What does integrated care mean from an older person’s perspective? A scoping review

Michael T. Lawless, Amy Marshall, Manasi Murthy Mittinty, Gillian Harvey

ABSTRACT

Objective To systematically map and synthesise the literature on older adults’ perceptions and experiences of integrated care.

Setting Various healthcare settings, including primary care, hospitals, allied health practices and emergency departments.

Participants Adults aged ≥60 years.

Interventions Integrated (or similarly coordinated) healthcare.

Primary and secondary outcome measures Using scoping review methodology, four electronic databases (EMBASE, CINAHL, PubMed and ProQuest Dissertation and Theses) and the grey literature (Open Grey and Google Scholar) were searched to identify studies reporting on older adults’ experiences of integrated care. Studies reporting on empirical, interpretive and critical research using any type of methodology were included. Four independent reviewers performed study selection, data extraction and analysis.

Results The initial search retrieved 436 articles, of which 30 were included in this review. Patients expressed a desire for continuity, both in terms of care relationships and management, seamless transitions between care services and/or settings, and coordinated care that delivers quick access, effective treatment, self-care support, respect for patient preferences, and involves carers and families.

Conclusions Participants across the studies desired accessible, efficient and coordinated care that caters to their needs and preferences, while keeping in mind their rights and safety. This review highlights the salience of the relational, informational and organisational aspects of care from an older person’s perspective. Findings are transferable and could be applied in various healthcare settings to derive patient-centred success measures that reflect the aspects of integrated care that are deemed important to older adults and their supporters.

INTRODUCTION

Evidence suggests that many older people are ‘falling through the gaps’ and experiencing fragmented care, particularly when they live with multimorbidity and frailty. In Australia, over 83% of the population aged over 75 years live with two or more chronic conditions and, in the USA, around half of the population aged over 75 years is reported to live with three or more chronic conditions. This group commonly deals with health and functional challenges and reports almost twice as many problems resulting from poorly integrated care compared with those without multimorbidity. This is because they typically see several healthcare providers for different medical conditions, take multiple medications, have numerous agencies involved in providing care and experience a higher incidence of hospitalisation. These circumstances can compromise patient care, further contributing to poorer health outcomes, reduced quality of life and increased healthcare utilisation and costs.

Care integration is proposed as a solution to such fragmentation, with the potential to improve patient experiences while minimising unnecessary use of healthcare resources. Definitions and terminology used...
to describe integrated care differ within the published literature. The WHO defines integrated care, or integrated health services delivery, as:

An approach to strengthen people-centred health systems through the promotion of the comprehensive delivery of quality services across the life-course, designed according to the multidimensional needs of the population and the individual and delivered by a coordinated multidisciplinary team of providers working across settings and levels of care.7 (p 10)

Attempts to improve care integration have been made through numerous policy and research endeavours, yet the extent to which such efforts have achieved wide-scale impact remains questionable. A key message from research to date is that there is no ‘one-size-fits-all’ approach to integrated care.8 Rather, successful design and implementation of integrated care models requires attending to contextual factors, including local enablers and barriers.9 Evidence to date indicates that the most successful implementation efforts are: (1) bottom-up rather than top-down; (2) driven by local need; and (3) have the support and engagement of all key stakeholders, particularly patients and their carers/families.3 This suggests the need to engage older adults, their families and carers, and care providers to achieve the most effective care coordination and integration.

To date, relatively few studies have focused on patients’ perspectives on integrated care, although some research suggests that there may be distinct differences between provider and patient narratives.9 Patients are more likely to emphasise the importance of relational aspects of care and the everyday consequences of living with their condition, as opposed to a clinical focus on managing specific health conditions. This reinforces the importance of understanding patients’ perspectives and views of integrated care, rather than focusing primarily on policy and service-level priorities—a point highlighted in previous research10 and the focus of this scoping review.

To our knowledge, no evidence synthesis has summarised the available literature on older adults’ views and expectations regarding integrated or similarly coordinated care. Starting with a focus on patient experiences rather than single-organisation or single-sector solutions,3 this review forms part of a larger programme of research that aims to coproduce and implement locally relevant approaches to improve integrated care for older adults at risk of repeated hospitalisation guided by a person-centred approach.11 12

Review questions

The review protocol has been published previously12 and sets out a plan to address the following questions:
1. How do older patients define their views and experiences of integrated care?
2. What are the barriers and enablers of quality integrated care from an older person’s perspective?
3. What is the quality of the literature on older patients’ perspectives on integrated care?
4. What are the potential implications for the design and implementation of integrated care programmes for older people?

