REVIEW ARTICLE

What is the experience of being readmitted to hospital for people 65 years and over? A review of the literature
Aim: The aim of this review is to explore the experience of a readmission to hospital from the perspective of older adults.

Methods: A systematic review with an interpretative approach was conducted. CINAHL, Embase, and Medline were consulted. The search took place during October 2016.

Results: Six studies with data collection between 2004 and 2013 fit the relevant criteria and included a total of 68 older adults. Two overarching themes were developed with relevant subthemes: experience during initial hospital stay distinguished by exclusion (Feeling powerless; Feeling disregarded; Perception of readiness for discharge) and patients experience uncertainty following discharge (Perception that community-based services are not available or adequate; Perception that hospital is the only safe place; Difficulty in adapting to a “new normal”).

Conclusions: A cycle of exclusion exists during the initial hospital stay and beyond. The experience of being readmitted to hospital is challenging, mostly perceived as negative, and existential, emotional and psychological well-being is not satisfactorily addressed by healthcare professionals.

Keywords: readmission; hospital; perception; feeling; patients; discharge; uncertainty

Impact statement

This review highlights patient voices and illustrates that they experience feelings of exclusion and uncertainty throughout admission and readmission to hospital.

Introduction and background

Current evidence indicates that older adults experience a high rate of unplanned readmission to hospital within 30 days of discharge (NHS England, 2015). Unplanned readmissions are associated with poor outcomes for patients (Walsh, 2014), have an impact on the wider health and social care system, can cause disruption to other forms of care (King’s Fund, 2010) and incur significant financial costs (Conroy & Dowsing, 2012). Hospital readmissions are rising despite efforts to implement preventative services (Walsh, 2014) and the latest published data for England shows readmissions accounted for 12% of hospital admissions in 2011/2012 (Health and Social Care Information Centre [HSCIC], 2015). A variety of factors have been noted to affect readmissions including increasing age (Walsh, 2014) or deprivation (Purdey & Huntley, 2013). Readmissions are unequally distributed across the population and rates are higher among older people (Oliver, 2015). In the UK adults admitted to hospital and over the age of 65 have a 15% readmission rate, a figure that is rising (Oliver, 2015) and higher than the rate for all readmissions at 11% (HSCIC, 2013). There is evidence that this trend is seen internationally (Li, Yong, Hakendorf, Ben-Tovim, & Thompson, 2015).

The vast majority of literature on readmissions relies on routinely collected health data such as readmission rates by age or diagnosis (Horwitz, 2016). Although these factors can be measured statistically, statistics alone do not give much in-depth understanding of this phenomenon. Indeed, there continues to be a real lack of understanding of how older people themselves experience readmissions. Patient experience is considered to be one of three dimensions of quality healthcare (Doyle, Lennox, & Bell, 2013) with positive patient experience associated with better health outcomes (Doyle et al., 2013). Furthermore, healthcare systems such as the National Health Service (NHS) actively include patient voices in their decision-making as they recognise their important value (NHS England, 2015). Despite knowing that patient experience is important, healthcare professionals are not always aware of what matters to patients (Edwards, Duff, & Walker, 2014). Thus, research undertaken from the perspective of the patient provides an enhanced understanding of their experience which may elucidate reasons for readmission and is consistent with providing and delivering quality healthcare. It is also consistent with nursing values (Knisely & Dracuker 2015; McCormack, Karlsson, Dewing, & Lerdal, 2010).

As healthcare professionals, we must listen to the voices of older adults in order to improve patient care (Kings Fund, 2008). Thus, this review will explore unplanned readmissions among older adults from the perspective of the patient using research that has been undertaken from a qualitative perspective.

Method

Aim

The aim of this interpretive review is to examine qualitative research papers to explore the experience of patients (aged 65 and over) who have been readmitted to a general hospital.
Search strategy

Search terms were discussed and confirmed with a specialist librarian; these were: Readmission, Rehospital*, Re-hospital*, Patient readmission, Reattend*, Re-attend*, Hospital readmission AND Experience*, Feel*, Perspective*, Attitude* AND Aged, Aged 80 and over, “older adult”, “old age”, Geriatrics, Geriatric*, Frail elderly, Elder*, “old person”, “old people”, “65 year”, “80 year”, “over 65”, “over 80”, Aged hospital patient, Very elderly. Search terms and relevant thesaurus terms were used in the following databases: CINAHL, Embase and Medline in October 2016. Title and abstract were searched in all databases.

