Scoping review of patients’ attitudes about their role and behaviours to ensure safe care at the direct care level

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Abstract

Background: To improve harm prevention, patient engagement in safety at the direct care level is advocated. For patient safety to most effectively include patients, it is critical to reflect on existing evidence, to better position future research with implications for education and practice.

Methods: As part of a multi-phase study, which included a qualitative descriptive study (Duhn & Medves, 2018), a scoping review about patient engagement in safety was conducted. The objective was to review papers about patients’ attitudes and behaviours concerning their involvement in ensuring their safe care. The databases searched included MEDLINE, CINAHL and EMBASE (year ending 2019).

Results: This review included 35 papers about “Patient Attitudes” and 125 papers about “Patient Behaviours”—indicative of growing global interest in this field. Several patterns emerged from the review, including that most investigators have focused on a particular dimension of harm prevention, such as asking about provider handwashing, and there is less known about patients’ opinions about their role in safety generally and how to actualize it in a way that is right for them. While patients may indicate favourable attitudes toward safety involvement generally, intention to act or actual behaviours may be quite different.

Conclusion: This review, given its multi-focus across the continuum of care, is the first of its kind based on existing literature. It provides an important international “mapping” of the initiatives that are underway to engage patients in different elements of safety and their viewpoints, and identifies the gaps that remain.

Keywords
direct care level, patient attitudes, patient behaviours, patient engagement, patient safety, scoping review
1 | INTRODUCTION

The need to improve patient safety continues to be paramount. Patient safety is recognized as a serious global public health issue, with approximately one in 20 patients harmed while receiving medical care (primary, secondary and tertiary care settings). When considering patient safety, it is important to understand the perspectives of patients and their families, and to recognize the need for more effective involvement of patients and families across the continuum of care. As recipients of health-care services, patients and their families are a valuable resource and contribute to our understanding of safety issues. To advance our knowledge about patient and family involvement in promoting health-care safety, as distinct from their health generally, a multi-phase study was conducted that included a qualitative study of patients' perspectives about their knowledge, comfort level and behaviours in helping their safety while receiving health care. This scoping review was phase 2, and the aim was to describe the literature about patients' and families' attitudes toward their role in health-care safety, as well as their reported behaviours in support of their safe care. A general multidimensional framework about patient and family engagement in health and health care underpinned this study.

2 | BACKGROUND

In preparation for the multi-phase study, a preliminary examination of the literature, which included findings from previous unpublished work, provided an initial understanding of the evidence about patient perspectives regarding health-care safety, and views on their involvement in safety processes. The findings from this preparatory work, which helped determine the need for a scoping review, included the following concepts.

2.1 | Communication

Patients and/or their families specifically alluded to communication as a vital part of patient safety. Participant views on this theme were common across primary care, ambulatory, hospital and home settings. In a Canadian telephone survey of 1,500 adults, respondents perceived that communication was associated with preventable medical errors. In the primary care studies, safety problems were linked to miscommunication and/or a lack in communication of test results.

2.2 | Trust

The concept of trust in the health-care system and providers was also a common theme related to patient safety as perceived by patients and families. The overwhelming need of “safety” in ill hospitalized patients was tempered by the knowledge of being able to trust staff. A group of 30 ambulatory patients receiving chemotherapy treatment recognized their own limitations in safety prevention, including deficits in knowledge, and therefore felt they had to trust the providers. In another example, some parents, who believed their children had experienced a medical error, expressed regret at having trusted a health-care professional.

2.3 | Being trusted, seen and valued

Patients and their families view being trusted for their knowledge, respected and valued for their contribution as integral to safe patient care across hospital, outpatient and home settings. Reports of patients’ experiences in primary care echoed the related theme of difficulties in interpersonal relationships with providers, particularly noting disrespect and insensitivity. A study of 20 individuals with Parkinson’s disease (or someone who spoke on their behalf—partner, wife, husband, daughter) and their experiences of medication errors most poignantly captures the lack of respect of patients’ knowledge related to their medication management and the de-valuing of their insights—the cost of which can jeopardize a patient’s safety.

2.4 | Patients as partners

Patients’ beliefs about participating in patient safety in different ways have also been investigated. Positive attitudes about safety engagement have been identified, with qualifiers such as the type of action required by the patient, practitioners receiving this level of involvement by patients and their families, as well as the setting. Tactics that patients (or family members of patients) use to protect themselves when interacting with the health-care system have been reported, including bringing family or friends to appointments and questioning unfamiliar medication. Patient involvement in ameliorating harm has also been documented. Generally, the most common approach to engage patients in health-care safety has taken the form of patient “tips” or strategies outlined in patient information or educational materials. There is evidence that some of these tips have been created with input from patients; however, this is not true of all recommendations, with a need for evidence-based content and evaluation.