MATERIALS AND METHODS

Scoping reviews are used to understand the existing breadth of research on a topic, identify gaps in existing literature and assess the need for further investigation.13 14 The scoping review methodology outlined by Arksey and O’Malley15 was employed, details of which are published in our protocol.12 The Preferred Reporting Items for Systematic Reviews and Meta-Analyses extension for scoping reviews checklist13 was used to guide reporting.

Identifying relevant studies

The search strategy aimed to locate published peer-reviewed studies and grey literature reporting on the views of older adults aged ≥60 years (male or female) who had received integrated or similarly coordinated care of any definition in any type of healthcare setting. MMM performed the initial search in four electronic databases (EMBASE, CINAHL, PubMed and ProQuest) and two grey literature databases (Open Grey and Google Scholar). Studies published from June 2008 to July 2019 in English language were included to ensure feasibility and relevance to the current healthcare context, that is, studies conducted after the publication of a consensus definition of integrated care by the WHO.12 No limitations were placed on study design, type of healthcare setting, geographical location, or the upper age and gender of the participants. Literature search strategies were developed using keywords pertinent to older patients and their perspectives. Appropriate variations in spelling and plurals were used in the search (table 1).

Study selection

Studies were selected via a three-step process. First, AM, GH and MMM independently screened titles and abstracts to determine inclusion status. A second screen of full-text articles (AM, ML) ensured that the studies met the inclusion criteria. Third, AM and ML assessed the remaining full-text records for eligibility. A third author (GH) assessed the articles when the other reviewers were uncertain about eligibility status. Disagreements between the reviewers were resolved through group discussion. Finally, we conducted bibliographic searching of the reference lists of the included articles to identify additional potentially relevant studies.

Data extraction

AM, ML and GH used a standardised form to extract relevant data and consulted regularly to help maintain uniformity during the extraction procedure. The following data were extracted: bibliographic information, aim(s) of study, additional research questions/objectives, study design characteristics, participant
characteristics, definition of integrated care provided, outcomes reported, most important findings containing patient voice, other relevant findings, conclusions, study limitations and author recommendations.

Quality appraisal
Although it is not customary to undertake a quality appraisal as part of a scoping review, because the purpose of this review was to inform recommendations for healthcare policy and practice, a formal quality assessment was deemed appropriate. The Joanna Briggs Institute critical appraisal checklists were used to assess the quality of final studies included in the review. We used these checklists due to their brevity, clarity and explicit focus on appraising the voices of participants and their role in generating and/or interpreting the research findings. These tools are designed to assess the methodological quality of studies and determine the extent to which studies have addressed the possibility of bias in study design, conduct or analysis. ML, AM and GH independently assessed the papers; scores were categorised into ‘low’ (1–3/10 for qualitative research; 1–2/8 for quantitative research), ‘average’ (4–7/10; 3–5/8) and ‘high’ (8–10/10; 6–8/8) ranges for each assessment tool. These ranges were used to define an overall quality rating for each article. In line with scoping review methodology and in contrast to systematic review methodology, articles were not excluded on the basis of methodological quality and in contrast to systematic review methodology, articles were included in the final review (figure 1).

RESULTS
Search results
Four hundred and thirty-six articles were retrieved from the initial search and, following the removal of duplicates (n=100), 336 articles were screened for eligibility. Fifty-six articles were selected for full-text review and, of these, 26 articles were included in the review. Reference list searching identified an additional four articles, which are included in the final review (figure 1).

Description of studies
Fifteen studies (50%) used a qualitative methodology, while the remainder used a quantitative methodology (20%, n=635–46), or a mixed methodology (20%, n=641–6). Three records (10%) were reports produced by two non-government organisations in the UK. Most of the studies were conducted in Europe (53%, n=16) or North America (40%, n=12). Sample sizes of patient groups or subgroups (ie, older adults) ranged from 4 to 15617 participants. The average age of participants in the older adult/patient groups ranged from 60 years due to the limited number of eligible studies...
reporting exclusively on older people. Participants had one or more of the following conditions: chronic obstructive pulmonary disease, cancer, coronary heart disease, diabetes, stroke, arthritis and asthma (table 2).