Inclusion criteria required that articles should be written in English language, published between 1996 and 2016 and were qualitative studies focusing on the experience of readmission of people aged 65 and over to a general hospital. Articles were excluded if they assessed an intervention or new care model, included people under the age of 65, focused on readmissions to mental health or oncology services, or solely explored the discharge process from hospital to home.

Titles and abstracts were screened and those that met the inclusion criteria were read in full. Sixty-two full text articles were obtained from which 6 were found to meet all inclusion criteria. A citation search was carried out in Web of Science™ but no further articles were found via this method. The reference lists of the selected articles were searched but no further articles were identified (Figure 1).

Quality appraisal

Appraising qualitative data can be undertaken in a variety of ways and there are no agreed criteria for determining good quality qualitative research. Aveyard, Payne, and Preston (2016) refer to a range of published guidelines which recommend assessing for credibility, resonance, significant contribution, ethics and coherence. The selected articles were assessed for these using the Joanna Briggs checklist (Joanna Briggs Institute, 2016) and were examined to establish if there was congruity between methodology and research question, data collection, data analysis, data interpretation and conclusions.

All of the selected articles had obtained relevant ethical approval, yet some limitations were noted. Only one of the articles
clearly located the researchers culturally and theoretically and addressed the influence of the researcher on the research (Dilworth, Higgins, & Parker, 2012). This lack of theoretical and cultural positioning could have an impact on assumptions made by the researchers thus affecting results. Congruence was noted between findings and conclusions in all selected articles. One article (Stephens et al., 2013) used field notes as opposed to audio-recorded data, but direct quotes from patient participants were included in the article. Yu, Lee, and Woo’s (2007) article is a “research in brief” as opposed to a full article and therefore contains less details.

Analysis

The six articles were analysed in order to capture a rich thematic description of the data. This was done following the methods outlined in Thomas and Harden (2008) and involved line-by-line coding of the data, construction of descriptive and then analytical themes. Each article was read multiple times and reviewed to determine (i) how participants explained their readmission and (ii) what themes exist around the readmission experience. Data were extracted into tables and grouped by initial code words by the first author. These tables were then regularly reviewed and themes were generated and refined using mind-maps over the course of approximately four weeks. These themes were reviewed and agreed upon via group discussion with all the authors.

Findings

Overall, there appears to be a “cycle of exclusion” experienced by many of the 68 participants in these studies. This begins during their initial hospital stay and continues until readmission. This period after their hospital stay is further distinguished by a perception of uncertainty about and exclusion from healthcare system factors.

The included studies were undertaken in Australia, the USA, Canada and Hong Kong. Further details, aims and findings are summarised in Table 1. Three of the articles (Slatyer et al., 2013; Stephens et al., 2013; Vat et al., 2015) incorporate experiences of older adults, caregivers and healthcare professionals. There were distinct sections enabling clear data extraction of the older adult perspective (Table 2).