Given the imperative of considering all approaches to reduce harm, as well as the importance of being patient- and family-centred, the preliminary work identified gaps and additional questions (including do patients intentionally think about their involvement in safety at the direct care level, and what does safety mean to them) necessitating and providing rationale to explore international evidence in a more fulsome, comprehensive way related to patient and family involvement in safety at the direct care level.
3 | METHODS

The scoping review was conducted to understand the breadth and depth of literature about patients’ engagement in safe care. The intention of the review was to gather as much relevant literature about patients’ attitudes toward having a role in preventing harm, as well as any safety behaviours they independently engage in or at the invitation of researchers or providers.

This scoping review was conducted using the search term “patient safety”, but other iterations were also considered given patients/families may understand it in different ways. Additionally, the preparatory work was informative in identifying that researchers typically investigate components of safety (eg handwashing), and therefore, these elements were part of the search strategy. A search of the JBI (Joanna Briggs Institute) Evidence Synthesis Journal and Cochrane database indicated that no scoping review had been conducted on this topic.

3.1 | Research questions

(a) What are patients’ and families’ attitudes and beliefs about their participation role in ensuring they receive safe care? (b) What are the behaviours indicative of harm prevention that patients and families engage in independently or at the direction of others [researchers; health-care providers]?

3.2 | Research design

Peters and colleagues indicate the appropriateness of a scoping review when “a body of literature...exhibits a large, complex or heterogeneous nature not amenable to a more precise systematic review.” A generic methodology was taken to scope the literature on this topic, and this was overseen by the first author (LD) as part of her doctoral thesis, who was responsible for the management of data collection and analysis. Decisions were made primarily by the first author, and the thesis committee provided assistance if there were disagreements. Scoping reviews are unique in that they are typically about broader topics; often include many different study designs; and do not usually involve an assessment of the quality of studies that are included in the review.

3.3 | Study inclusion criteria

The PCC format for scoping reviews is an acronym for P = population; C = concept; and C = context. The review was about a population which included all individuals engaged in the health-care system as patients, family members of patients or health-care consumers (no age criteria). The concepts were twofold, but both included the focus on safety actions as distinct from caring for one’s health. First, it was the patients’ and families’ attitudes, opinions and beliefs about taking an active role to ensure safe care at the direct care level. Second, it was the patients’ and family members’ behaviours or actions that demonstrated active engagement and involvement to prevent harm at the direct care level. The safety behaviours or actions were either self-determined or directed as part of a study intervention. The review included evidence of whether patients and family members performed requested safety behaviours, or if they demonstrated self-determined strategies, to prevent harm while receiving health care. The context included all sectors of health care including acute care, ambulatory care and long-term care. Home settings (people’s own home) were seen as a separate context and as such were excluded, unless they were included as part of multiple sites in a relevant paper. Direct care level was defined as any clinical care interaction, as based on the study framework.

3.4 | Types of literature

This review included systematic reviews, quantitative studies, qualitative studies, mixed-methods studies, scoping reviews, literature reviews, quality improvement projects, as well as opinion and discussion papers. Study or systematic review protocols were also included in order to give perspective on the interest in this topic and anticipated future results. It is acknowledged that some may not be actioned into full reviews but this does reflect current research activity. Studies published in abstract form (such as conference proceedings) were not included.

3.5 | Search strategy

A three-step search strategy was used that considered published and unpublished papers written in English. An initial limited search of MEDLINE, CINAHL and EMBASE was undertaken followed by analysis of the text words contained in the title and abstract and the index terms used to describe the article. A second search using all identified keywords and index terms was undertaken across all included databases. Third, the reference lists and bibliographies of select reports and articles were searched for additional studies, and selections made based on title. Papers written only as abstracts were excluded. Two separate search strategies for the two research questions were conducted given each had a different focus. The databases searched (during 2016-2019) included the following: JBI (Joanna Briggs Institute) Evidence Synthesis (1998-2019); PsycINFO (1806-2019); CINAHL (1981-2019); MEDLINE (1946-2019); EMBASE (1947-2019); and ProQuest Dissertations and Theses Global (1861-2019). Table 1 is a list of the initial keywords used for the first search about patient attitudes and beliefs, as well as the second search on patient behaviours.

The review also included searching of organizational, governmental and advocacy group websites, as well as communications...
with patient safety experts for relevant papers and texts. The websites of 44 organizations (13 local and provincial organizations; 11 national organizations; and 20 international organizations [eg Institute for Healthcare Improvement, World Health Organization]) were searched focusing on the study objectives (see Appendix S1 for the site listing). In the searching of websites, given their design, it was most effective to search using general terms, such as “patient safety” or “patient beliefs on role in safety”, or as based on the provided categories and headings.