Six articles (20%) provided a definition of integrated care. Although definitions varied, they cited common elements and/or principles including: comprehensive services, coordinated care, patient focus, multidisciplinary and/or interprofessional teamwork, effective information systems, optimised resource use, and appropriate organisational culture and leadership. None of the studies explicitly referenced the WHO definition of integrated care. Reflecting the diffuse research literature on integrated care across academic journals and disciplines and the inconsistent terminology used to describe integrated care programmes, initiatives, settings and/or evaluations, several other concepts and terms were mentioned or defined in the articles in relation to care integration: ‘care coordination’, ‘continuity of care’, ‘collaborative self-management’, ‘person-/patient-centred care’.

Quality of the included articles
The methodological quality of the included studies was rated as ‘average’ (15 studies) to ‘high’ (12 studies); quality appraisal was not performed on the three reports. Common limitations of the qualitative papers were: lack
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<th>Author(s) and publication year</th>
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<th>Participant characteristics</th>
<th>Definition of integrated care used in study</th>
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<tbody>
<tr>
<td>Algilani et al (2017)</td>
<td>An interactive ICT platform for early assessment and management of patient-reported concerns among older adults living in ordinary housing—development and feasibility</td>
<td>Sweden</td>
<td>To develop and test feasibility and acceptability of an interactive ICT platform integrated in a tablet for collecting and managing patient-reported concerns of older adults in home care</td>
<td>Mixed-methods design combining interviews with older adults and healthcare professionals, and logged quantitative data</td>
<td>n=8 Swedish-speaking older adults registered in and receiving assistance and/or regular contact with a nurse in the healthcare system</td>
<td>None provided</td>
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<tr>
<td>Baillie et al (2014)</td>
<td>Care transitions for frail, older people from acute hospital wards within an integrated healthcare system in England: a qualitative case study</td>
<td>UK</td>
<td>To investigate care transitions of frail older people from acute hospital wards to community healthcare or community hospital wards, within a system that had vertically integrated acute hospital and community healthcare services</td>
<td>Multimethod qualitative study including individual interviews and focus groups</td>
<td>n=17 key hospital ward staff (GPs, adult community healthcare lead, district nurses, community physiotherapist, community occupational therapist); n=36 ward staff; n=4 older adults (75% female; age range 78–98 years) undergoing care transitions</td>
<td>'Integration is the combination of methods, processes and models that aim to achieve integrated care, which is an organising principle for care delivery that aims to improve patient care through better coordination'</td>
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<tr>
<td>Bayliss et al (2008)</td>
<td>Processes of care desired by elderly patients with multimorbidities</td>
<td>USA</td>
<td>To explore processes of care desired by elderly patients who have multimorbidities that may present competing demands for patients and providers</td>
<td>Qualitative design using individual interviews</td>
<td>n=26 non-for-profit Health Maintenance Organization members (50% female) aged 65–84 years</td>
<td>None provided</td>
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<tr>
<td>Berendsen et al (2009)</td>
<td>Transition of care: experiences and preferences of patients across the primary/secondary interface—qualitative study</td>
<td>Netherlands</td>
<td>To explore the transition of care at the primary–secondary interface with reference to the impact of patients' ability to make choices about their secondary care providers</td>
<td>Exploratory qualitative design with semistructured focus groups</td>
<td>n=71 Dutch patients who had been referred to a specialist within the past 2 years</td>
<td>None provided</td>
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<tr>
<td>Blom et al (2016)</td>
<td>Effectiveness and cost-effectiveness of a proactive, goal-oriented, integrated care model in general practice for older people. A cluster randomised controlled trial: integrated systematic care for older people—the ISCOPE study</td>
<td>Netherlands</td>
<td>To assess the effectiveness and cost-effectiveness of a simple structural monitoring system to detect the deterioration in somatic, functional, mental or social health of individuals aged 75 and over followed by the execution of a care plan for those people with a combination of somatic, functional, mental and social problems</td>
<td>Observer-blinded cluster randomised controlled trial</td>
<td>n=7285 older adults with complex health problems attending general practices in Leiden, Netherlands</td>
<td>None provided</td>
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<tr>
<td>Burridge et al (2016)</td>
<td>Making sense of change: patients' views of diabetes and GP-led integrated diabetes care</td>
<td>Australia</td>