<table>
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<th>Table 1.</th>
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<td><strong>Aims</strong></td>
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| Dilworth et al. (2012) | The aims of the study were  
• to explore the experience of older people who had been readmitted to hospital following discharge from hospital  
• to investigate the circumstances that lead to their readmission | • Qualitative descriptive design  
• Face-to-face, semi-structured, in-depth interviews  
• Data analysis: inductive approach, coding and theme generation (can I call this thematic analysis?) | 3 older adults, readmitted to hospital within 28 days of discharge from a large tertiary referral hospital in Australia, all English speakers | Australia/hospital | Main themes: being left out, being cared for and feeling let down |
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<tr>
<th>Study</th>
<th>Aims</th>
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<th>Sample</th>
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<td>Slatyer et al. (2013)</td>
<td>The study aimed to describe the factors contributing to early ED re-presentation from the perspectives of (1) patients aged 65 years and older who experienced a presentation to that hospital's AMU; (2) family caregivers who provided care to such patients between the discharge and re-presentation; (3) hospital and community-based health professionals involved in their care</td>
<td>Qualitative descriptive design; Narrative approach; Individual, audio-recorded, semi-structured interviews, conducted face-to-face or by telephone, as preferred, within three weeks of representing; Constant comparison method to descriptive level for data analysis</td>
<td>12 older adults, 6 female and 6 male, who returned to the study hospital's ED within 28 days of discharge. 10 of these participants were interviewed. Additionally, 15 caregivers were included in the study along with 35 health workers</td>
<td>Australia/hospital</td>
<td>Four themes emerged: the health trajectory; communication challenges; discharge readiness and; the decision to return. Re-presentation to hospital was seen as part of a declining health trajectory. Patients and caregivers left hospital with limited understanding of health problems; therefore, they were ill-prepared for future health crises</td>
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<td>Stephens et al. (2013)</td>
<td>To better understand patient and provider perspectives of the transitional care needs and challenges faced by re-hospitalised veterans</td>
<td>Grounded theory; Semi-structured face-to-face interviews; Thematic analysis</td>
<td>25 older veterans (all male?) readmitted within 90 days to medical/surgical units in a Veterans Affairs medical centre in the USA. Average age 68; 32% Black, 4% Hispanic/Latino; 4% Native American; 60% White. 73% have a history of mental health/substance abuse. Average number of readmissions in prior 90-day period: 2.6. Average days inpatient in prior 90-day period: 32. Average length of stay (days): 7.14 healthcare providers were included in this study. Interviews conducted at the bedside in hospital</td>
<td>USA/hospital</td>
<td>Patients identified three themes that led to their rehospitalisation: knowledge gaps and deferred power, difficulties navigating the system, and complex psychiatric and social needs. Healthcare providers identified the following themes: substance use and mental illness, lack of social or financial support and homelessness, premature discharge and poor communication, and non-adherence with follow-up</td>
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| Uscatescu, Turner, and Ezer (2014) | The question guiding the study is:  
- What are the experiences of older adults after discharge from the ED that led to early return visits? | Qualitative descriptive design  
- Semi-structured face-to-face audio-recorded interviews in English or French (all researchers bilingual). In 4 interviews both patient and caregiver present  
- “Member checks” conducted  
- Thematic analysis | 15 English- or French-speaking participants readmitted to a Canadian ED within 14 days were interviewed. Seven women and eight men. Mean age 80. 11 of the 15 participants lived with family members, had high school education or less, and had a family physician | Canada/hospital | Three major themes emerged: “Managing the symptoms”, “Care during the Initial ED Visit” and “Who I Am”. The findings suggest that the main reason for older adults’ return to the ED is the severity of the symptoms they experienced |
| Vat, Common, Laizner, Borduas, and Maheu (2015) | This study aims to:  
- understand patients’ reasons for returning to ED following a hospitalisation on an internal medicine unit  
- Compare these reasons against the risk assessment tools completed by the healthcare professional prior to discharge | Qualitative, descriptive study  
- Individual, semi-structured, audio-recorded interviews in English or French at the hospital  
- Inductive thematic content analysis according to Burnard’s (1991) method of analysing interview transcripts  
- “Member checks” conducted | Eight patients readmitted to the ED of a major teaching hospital in Canada within 14 days of discharge. Six women, two men | Canada/hospital | Patients attributed their return to hospital to being discharged too soon, feeling weak at discharge, having limited help to manage their chronic illness and not having enough discharge instructions. Their reasons for returning were different to the ones predicted by the clinicians who used risk prediction tools |
Yu et al. (2007)
The aim of this study was to investigate the phenomenon of recurrent hospital readmission from the perspective of older Chinese COPD patients.

**Methodology and method**
- Exploratory qualitative design
- In-depth, unstructured, face-to-face, audio taped interviews
- Thematic content analysis

**Sample**
Five participants, all male, aged between 70 and 81 years, living with COPD, with multiple readmissions (readmission number range from 10 to 22) to a regional hospital in Hong Kong during 2004.

**Setting**
Hong Kong/not stated

**Findings**
Main themes: perceived powerless to manage the disease after discharge, lack of confidence in community-based care, tense relationship between caregiver and recipient, satisfaction with social atmosphere in hospital. The findings highlight that the illness behaviour in seeking hospital readmission is framed in the Chinese patients' perceptual, social and cultural schemata.