**TABLE 1** Initial keywords for scoping review search strategy

| Patients/ Family | Attitudes | Involvement | Safety |
|------------------|-----------|-------------|--------|
| *patients*; *family*; *siblings*; *parents* | attitude; beliefs; opinions; patient perception; patient perspective; values; understanding | patient participation; role; consumer participation; patient involvement; contribution to health care; patient advocacy; consumer advocacy; advocacy | safety; patient safety; medical errors; safety management; adverse events; vigilant |

**Initial keywords used for the second search about Patient Behaviours**

| Patients/ Family | Behaviours | Safety |
|------------------|------------|--------|
| *patients*; *family*; *siblings*; *parents* | behaviours; self-efficacy; patient education; patient empowerment; patient participation; consumer participation; patient involvement; patient engagement; patient advocacy; speaking up | patient safety; health-care error—prevention and control; safety management; adverse health-care event; handwashing; medication safety; surgical safety; infection control—prevention and control; patient handoff; safeguarding |

Note: The terms within each column were entered using ‘OR’, and the full sets of terms in each were combined across columns using “AND”. **“exp”** used with these terms to provide full scope of definition.

**FIGURE 1** PRISMA diagram
3.6 | Procedure

The associated university research ethics board granted ethics approval for the multi-phase study, which included this scoping review (HSREB 6007637, NURS-299-12).

3.7 | Data collection and extraction

A standard approach was used to extract quantitative, qualitative and textual data from the literature about patients’ and families’ attitudes and behaviours related to their role in ensuring safe care. For all papers, general items, such as author(s), location of first author, title of document, name of organization or association (as relevant) and year of publication, were extracted. As relevant, data extraction also included, although not limited to, methodology, methods (eg interventions, sample, setting, data collection methods) and findings.

3.8 | Analysis

The analysis of the extracted data from the papers (formatted as tables) for this phase occurred in several ways. A descriptive approach was taken to determine the quantity of publications by type, author, location of author, year of publication and topic (ie attitudes versus behaviours; behaviours were further sub-divided into specific categories). Additionally, and as applicable, the results from the obtained literature were examined as a collective (published and unpublished papers together) using a content analysis approach for overall themes or patterns. This analysis was guided by the process as outlined by Miles, Huberman and Saldana, including an initial coding process, followed by second cycle categorization into the larger groupings/themes. The results of this review are presented in narrative form. Study designs are reported as published by the author(s).

4 | RESULTS

This review includes a final set of 151 individual papers: 35 about Patient Attitudes and 125 about Patient Behaviours (Note: nine papers were in both sets given their multi-focus). A PRISMA diagram, as reflective of the search strategy for this review, is presented in Figure 1.

Of the 151 papers included in the final set of this review, there were 13 systematic reviews; one scoping review; six systematic review or study protocols; 53 quantitative studies; 29 qualitative studies; eight mixed-methods studies; eight literature reviews; 29 opinion or discussion papers; and five quality improvement projects. Table 2 depicts a listing of the number of publications by the first author’s location, and Table 3 provides the distribution of the publication dates.

| TABLE 2 | Scoping review final set: by location of first author |
| Location | Number of publications (n = 151) |
|----------|---------------------------------|
| United States of America | 52 |
| United Kingdom | 29 |
| Australia | 18 |
| Canada | 16 |
| Switzerland | 8 |
| China | 6 |
| Denmark | 4 |
| Korea/Republic of Korea | 3 |
| Germany | 2 |
| Italy | 2 |
| Sweden | 2 |
| Belgium | 1 |
| Czech Republic | 1 |
| Finland | 1 |
| Iran | 1 |
| Israel | 1 |
| Japan | 1 |
| Norway | 1 |
| Singapore | 1 |
| Spain | 1 |

| TABLE 3 | Scoping review final set: publication date distribution |
| Year of publication | Number of publications (n = 151) |
|----------------------|---------------------------------|
| 2019 | 8 |
| 2018 | 14 |
| 2017 | 21 |
| 2016 | 11 |
| 2015 | 9 |
| 2014 | 17 |
| 2013 | 14 |
| 2012 | 10 |
| 2011 | 14 |
| 2010 | 12 |
| 2009 | 2 |
| 2008 | 2 |
| 2007 | 7 |
| 2006 | 3 |
| 2005 | 0 |
| 2004 | 3 |
| 2003 | 3 |
| 2002 | 1 |
| 2001 | 0 |
| 2000 | 0 |
4.1 | Descriptive summary: Patient attitudes

Thirty-five papers about patient attitudes or feelings about being involved to protect themselves when receiving health care were identified (see Appendix S2). This included three systematic reviews, eight quantitative studies, 13 qualitative studies, one mixed-method study, one literature review, nine text and opinion pieces, and one study protocol. Publication dates ranged from 2004 to 2019 [most in: 2011 (n = 6)]. The countries of the first authors of the publications were as follows: United States of America (n = 13), United Kingdom (n = 9), Switzerland (n = 4), Canada (n = 3), Australia, Czech Republic, Denmark, Iran, Korea and Sweden (n = 1).

