physician language is “informative and directive” versus the patients’ “expressive, subjective, and experiential” narrative: “Children and their doctors have two separate orientations toward pain and employ two separate vocabularies to describe pain that are inextricably linked to these orientations...[They] appear as tacit and embedded approaches to disease and illness” [104]. Nutkiewicz is describing the challenge that is central to understanding patient language and the patient’s part in the conversation that is health care: to “name the illness in a way that is meaningful to physician and patient, wherein the patient’s experience of illness is validated and accepted untransformed [italics added] and then later reconciled with the physician’s diagnostic categories” [105].
DISCUSSION

This literature review has identified a serious lack of information about consumer contributions to controlled vocabulary, which appears to be a seriously under-researched area inside and outside of health care. The growing interest in folksonomy research among library scientists, information scientists, and computer scientists is a positive sign—but only a sign—that this deficiency is potentially reversible. Health care researchers need to engage with laypeople in their roles as patients to ensure that information systems can truly support health communication between laypeople and health care providers.

It can be argued that systems that deliver information run the risk of reinforcing the information designer’s communication biases. For example, Beach has argued that research into the medical interview resembles the interview itself: It reiterates “medical authority and the institutional character of professional/lay communication” [106]. The same might be said of research into patient language, inevitably constricted by the fact that the patient is a patient, being observed and recorded by people seeing the individual as a patient. Sarangi contends that the assessment of “patient talk” has in fact meant attention to “responses to physician questions” [107]. As Keselman et al. have written, “Paradoxically, there is voluminous literature on the information needs of health care professionals but very little on those of patients and little about the needs of the general public. In practice, systems design is typically guided by the providers’ perception of patients’ information needs, rather than by actual needs assessment” [10]. This shows the effect of what Foucault called the “clinical gaze”: By defining the person as a patient, the physician also defines the direction and the content of the subsequent conversation [108].

Making internalized understanding externally visible for the use of external others has always been a difficult task. Decades of informatics work on clinical data standards makes clear that symptoms, like other expressions of the lived patient experience, are both “unconscious and procedural...hard to formalize and communicate to others” [109]. Forsythe commented on the same problem for representing clinical information: “the tacit, taken-for-granted, non-standardized information so essential to comprehension in particular situations” [110]. She used ethnographic methods precisely because of their value in eliciting implicit knowledge for explicit representation.

What are the consequences of Foucault’s clinical gaze for information systems that serve the needs of patients and consumers? Oudshoorn and Somers looked at three Dutch patient organizations and the websites they built. One site, maintained by health professionals, focused on clinical depression. The second and third were maintained by patients, people living with cancer and repetitive stress injuries (RSI). A dichotomy was found between “implicit” and “explicit” techniques used for knowledge represen-
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**AUTHOR’S AFFILIATION**

Catherine A. Smith, MA, MILS, MSIS, PhD, casmith24@wisc.edu, Assistant Professor, School of Library and Information Studies, University of Wisconsin-Madison, 600 North Park Street, #4255, Madison, WI 53706

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