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Experience during initial hospital stay distinguished by exclusion

Participants felt that they were not involved in decision-making, information was not shared or discussed and they felt excluded from the discharge process. These examples point to a “cycle of exclusion” that begins during the initial hospital visit and continues into the discharge process.

Feeling powerless

Participants noted feeling uninformed about their own care and health and described feeling “powerless, unheard and disrespected” (Dilworth et al., 2012, p. 283). Stephens et al. (2013) report patients deferred power to the healthcare professionals looking after them resulting in a lack of perceived control or ownership over their care. Similar examples are seen in Dilworth et al.’s (2012) study with some participants not questioning medication changes that later affected their health.
and contributed to their readmissions. Furthermore, one participant commented they could not discuss their care with the medical team due to self-perceived lack of education and understanding of medical language used in discussion. Powerlessness was also experienced through not having control over changes to plans and treatments or not receiving adequate information from which to make informed decisions. Patients noted they did not have information about what was happening to them, they had unanswered questions, and treatment changes were not explained or discussed (Dilworth et al., 2012). One participant stated feeling they were in a “no man’s land” where a lack of information was coupled with multiple changes to plans and treatments (Dilworth et al., 2012).

Feeling disregarded
Patients’ knowledge of themselves, their values and preferences were ignored (Dilworth et al., 2012). Participants reported feeling disregarded multiple times during the initial hospital stay and discharge decision-making period (Dilworth et al., 2012). They described communication with staff where their needs and wishes were not acknowledged (Slattery et al., 2013; Stephens et al., 2013; Uscatescu et al., 2014). Some noted they had limited recall of communication with healthcare staff and that the busy, stressful environment affected their understanding of their complex healthcare situations (Slattery et al., 2013). A lack of information sharing and patients receiving mixed messages were also noted (Dilworth et al., 2012). In some instances, different members of the healthcare team (doctors and nurses) provided diverse pieces of information, and sometimes included conflicting advice about treatment, discharge or care plans (Dilworth et al., 2012). Participants noted they did not fully understand their conditions (Slattery et al., 2013), could not recall all their medication or care plan details (Stephens et al., 2013), or left the hospital with no explanation as to the cause of their symptoms (Uscatescu et al., 2014). Being disregarded led to some people feeling frustrated and let down (Dilworth et al., 2012) or harmed (Dilworth et al., 2012; Uscatescu et al., 2014). For example, one participant experienced renal failure when a doctor disregarded their warnings about Non-Steroidal Anti-Inflammatories and prescribed them regardless (Dilworth et al., 2012).

Perception of readiness for discharge
Patients did not feel ready for discharge even when positive about going home. Vat et al. (2015, p. 3609) report participants felt they were rushed out, “discharged too soon”, “definitely too early” or “not prepared to go home”. These participants felt the decision to be discharged should be based on “their level of autonomy and physical capacity” and not on their length of stay (Vat et al., 2015, p. 3609). One participant discussed he knew he was not ready for discharge (Yu et al., 2007). This participant immediately consulted a private doctor on discharge and was readmitted to hospital (Yu et al., 2007). In some instances, participants affirmed their feelings of not being fully recovered or well enough to go home were ignored (Dilworth et al., 2012). Two participants mentioned they had been looking forward to going home despite not having felt ready or healthy enough to go (Yu et al., 2007).

Not feeling ready to go home could be associated with readmissions or the fear of readmission. Vat et al. (2015) noted their participants came back to the hospital as they did not feel ready to go home in the first place. Elsewhere, a participant represented to hospital as they had been sent home without knowing why they were in pain (Uscatescu et al., 2014). Others came back to hospital after experiencing anxiety and worry about having gone home with their condition (Slattery et al., 2013). Furthermore, patients felt their readmission may have been prevented if their clinical condition had been considered as opposed to their length of stay (Yu et al., 2007).

Patients experience uncertainty following discharge
Immediately on discharge, some individuals experienced a feeling of uncertainty. Some felt they could not rely on community support, others had difficulty accessing it. For others the feeling of uncertainty appears linked to the contrast between how they felt in hospital versus home; or their ability to manage living with the effects of a diagnosis or treatment; a “new normal”. Therefore, the following subthemes were developed: Perception that community-based services are not available or adequate; Perception that hospital is the only safe place and Difficulty in adapting to a “new normal”.