Of those 35 papers, three were systematic reviews. While the McVeety and colleagues review (n = 14 qualitative studies) was primarily about the patients’ and family members’ perspectives having encountered an adverse event, all three reviews addressed patients’ and family members’ feelings about safeguarding and beliefs about safety actions. The combined total of included studies for the three reviews was 79, with a total overlap of 24 studies. One literature review also contained some of the same articles (n = 4) as those in the Doherty and Stavropoulou systematic review. In the qualitative review by McVeety and colleagues, the number of participants ranged from 10 to 172 and included interviews (individual and focus group) in the hospital and home setting, as well as part of conference workshops. The review by Doherty and Stavropoulou included studies that had a range of participants from 4 to 2,078 patients, and included patients who were discharged from a hospital, as well as ambulatory patients. The Schwappach review sample sizes ranged from 72 to 2,078 participants and included a variety of methods including face-to-face interviews, focus groups, and mailed or telephone surveys. The remaining 31 papers in this set of 35 were not included in these three systematic reviews but are outlined as follows.

Thirty-one papers about patient attitudes contained the following study designs. (a) Nine quantitative studies: The most common methodology was cross-sectional exploratory studies using a survey with sample sizes ranging from 50 to 1,053 patients, including medical-surgical patients, oncology patients and community dwelling seniors. (b) Eleven qualitative studies: Typically focus groups (range of 2-8) or interviews were used, and participants included those with a recent overnight hospital admission as well as hospitalized oncology patients. Several ethnographic studies that included observations as well as interviews were conducted. (c) One mixed-method study was about patient attitudes. In this Canadian study, survey methods and focus groups were conducted with patients (217 survey responses and 10 in focus groups). (d) One study protocol for an exploratory qualitative study about patient and provider perspectives about the role of patients in safe care was also found. (e) Nine text and opinion papers: These items include books, opinion articles and a white paper. The authors are from the United Kingdom, United States of America, Canada and Switzerland. The publications are either authored by patients or reflect their stories and narratives using quotes.

4.2 | Descriptive summary: Patient behaviours

One hundred twenty-five papers related to patient behaviours, either generally or specific actions that patients are involved in, both independently or when directed by others (eg researchers; providers), to ensure their safe care. This included 12 systematic reviews [handwashing; patient behaviours—general; patient handover; medication safety; technology] [number of studies included in the reviews ranged from six to 68], 44 quantitative studies, 23 qualitative studies, eight mixed-method studies, one scoping review, eight literature reviews, five systematic review or study protocols, 20 text and opinion pieces, and five quality improvement projects. The publication dates ranged from the years 2002 to 2019. The country of the first author of the papers was predominantly the United States of America, the United Kingdom, Australia or Canada.

The 125 papers related to patient behaviours were categorized according to specific topics for ease of reporting and to discern the range of study focus. Table 4 is a list of these categories, and the type of publication and date range for each. Appendix S3 is a detailed listing of the papers included in this subset. The findings between published and unpublished papers in many cases were complementary (dependent on topic), and although not included in this article, the retrieved grey literature about patient safety advisories/tips was immense.

5 | OVERALL THEMES—PATIENT ATTITUDES AND BEHAVIOURS

5.1 | Patient attitudes

Three patterns emerged from the 35 papers about patients’ and family members’ attitudes and beliefs toward having an active role in promoting their safe care.

5.1.1 | Lack of evidence about patient attitudes

The first pattern is the paucity of evidence specifically about patients’ attitudes toward their role generally in regard to safety at the direct care level. Most investigators have focused on a particular dimension of harm prevention, such as patients’ attitudes toward and comfort in asking providers about handwashing (eg four of 13 studies reported in Schwappach review; Davis et al), or intervention strategies, and made inferences from those results to patient safety overall. Those primary studies with a more purposeful focus on understanding patients’ attitudes and thoughts about whether they have a role in safety at the direct care level were few in number, and were sometimes combined with general care activities, or focused on parents. The general findings from these investigations included viewing their role in safety “as a right, not an obligation”, to seeing their role as a
| Category                              | Number in set | Publication data range | Study design or publication type | Quality improvement project |
|---------------------------------------|---------------|------------------------|----------------------------------|-----------------------------|
|                                       |               |                        | Systematic review                 |                             |
|                                       |               |                        | Quantitative                      |                             |
|                                       |               |                        | Qualitative                       |                             |
|                                       |               |                        | Mixed-methods                     |                             |
|                                       |               |                        | Scoping review                    |                             |
|                                       |               |                        | Literature review                 |                             |
|                                       |               |                        | Systematic review or study protocol|                             |
|                                       |               |                        | Opinion/white paper               |                             |
| Handwashing                           | 29            | 2006-2019              | 2                                | 22                          | 1 1 3 3 5 20 5 |
| Patient Behaviours—General            | 26            | 2002-2018              | 5                                | 4                           | 3 2 2 2 10 10 |
| Patient Handover                      | 15            | 2011-2019              | 2                                | 1                           | 6 2 2 1 1 |
| Medication Safety                     | 13            | 2006-2019              | 2                                | 3                           | 4 2 2 2 |
| Communication                         | 8             | 2003-2018              | 3                                | 3                           | 2 2 2 2 2 |
| Patient Reporting                     | 5             | 2007-2017              | 3                                | 1                           | 1 1 |
| Advocacy—Parents/Others              | 5             | 2004-2018              | 4                                | 1                           | 1 1 |
| Ambulatory and Primary Care           | 4             | 2010-2017              | 1                                | 1                           | 1 1 1 |
| Health Literacy and Safety Tips       | 4             | 2003-2012              | 3                                |                             | 1 1 |
| Health-care-associated Infections     | 3             | 2015-2016              | 1                                | 1                           | 1 1 |
| Surgical Safety                       | 3             | 2010-2011              | 1                                |                             | 1 1 1 |
| Technology                            | 3             | 2003-2019              | 1                                | 1                           | 1 1 |
| Rapid Response Team Activation        | 2             | 2014, 2018             | 1                                |                             | 1 1 |
| Diagnostic Imaging/X-ray Imaging      | 2             | 2013, 2017             | 1                                |                             | 1 1 |
| Newborn Fall Prevention               | 1             | 2018                   | 1                                |                             | 1 1 |
| Advocacy—General                     | 1             | 2013                   | 1                                |                             | 1 1 |
| Blood Transfusion                     | 1             | 2011                   | 1                                |                             | 1 1 |
| Total                                 | 125           | 12 44 23 7 1 8 5 20 5 | 20 5                            | 1 1 1 1 1 1 1 1 1 1 1 1 1 1 |
"shared responsibility" with providers. Most importantly, the mapping of this literature identified limitations in our current understanding in this regard, both in terms of depth of evidence and clarity of concepts.