<td>To investigate patients' perceptions and experiences of type 2 diabetes, self-care and engagement with a GP-led integrated diabetes care model</td>
<td>Qualitative interviews, thematic analysis using normalisation process theory</td>
<td>n=30 older patients with type 2 diabetes</td>
<td>None provided</td>
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<tr>
<td>Cheng (2017)</td>
<td>Emotions, significance and improvement expectations: the personal matter of a patient's hospital stay</td>
<td>Canada</td>
<td>To learn about patients' perspectives of hospital care to gain insights about the specifics of patient-centred care</td>
<td>Qualitative analysis of National Research Corporation Canada adult inpatient survey responses</td>
<td>n=1638 responses from patients and hospital discharges from 22 units</td>
<td>None provided</td>
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<tr>
<td>Cook et al (2017)</td>
<td>Older UK sheltered housing tenants’ perceptions of well-being and their usage of hospital services</td>
<td>UK</td>
<td>To examine sheltered housing tenants’ views of health and well-being, the strategies they adopted to support their well-being and their use of health and social care services through a health needs assessment</td>
<td>Parallel, three-strand mixed-methods approach encompassing tenants’ perceptions of health and well-being, analysis of the service’s health and well-being database and analysis of emergency and elective hospital admissions</td>
<td>n=978 tenants living in sheltered housing</td>
<td>None provided</td>
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<tr>
<td>Cowie et al (2009)</td>
<td>Experience of continuity of care of patients with multiple long-term conditions in England</td>
<td>UK</td>
<td>To examine patients’ experiences of continuity of care in the context of different long-term conditions and models of care, and to explore implications for the future organisation care of long-term conditions</td>
<td>Qualitative design with semistructured interviews</td>
<td>n=33 patients with multiple long-term conditions</td>
<td>Definition of ‘continuity of care’ provided</td>
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<tr>
<td>Derksen et al (2012)</td>
<td>A local consensus process making use of focus groups to enhance the implementation of a national integrated health care standard on obesity care</td>
<td>Netherlands</td>
<td>To understand experiences and expectations of healthcare professionals and patients concerning opportunities and barriers for local overweight/obesity care</td>
<td>Exploratory qualitative study using focus groups and individual interviews</td>
<td>n=24 older adults living independently in Zwolle, Netherlands</td>
<td>‘Integrated care includes prevention, screening, diagnosis, treatment, relapse prevention and long-term care’ no source. ‘The central aim of recent guidelines and integrated health standards is the organisation of patient oriented care and the support of patients’ self management’</td>
</tr>
<tr>
<td>Ebrahimi et al (2017)</td>
<td>Effects of a continuum of care intervention on frail elders’ self-rated health, experiences of security/safety and symptoms: a randomised controlled trial</td>
<td>Sweden</td>
<td>To evaluate the effects of the intervention on self-rated health, experiences of security/safety and symptoms</td>
<td>Non-blinded controlled trial</td>
<td>n=161 frail older adults at high risk of further care consumption</td>
<td>None provided</td>
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<tr>
<td>Freeman and Hughes (2010)</td>
<td>Continuity of care and the patient experience: an inquiry into the quality of general practice in England</td>
<td>UK</td>
<td>To examine continuity of care in general practice, with a particular emphasis on understanding ‘good continuity’ from the patient’s point of view, considering the different types of continuity distinguished by researchers and their relationship to other aspects of quality in primary care, and assessing the state of the art of measuring continuity of care</td>
<td>Mixed-methods design involving a search of published research and other relevant documents, following up leads from key sources and individual interviews with GPs and other members of practice teams</td>
<td>n=3 practice managers, n=8 GPs (including partners and trainees), n=6 receptionists and n=2 nurses</td>
<td>Definition of ‘continuity of care’ provided</td>
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<td>Hepworth et al (2013)</td>
<td>'Working with the team': an exploratory study of improved type 2 diabetes management in a new model of integrated primary/secondary care</td>
<td>Australia</td>
<td>To explore how a new model of integrated primary/secondary care for type 2 diabetes management related to improved diabetes management in a selected group of patients</td>
<td>Qualitative research design with semistructured interviews and critical case sampling</td>
<td>n=10 patients with type 2 diabetes attending the Brisbane South Complex Diabetes Service</td>
<td>None provided</td>
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<tr>
<td>Jackson et al (2012)</td>
<td>Patient journey: implications for improving and integrating care for older adults with chronic obstructive pulmonary disease</td>
<td>Canada</td>