Perception that community-based services are not available or adequate
Participants perceived healthcare professionals in the community could not be trusted or relied upon (Uscatescu et al., 2014; Vat et al., 2015) due to both a lack of availability and perceived competence or clinical skill. Unavailability of resources in the community setting was noted when patients could not access community services (Stephens et al., 2013; Vat et al., 2015), some could not obtain a GP or physiotherapy appointment for over two-weeks post-discharge (Dilworth et al., 2012). Several participants stated it took too long to be seen (Uscatescu et al, 2014) or spoke of doctors and community-based nurses who did not show up (Vat et al., 2015). Others were told they would have a home assessment at an unknown point in the future (Dilworth et al., 2012). Some noted that not all community-based resources were available (Vat et al., 2015). Difficulties were also experienced when trying to make appointments or get medication (Stephens et al., 2013). Certain groups of patients, such as homeless or socio-economically disadvantaged individuals, also had difficulty accessing services due to an inability to pay for transport (Stephens et al., 2013). The absence of community-based resources is also illustrated by Uscatescu et al. (2014) who note only one participant in the study was able to access a follow-up appointment, the others were readmitted before they could attend one. Similarly, Slattery et al. (2013) described a patient’s condition deteriorated whilst they had been waiting for a GP appointment and then had to return to hospital.

In addition, community healthcare professionals were not trusted or relied upon due to the perception they could not manage patients’ clinical conditions (Vat et al., 2015; Yu et al., 2007). Some stated their GPs did not understand their condition (Vat et al., 2015) or were not competent to deal with it (Uscatescu et al., 2014) whilst others thought their condition could not be “cured” in the community (Yu et al., 2007).

Perception that hospital is the only safe place
The view that hospital is a safe place in contrast to the uncertainty of home is illustrated with some participants describing doubt about their ability to manage at home. All the participants (n = 3) in Dilworth et al.’s study (2012) described this, feeling their return to hospital was unavoidable and that they needed to return as they required help or felt unwell. Similar situations are described elsewhere (Slattery et al., 2013) with individuals feeling uncertainty because they did not understand the complexity of their care or had a limited understanding of their health which contributed to feeling they could not manage (Slattery et al., 2013). In some instances, participants felt hospital was a positive setting and their only lifesaver, with some noting their only resource in the face of death was to return to hospital describing this as an “obligation with no alternative” (Uscatescu et al., 2014, p. 37). Yu et al. (2007) noted patients felt powerless to manage COPD after discharge and they lacked confidence in their ability to manage symptoms. Some participants reported a feeling of impending death which meant...
they did not feel they had any other option but to return to hospital (Yu et al., 2007). Hospital as a safe social space was experienced by some patients who had formed positive and supportive bonds with healthcare professionals and other patients (Yu et al., 2007). This contrasted with their uncertainty over what services were available in the community and a feeling of being a burden to their families. Participants in this study stated they experienced a positive social atmosphere in hospital and did not feel lonely there (Yu et al., 2007).

Difficult in adapting to a “new normal”

Uncertainty was experienced by patients in terms of expectations of themselves and their ability to adapt to their altered situation as a result of their diagnosis or treatment, which we refer to as a “new normal”. There is evidence that patients are experiencing new uncertainty that they need to learn to live with, but they do not feel they have received adequate support from healthcare professionals to do so. Patients described not feeling back to normal after discharge, or feeling isolated and some experienced acute or unresolved symptoms (Uscatescu et al., 2014). Some patients attributed this to not receiving discharge information or instructions which meant they could not tell if their health was getting better or worse when they went home, they did not know what to expect (Vat et al., 2015). For others this was because they had limited recall of conversations with healthcare professionals (Slatyer et al., 2013), thus they did not have clarity over what to expect on discharge. A lack of confidence about how to control symptoms was also cited by some as a reason for readmission (Uscatescu et al., 2014; Yu et al., 2007). A number of patients even noted they felt anxious and scared (Slatyer et al., 2013).