5.1.2 | Patients’ opinions about their role vary

Second, a pattern emerging from the existing evidence indicates patients have varied opinions and perceptions about having an active role in error prevention. There is no consistent, unequivocal agreement among patients and family members about their level of involvement or role in safety. For some patients, they see their role as a passive one and that safety is not their responsibility. For others, they believe patients must participate to help with harm prevention. Generally, different factors, values, understanding and experiences influenced individuals’ attitudes toward their role and involvement in safety in different ways.

There are individuals who believe that safety is the obligation of the health-care provider and that patients do not have that responsibility. As found by some investigators, when the patient (philosophically) views their role as a passive one, they are not willing to be involved in safety and are described as having more submissive attitudes toward safety, as well as believe their only focus is to get well. In one qualitative study, participants expressed fear that the responsibility of safe care would shift to patients and that the factors contributing to errors were beyond their control, and therefore, they could have little influence.

There are individuals who have a different view, believing that patients should be engaged in ensuring safe care, and that they can effectively contribute to safety. Patients have expressed the viewpoint that they know themselves best, and therefore need to be engaged and involved in sharing that knowledge and ensuring it is not dismissed. In a 2011 cross-sectional survey of 1,053 patients, 95% agreed that patients should be educated about how to prevent errors, while other investigators have reported that having the opportunity to discuss safety issues with care was rated favourably by patients. In a small sample of 50 internal medicine patients, 90% wanted to review their medication list for accuracy and 94% believed patient involvement and review of medications had the potential to reduce error. Even when patients had difficulty understanding the specifics of what safety engagement involved, for this group of participants, the willingness to participate, whatever the underlying motivation or inspiration, was evident.

5.1.3 | Patient belief may differ from their action

The third pattern relates to the second, specifically to those patients and consumers who believe they have a role in promoting safe care. While patients may indicate support and favourable attitudes toward safety involvement generally, intention to act or actual behaviours may be different. The Theory of Planned Behaviour has been used as a conceptual framework by several investigators when assessing patient attitudes about engagement in safety. The premise of the theory is that an intention to act is influenced by the attitude toward that behaviour, the perceived influence of others and the perception of ease or difficulty of the behaviour. Behavioural intent is predictive of behaviour. Schwappach suggested that patient engagement in patient safety can be viewed as "...a special case of health-promoting behaviour" (p. 122), as such, the theory can be a useful one in explaining and predicting involvement. In conceptualizing patient safety in this way, it is helpful in identifying that a patient’s positive attitude about safety engagement does not necessarily equate to actual action. A number of investigators have demonstrated that only considering attitude is not sufficient and that other elements are influential, including why a patient will ask about their medications but not a provider about handwashing. The 21-item tool to measure views about safety tasks by Elder et al is an example of how investigators are not only asking individuals how comfortable they are with certain activities, but how often they have completed them. Overall, it is evident that one must be cautious of oversimplifying a positive attitude about patient engagement in patient safety without consideration and examination of other potential influencing factors and qualifiers.

5.2 | Patient behaviours

Four patterns emerged reflecting patients’ and family members’ behaviours and actions in promoting their safe care.

5.2.1 | Escalation in patient engagement research

The first is the increased interest in this topic in recent years, and the evolving range and diversity of how patients are being involved to support the safety of their care. The action of patients asking providers whether they have washed their hands has been a predominant focus of research in this field. Davis and colleagues conducted a systematic review of the evidence on the effectiveness of strategies that increase patient participation in reminding providers about hand hygiene. Their review included a total of 28 articles, and while they reported a number of strategies being examined to increase this type of patient involvement, they found the studies lacked methodological rigour, and cautioned that because most studies were designed to examine patients’ intention about asking, their actual behaviour may be different.