<td>To summarise the experiences of four patients with COPD as they interacted with the healthcare system over a 3-month period following hospital discharge, with a view to informing the development of a more integrated approach to service delivery and improved quality of care</td>
<td>Case study methodology using semistructured interviews and patients’ logs</td>
<td>n=3 older adults with a primary or secondary diagnosis of COPD who were discharged home or to seniors’ housing</td>
<td>‘Principles to achieve a fully integrated health system...include (a) comprehensive services across the care continuum; (b) patient focus; (c) geographic coverage and rostering; (d) standardized care delivery through interprofessional teams; (e) performance management; (f) information systems; (g) organizational culture and leadership; (h) physician integration; (i) governance structure; and (j) financial management’</td>
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<tr>
<td>Jeon et al (2010)</td>
<td>Achieving a balanced life in the face of chronic illness</td>
<td>Australia</td>
<td>To develop an in-depth understanding of the experience of patients and family carers affected by chronic illness that will be the basis on which to propose policy and health system interventions that are patient centred</td>
<td>Qualitative design with semistructured in-depth interviews</td>
<td>n=52 patients (46% female, 67% aged ≥65 years, 21% Indigenous Australians, 21% culturally and linguistically diverse (CALD)) and n=14 carers (93% female, 50% aged ≥65 years, 0% Indigenous, 36% CALD)</td>
<td>None provided</td>
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<tr>
<td>Johnston et al (2009)</td>
<td>Designing and testing a web-based interface for self-monitoring of exercise and symptoms for older adults with chronic obstructive pulmonary disease</td>
<td>USA</td>
<td>To describe our process of developing a set of integrated tools to support collaborative symptom and exercise monitoring for patients with chronic obstructive pulmonary disease (COPD) who may be experiencing breathing difficulties</td>
<td>Mixed-methods four-phase design involving semistructured interviews, a targeted review of publicly available self-monitoring tools, software development and field usability testing</td>
<td>n=14 patients with COPD</td>
<td>None provided</td>
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<td>Jubelt et al (2014)</td>
<td>Patient ratings of case managers in a medical home: associations with patient satisfaction and health care utilization</td>
<td>USA</td>
<td>To measure the association of patient perceptions of patient-centred medical home case manager performance with overall satisfaction and subsequent healthcare utilisation</td>
<td>Retrospective cohort study of patients within an integrated healthcare system</td>
<td>n=1415 patients with clinically complex conditions</td>
<td>None provided</td>
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<tr>
<td>Liss et al (2011)</td>
<td>Patient-reported care coordination: associations with primary care continuity and specialty care use</td>
<td>USA</td>
<td>To investigate the association between care coordination and continuity of primary care and differences in this association by level of specialty care use</td>
<td>Cross-sectional study involving survey information on patient experiences and automated healthcare utilisation data</td>
<td>n=2051 Medicare enrollees with select chronic conditions in an integrated healthcare delivery system in Washington State, USA</td>
<td>Definition of ‘continuity of care’ provided</td>
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<tr>
<td>National Voices (2012)</td>
<td>Principles for integrated care</td>
<td>UK</td>
<td>To describe ‘success’ from the perspective of patients and to discuss measures of success</td>
<td>Report with literature review</td>
<td>NA</td>
<td>Integrated care mentioned but not defined</td>
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<tr>
<td>National Voices (2013)</td>
<td>Integrated care: what do patients, service users and carers want?</td>
<td>UK</td>
<td>To summarise views of patients, service users and carers regarding what they want from integrated care, considering implications for education, training and public health</td>
<td>Report with literature review</td>
<td>NA</td>
<td>Integrated care mentioned but not defined</td>
</tr>
<tr>
<td>Osborn et al (2014)</td>
<td>International survey of older adults finds shortcomings in access, coordination and patient-centred care</td>
<td>Eleven countries, including Australia, USA, UK, Canada, New Zealand</td>
<td>To assess how the health system performs, with a particular focus on access to care, chronic conditions and care coordination, patient engagement, social care needs and end-of-life care planning</td>
<td>Computer-assisted telephone interviews of nationally representative random samples in 11 countries</td>
<td>n=15617 older adults aged ≥65 years contacted by market research firms via mobile and/or landline phone</td>
<td>None provided</td>
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<tr>
<td>Rimmer et al (2015)</td>
<td>The design and initial patient evaluation of an integrated care pathway for faecal incontinence: a qualitative study</td>
<td>UK</td>
<td>To describe a novel integrated care pathway for the management of faecal incontinence (FI) and examine the experiences of patients with FI in relation to this pathway</td>