Other participants contrasted the independence they experienced before their initial admission with how they felt on discharge; they noted this led to feelings of uncertainty and in some cases readmission (Dilworth et al., 2012). This was echoed in other articles, where some patients described they did not want to accept this situation or lose their independence (Slatyer et al., 2013), thus they did not have clarity over what to expect on discharge. This was echoed in other articles, where some patients described they did not want to accept this situation or lose their independence (Slatyer et al., 2013), thus they did not have clarity over what to expect on discharge. For others the “new normal” was yet another issue they needed to adapt to alongside complex life-changing events, such as being a veteran, experiencing homelessness and living with psychiatric conditions (Slatyer et al., 2013).

Discussion

This review highlights the experiences of older adults who are readmitted to hospital. Overall, the themes in this review indicate that returning to hospital can be the consequence of a prior experience in which individuals feel excluded from decision-making, are unprepared for discharge and perceive a lack of support at home which paradoxically can lead to the perception that hospital is the safe place to be. This review illustrates the psychological and emotional experiences that can run alongside the physical health harms, which include readmission, as described by the Kings Fund (2014).

There is evidence in the wider nursing literature that many patients feel ill-prepared to go home (Annema, Luttik, & Jaarsma, 2009; Kangovi et al., 2012). Participants in Annema et al.’s study (2009) did not consider they received adequate help from healthcare professionals with this transition. Not feeling prepared to go home was also found to be associated with readmissions among older adults (Annema et al., 2009; Coffey & McCarthy, 2013). The inevitability of a return has been highlighted elsewhere (Jefferds, Dhall, Cardoso, & Bell, 2014). It therefore seems logical to argue that feeling ill-prepared to go home might increase the likelihood of a readmission.

Effective discharge planning is recognised as a factor in readmissions and patient satisfaction (Gonçalves-Bradley, Lannin, Clemson, Cameron, & Discharge, 2016); however, the strength of this evidence is weak (Gonçalves-Bradley et al., 2016; Preyde, Macaulay, & Dingwall, 2009). Nevertheless, best practice guidance for nurses on discharge planning is available (Lees, 2013; National Institute for Health and Care Excellence, 2015a) and hospitals have local processes and guidelines to support nurses. Evidence from this review through the subthemes of feeling disregarded, powerless and perception of readiness for discharge indicates that discharge planning can impact on readmissions. However, our review also suggests that patients do not feel included throughout the hospital stay; they describe a “cycle of exclusion” that indicates that their care is not being undertaken in a holistic manner.

This review identified that many patients struggle to receive the support they need once at home. Having difficulty obtaining support in the community is also an issue observed in other peer-reviewed (Lees, 2013) and organisational (Age UK, 2012; Healthwatch, 2015; Royal Voluntary Service [RVS], 2013) research. Adequate community support and follow-up is also deemed important by older adults themselves (Healthwatch, 2015; RVS, 2013). It is logical to argue that closer patient involvement in discharge planning might lead to greater awareness and perceived accessibility of community services which may help to support patients once discharged.

Context and power are essential to understanding the findings contained in these studies. For example, some respondents noted they felt powerless to manage their condition (Yu et al., 2007) or that they did not have the knowledge or authority to question the decisions made by healthcare professionals (Dilworth et al., 2012). This indicates opportunities for creating authentic and therapeutic relationships may have been missed. A respondent noted they wanted to be spoken to about their condition using lay language (Dilworth et al., 2012) highlighting that power can be exerted through the language we use as professionals (Kuokkanen & Leino-Kilpi, 2000). This is reflected in other work that describes patients feeling like objects, with discussions controlled by healthcare professionals, use of medical language and communication being “about” instead of “with” them (Efraimsson, Rasmussen, Gillie, & Sandman, 2003).

Power can be viewed in the ability of people to exercise choice (Laverack, 2016), thus having no choice is synonymous with powerlessness. This is pertinent in light of the respondents’ views that they felt powerless or disregarded in decision-making throughout their stay and readmission. As nurses, it is our role to help create therapeutic relationships with patients, creating environments where equality of worth is central and where patients feel able to be active and equal participants (Kuokkanen & Leino-Kilpi, 2000). By understanding all the elements of transition processes, nurses can nurse patients accordingly and take their individual circumstances into account (Meleis et al., 2000). An example of this type of care in practice is delivered by Acute Clinical Team services run in Wales. Skilled Advance Nurse Practitioners assess, diagnose and care for older adults in the community (Griffiths & Davies, 2017). Patients are considered partners in care management, and initial feedback suggests this service prevents hospital admission though more evaluation is required (Griffiths & Davies, 2017).