In addition to asking providers about handwashing, researchers have investigated how patients can participate and take action to support safe care related to other care processes, including patient
handover and medication safety. More recently, researchers have examined areas such as diagnostic imaging and primary care to determine whether there are opportunities for patients to participate in safe care in these settings. It seems clear that, with the increasing attention given to patient and family engagement generally and the continued focus on harm prevention, there is greater interest to determine how patients can engage and participate in safety-specific practices.

5.2.2 | Patients are engaged

The second pattern was the indication that patients are engaging in behaviours that promote safe care, either independently in their own ways or as requested to varying degrees and that they see and are aware of safety practices and strategies occurring in the health-care environment. The literature is limited, but researchers have reported that patients have developed strategies to protect themselves, such as taking notes; asking a family member to ask a question on their behalf; seeking information from the Internet; talking with other patients; learning by listening to providers educate each other; and speaking up when concerned and may not even recognize the activities they do as safety measures. Additional research is needed, but current evidence suggests that patients can and do perform actions to promote their safe care, and positive outcomes related to their involvement have been described. Australians Hor and colleagues believe that patients are already involved and that instead of asking whether patients ought to be involved with safety, we should be asking how we can make patient-provider collaborations for safety more effective.

5.2.3 | A focus on influencing and impeding factors to engagement

A third pattern was patients' and family members' behaviours about promoting safe care. Investigators have examined factors that influence or impede patients' participation in safety initiatives. Factors that may support and enhance patient involvement in safety include the following: perceived risk, provider encouragement, perceived self-efficacy and health-care setting (eg primary care is seen as key in developing trusting relations with implications to safety and is seen as more feasible to engage, yet seen as paramount for family members in hospital settings). Issues that may negatively influence an individual's ability to engage in activities to support safe care include the following: severity of one's illness, perception of staff work pressure, lack of awareness of the benefit of their involvement, engaging in a task perceived as challenging authority (versus factual, information-sharing tasks) and belief that one's role should be passive. There is recognition that there will be different motivators and inhibitors that will affect patients' behaviours related to participating in the safety aspects of their care, and while understanding the motivators and reasons for engagement was not a focus of this review, further unpacking influencing factors such as whether one's preference to engage is anxiety or fear-based that safe care could be compromised versus acting based on preference alone is warranted.

5.2.4 | The role of family

The final pattern was the role of family in advocating for and protecting the well-being of another. Parents whose children are requiring health care report and demonstrate a vigilance and protectiveiveness to ensure they are safe. Of 130 parents of hospitalized children, 63% agreed or strongly agreed that they needed to safeguard their child against potential errors. Hurst's critical ethnography of 12 mothers of hospitalized premature babies wrote of how mothers' primary action was to watch over and observe their baby and the care processes. Parents advocated on behalf of their child and were prepared to ask about and challenge processes if needed; however, they were also sensitive and cautious as to what and how they questioned. Parents viewed their role as a protector—a responsibility described as "both their right and their job". In the adult setting, the family role was equally important and members as engaged. Family members acted as observers, noted in the study by Kim, where family members report watching care practices, such as provider handwashing (63% report they agree/strongly agree that they observe provider handwashing; n = 173). Family members reported speaking up about worrying symptoms they see exhibited in their loved one and of their comfort when action is taken by the provider as a result. Patients also called on family members to assist them in ensuring safe care, including asking a family member to question providers on their behalf and the role of family members increase if individuals are too ill or cognitively impaired. Investigators have included family members in studies about safety and error-prevention strategies, though the investigation of family members and their behaviours related to ensuring safe care as a primary focus is limited. Though not the intended focus of this study, the responsibility of nurses to advocate for patients and to support and encourage them in their own advocacy is also an important consideration.

6 | DISCUSSION

There are important considerations as a result of this review. First, this review provides confirmation of the interest internationally, in varying ways and degrees, about patient and family engagement in safety. The number of publications written in English in peer-reviewed journals suggests that many are investigating, writing and commenting on patient involvement in safety. The results of this analysis are one indication, with 21 publications from 2000 to 2009, increasing to 130 from 2010 to 2019. It could be as a result of both overall efforts in patient engagement in health care generally, and in harm prevention. Those organizations that have patient safety...
as their mandate have also expanded their focus to include patient involvement, such as the World Health Organization’s Patients for Patient Safety (an advocacy programme to elevate and profile the rights and viewpoints of patients and families regarding their health, and including collaboration with providers in advancing quality person-centred health care). Additionally, the advantage of mapping this topic provides insight into how patient safety engagement is being parsed, defined or promoted—not always leading to success. The most common focus regarding behaviour was handwashing and specifically, encouraging patients to ask providers whether they have washed their hands prior to providing care. Given substantive limitations in establishing consistent and sustained effectiveness and success rates, continued attempts to engage patients in this area are questionable, or at the very least, a focus on patient-identified strategies may have more beneficial effects on the rates.