To be ill has meaning to the person with the illness (Lindburg, Horberg, Persson, & Ekebergh, 2013) and addressing this meaning with patients should form part of routine nursing care. This review demonstrates this psychological and emotional care is often not addressed by healthcare professionals. This is not surprising as these aspects of care have been found to be some of the most commonly reported activities of care left undone with 46% of nurses sampled citing they did not have time for “comfort/talking with patients” and 34% not able to develop or update nursing care plans or care (Ball et al., 2016). Furthermore, other research has found that older adults can rationalise the need for extra support or time in hospital but their existential needs or feelings around grief and loss are not always addressed (Lindburg et al., 2013). Psychological adjustment is also required by patients when they are discharged from hospital but again this element of patient care is often overlooked.
by healthcare providers (Lees, 2013). Helping people to address existential matters and manage and understand changing expectations, of themselves as individuals and of healthcare services, are vital and form part of delivering holistic person-centred care.

One strength of this study lies in the consistent thematic findings despite heterogeneity between studies. The size, composition, geographic and clinical location all vary as do the readmission timeframes which range from 14 days to 1 year. The healthcare systems represented also vary. This review adds strength to the argument around the need to deliver holistic person-centred care and to value the nursing time and skill dedicated to this. It should not be seen as acceptable to rush or omit this aspect of care.

The articles selected for this review each have their own limitations thus these findings should be viewed accordingly. Only articles written in English were included; interviews took place in a hospital setting, not a setting of the participants choosing; there is a lack of noticeable diverse voices (including women, people from different black or minority ethnic groups, LGBTQ and different socio-economic backgrounds); and a lack of data from the UK is apparent. Additionally, only half of the selected articles wholly focused on the patient experience, the others incorporated views from caregivers and health professionals (Slater et al., 2013; Stephens et al., 2013; Vat et al. 2014). Furthermore, a sole researcher carried out the search and analysis which could potentially bias findings.

Impact statement

There are two main outcomes from this review. Firstly, this review highlights the importance of the patients’ voices that are otherwise “silenced” (Serrant-Green, 2011) in current research on readmissions to hospital. Listening to patient voices is vital for healthcare professionals and enables the provision of person-centred care (Schwind, Fredericks, Metersky, & Porzuczek, 2016). Furthermore, viewing patients holistically and considering the way they describe their healthcare trajectories within and outside of hospital settings is important when we are providing care (Schwind et al., 2016).

Secondly, the themes identified in this review are important as they have not been identified in individual qualitative studies. They illustrate that patients often feel excluded and uncertain during readmission, indicating that holistic care is not always being provided. This emphasises the value and importance of human connection in nursing practice. Sharp, McAllister, and Broadbent (2016) have identified that this can improve patients’ experience of care and that incorporating personal, emotional and spiritual elements within nursing can have a positive impact on patient empowerment and recovery. Understanding how and why patients feel excluded and uncertain at readmission is therefore of paramount importance.

Conclusion and recommendations

This review was carried out to explore the experiences of readmission to hospital among older adults. Individual themes in this review are echoed in other research. However, when taken collectively the themes synthesised illustrate a negative experience and a cycle of exclusion for many. In addition, individuals’ existential, emotional and psychological well-being was not adequately addressed. This review highlighted the continued lack of evidence that exists on returning to hospital in the words of older people themselves. Future research should address the continued paucity of data from the perspective of older people and incorporate more diverse voices. Research should also be conducted on how to help manage patients’ and families’ expectations. It would also be interesting to consider how readmission experience could vary depending on the healthcare system in place as this can influence how care is paid for and what services are available. How a lack of resources could impact on healthcare professionals’ abilities to deliver person-centred care must be addressed as part of future research.

The following recommendations are made in light of the findings from this review:

- The nursing skill and time dedicated to delivering holistic person-centred care must be valued by clinical, educational, research and policy organisations and prioritised in clinical settings.
- Future research should include the voices of older people themselves and ensure diversity is represented, for example, through participatory research methods.
- Discharge processes should incorporate adequate psychological and emotional preparation for the transition.
- Information and self-care advice should be given in a way that is relevant to people and the reality of their home life.
- Specialist services that bridge acute and community settings should be further evaluated.

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