Second, there is a need to better understand patients’ attitudes and beliefs about engagement at the direct care level in safety holistically across the continuum of care and to include a more specific focus on family members’ opinions about how they feel about having a role in safeguarding. Investigators have made inferences from one or two safety practices (eg asking about handwashing) and have generalized to how patients feel on all aspects of personal safety. Also, there needs to be clarity, and consistency in meaning and approaches to distinguishing how patients believe they should be, want to be, are able to be and need to be involved in safety. This also includes how they report being and are observed to be involved in safety, and all of which may vary across the continuum of the patient’s condition. Related to this is awareness that beliefs and attitudes may not translate to actions—saying one agrees philosophically with the premise does not mean they will act that behaviour in reality. Conversely, it may be that some consumers do not fundamentally agree that patients should have a role but engage because they have been asked to be part of the process. It is these complexities and nuances that make investigating this topic a difficult one. Overall, based on the review results, while there is the belief and attitude of many patients and family members that patients should and want to participate in partnering to ensure safe care, safety is principally seen as the responsibility of providers.

Third, most of the existing evidence is about hospitalized adult medical-surgical patients, and primary care has not been extensively studied. Further, the patients’ and family members’ role in supporting safe care is different and evolves or changes during the continuum of care (eg from emergency to critical care to a medical unit) and in different settings (ambulatory versus inpatient) and this context is important to understand. Additionally, it is important to ensure the best study methodologies, dependent on the research study question(s), are used to investigate these complex topics, and not approached as an add-on or out of convenience (eg survey)—an observation of some existing literature. It is offered that approaches such as an exploratory sequential mixed-methods design or multi-phased studies may best address this complexity, but integral to this work must be rigorous designs that capture the richness and depth of patient and family insights.

Fourth, the findings of the scoping review support the concept that there are many issues to consider when one examines patient engagement in patient safety at the direct care level. Additionally, these issues can change and evolve over time, and are unique to each patient. The review by Davis and colleagues about factors affecting participation in patient engagement in safety illustrated similar concepts to the qualitative study of this multi-phase research, including patient-related characteristics, illness-related aspects, health-care professional-related characteristics, health-care–related characteristics and task-related characteristics. However, it does not address issues such as patient’s temperament, the choices and decisions they are making about their circumstance or specific situations, or safety behaviours that patients engage in on their own. As well, previous experience with a safety incident did not consistently mean patients were engaged. Some investigators have found inconsistency in demographic characteristics, while others suggest predictable influences.

Finally, investigators have typically examined safety practices that they have identified, without consideration of patient preferences and without using a participatory action research approach. In most cases, the uptake of these behaviours has been variable and lacking consistency. Further, in many cases, an indirect finding has been that patients engage in strategies of their own. A more effective approach when considering patient engagement in patient safety may be to determine patients’ understandings and preferences at the point of contact with the health-care system and strengthen and enhance those self-identified strategies. An efficient, user-friendly mechanism for determining this will be required. Vincent and Spath write of the complexity of involvement and the importance of discerning patient preferences, and it is offered that greater emphasis needs to be taken in this regard to most effectively involve patients and family members in ways that are right for them and in support of their safe care at the direct care level.

6.1 Study strengths and limitations

It is offered that, as a component of a larger investigation, the strength of this study is its depth and breadth, particularly related to the focus on both attitudes and behaviours across the continuum of care, and cross-referenced with findings from the qualitative study phase. To our understanding, no other review has been conducted with this comprehensive approach. It provides perspective internationally about initiatives and efforts that are underway to engage patients in different elements of safety, and illuminated the gaps that remain. It is acknowledged that a study limitation may include the oversight of papers not identified in the applied search strategy.

7 Conclusion

This review was about mapping what is known of patients’ and families’ attitudes regarding their role in safety at the direct care level, as well as their reported behaviours in support of their safe
care. The review included 151 papers, and among the findings included patients’ belief in having a role in safer care (although not for everyone), with indication of the need for further investigation in this regard, as well as degrees of variability in taking action. The review also provided perspective of the rapidly evolving interest in this topic, particularly as it relates to behaviours generally, and more specifically about the involvement of patients in asking providers about handwashing, although involving patients in the research process and specifically in patient-identified engagement safety strategies is needed. We must appreciate that many patients are engaged while recognizing their own and system limitations, and better position ourselves as researchers, policymakers and providers in understanding and implementing approaches collaboratively as relevant and feasible.

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CONFLICT OF INTEREST
None of the authors have any conflict of interest to declare.

DATA AVAILABILITY STATEMENT
Data sharing is not applicable to this article as no new data were created or analysed in this study.

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REFERENCES
1. Panagioti M, Khan K, Keers RN, et al. Prevalence, severity, and nature of preventable patient harm across medical care settings: systematic review and meta-analysis. BMJ. 2019;366:l4185.
2. Gandhi TK, Kaplan GS, Leape L, et al. Transforming concepts in patient safety: a progress report. BMJ Qual Saf. 2018;27(12):1019-1026.
3. Duhn L, Medves J. A 5-facet framework to describe patient engagement in patient safety. Health Expect. 2018;21(6):1122-1133.
4. Carman KL, Dardess P, Maurer M, et al. Patient and family engagement: a framework for understanding the elements and developing interventions and policies. Health Affairs (Project Hope). 2013;32(2):223-231.
5. Duhn L. A comparison of patient safety perspectives of patients, families and healthcare professionals. In: Queen’s University; 2011.
6. Bernstein M, Potvin D, Martin DK. A qualitative study of attitudes toward error in patients facing brain tumour surgery. Can J Neurol Sci. 2004;31(2):208-212.
7. Burroughs TE, Waterman AD, Gallagher TH, et al. Patients’ concerns about medical errors during hospitalization. Jt Comm J Qual Patient Saf. 2007;33(1):5-14.
8. Dowell D, Manwell L, Maguire A, et al. Urban outpatient views on quality and safety in primary care. Longwoods Rev. 2005;3(1):2-8.
9. Elder NC, Jacobson CJ, Zink T, Hasse L. How experiencing preventable medical problems changed patients' interactions with primary health care. Ann Fam Med. 2005;3(6):537-544.
10. Entwistle VA, McCaughan D, Watt IS, et al. Speaking up about safety concerns: multi-setting qualitative study of patients’ views and experiences. Qual Saf Health Care. 2010;19(6):e33.
11. Hupcey JE. Feeling safe: the psychosocial needs of ICU patients. J Nurs Scholarsh. 2000;32(4):361-367.
12. Kuzel AJ, Woolh SF, Gilchrist VJ, et al. Patient reports of preventable problems and harms in primary health care. Ann Fam Med. 2004;2(4):333-340.
13. Mazor KM, Goff SL, Dodd KS, Velten SJ, Walsh KE. Patients’ perceptions of medical errors. J Patient Saf. 2010;6(2):102-107.
14. National Patient Safety Foundation at the AMA. Public Opinion of Patient Safety Issues: Research Findings. In. Boston, MA1997.
15. Rathert C, Brandt J, Williams ES. Putting the ‘patient’ in patient safety: a qualitative study of consumer experiences. Health Expect. 2011;15(3):327-336.
16. Saxton M, Curry MA, Powers L, Maley S, Eckels K, Gross J. “Bring My Scooter So I Can Leave You” A Study of Disabled Women Handling Abuse by Personal Assistance Providers. Violence Against Women. 2001;7(4):393-417.
17. Saxton M, McNeff E, Powers L, Curry MA. We’re all little John Waynes: a study of disabled men’s experience of abuse by personal assistants. J Rehabil. 2006;72(4):3.
18. Schwappach DLB. “Against the silence”: development and first results of a patient survey to assess experiences of safety-related events in hospital. BMC Health Serv Res. 2008;8:59.
19. Schwappach DLB, Wernli M. Am I (un) safe here? Chemotherapy patients’ perspectives towards engaging in their safety. Qual Saf Health Care. 2010;19(5):e9.
20. Vanderheyden L, Northcott H, Adair C, et al. Reports of preventable medical errors from the Alberta Patient Safety Survey 2004. Healthcare Quart. 2005;8(5p):107-114.
21. Weingart SN, Price J, Duncombe D, et al. Patient-reported safety and quality of care in outpatient oncology. Jt Comm J Qual Patient Saf. 2007;33(2):83-94.
22. Wolosin RJ, Vercler L, Matthews JL. Am I safe here?: improving patients’ perceptions of safety in hospitals. J Nurs Care Qual. 2006;21(1):30-40.
23. Pandhi N, Schumacher J, Flynn KE, Smith M. Patients’ perceptions of safety if interpersonal continuity of care were to be disrupted. Health Expect. 2008;11(4):400-408.
24. Rathert C, Huddleston N, Pak Y. Acute care patients discuss the patient role in patient safety. Health Care Manage Rev. 2011;36(2):134-144.
25. Walrath JM, Rose LE. The medication administration process: patients’ perspectives. J Nurs Care Qual. 2008;23(4):345-352.
26. Agoritsas T, Bovier PA, Perneger TV. Patient reports of undesirable events during hospitalization. J Gen Intern Med. 2005;20(10):922-928.
27. Buetow S, Henshaw J, Bryant L, O’Sullivan D. Medication timing errors for Parkinson’s disease: perspectives held by caregivers and people with Parkinson’s in New Zealand. Parkinson’s Dis. 2010;2010:432983.
28. Swahnberg K, Wijima B, Hearn J, Thapar-Björkert S, Berterö C. Mentally pinned: men’s perceptions of being abused in health care. Int J Mens Health. 2009;8(1):60.
29. Davis RE, Koutanji M, Vincent CA. How willing are patients to question healthcare staff on issues related to the quality and safety of their healthcare? An exploratory study. Qual Saf Health Care. 2008;17(2):90-96.
30. Davis RE, Sevdalis N, Vincent CA. Patient involvement in patient safety: How willing are patients to participate? BMJ Qual Saf. 2011;20(1):108-114.