<td>Focus groups and narrative interviews</td>
<td>n=13 patients with FI</td>
<td>‘Integrated care pathways (ICPs) are multidisciplinary plans that predict the course of events in the treatment of patients with similar problems. The aim of an ICP is to enhance the quality of care by improving patient outcomes, promoting patient safety, increasing patient satisfaction and optimising the use of resources’</td>
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<tr>
<td>Roland et al (2012) [44]</td>
<td>Case management for at-risk elderly patients in the English integrated care pilots: observational study of staff and patient experience and secondary care utilisation</td>
<td>UK</td>
<td>To report the outcome of intensive case management for older adults at risk of emergency hospital admission</td>
<td>Mixed-methods approach with preintervention and postintervention survey questionnaires sent to health and social care staff directly involved or impacted by the intervention; patient questionnaires; analysis of hospital utilisation using existing data, analysed by difference in differences analysis</td>
<td>n=460 patients who were part of the integrated care pilot who received the case management intervention and returned both pre/postquestionnaires</td>
<td>'([Integrated care is intended] to achieve more personal, responsive care and better health outcomes for a local population')</td>
</tr>
<tr>
<td>Ryan et al (2013) [45]</td>
<td>Comparing patient and provider perceptions of home- and community-based services: social network analysis as a service integration metric</td>
<td>Canada</td>
<td>To examine and compare provider and patient perceptions of teamwork and collaboration among the home and community-based care providers in four case studies</td>
<td>Case study design involving social network visualisations</td>
<td>n=4 community-dwelling frail older adults</td>
<td>None provided</td>
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<tr>
<td>Sada et al (2011) [31]</td>
<td>Primary care and communication in shared cancer care: a qualitative study</td>
<td>USA</td>
<td>To explore the perceptions of primary care physicians and oncologists' roles, responsibilities and patterns of communication related to shared cancer care in three integrated health systems that use electronic health records</td>
<td>Qualitative design with semistructured interviews</td>
<td>n=10 male patients with cancer</td>
<td>Definition of ‘shared care’ provided</td>
</tr>
<tr>
<td>Sharma (2014) [40]</td>
<td>Integrated care of the diabetic-oncology patient</td>
<td>USA</td>
<td>To measure the effect on emergency department, observation and hospital admissions for patients with cancer with diabetes who were seen by the Diabetic Oncology Program (DOP). A secondary aim included evaluation of patient satisfaction with care coordination and patient empowerment with diabetes self-management for the patients who were seen by the DOP.</td>
<td>Before and after study involving analysis of claims data of adult patients with cancer with diabetes before and after the DOP</td>
<td>n=98 patients with a diagnosis of cancer, historical or active hyperglycaemia, pre-diabetes, or any type of diabetes, attending hospital-affiliated oncology practices, under active treatment</td>
<td>Definition of ‘care coordination’ provided</td>
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<td>Lawless MT, et al. 2020</td>
<td>Specialised (2020)</td>
<td>UK</td>
<td>To explore the experience and perceptions of patients with cystic fibrosis (CF) and staff regarding palliative care and the acceptability of this as a service early in the patient's disease trajectory</td>
<td>Mixed-methods design involving a focus group, a national survey and patient interviews</td>
<td>n=8 patients with CF</td>
<td>'Integrated care relates to principles for delivery of care that aims to improve the patient's experience through improved coordination of care. Integration is the bringing together of methods, processes and models that help bring this about'</td>
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<tr>
<td>Stevens (2014)</td>
<td>An exploration of early palliative care in adult patients with cystic fibrosis and healthcare professionals</td>
<td>UK</td>
<td>To explore the experience and perceptions of patients with cystic fibrosis (CF) and staff regarding palliative care and the acceptability of this as a service early in the patient's disease trajectory</td>
<td>Mixed-methods design involving a focus group, a national survey and patient interviews</td>
<td>n=8 patients with CF</td>
<td>'Integrated care relates to principles for delivery of care that aims to improve the patient's experience through improved coordination of care. Integration is the bringing together of methods, processes and models that help bring this about'</td>
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<tr>
<td>Toscan et al. (2012)</td>
<td>Integrated transitional care: patient, informal caregiver and health care provider perspectives on care transitions for older persons with hip fracture</td>
<td>Canada</td>
<td>To determine the core factors related to poorly integrated care when patients with hip fracture transition between care settings</td>
<td>Qualitative focused ethnographic study using individual interviews and repeated observations</td>
<td>n=6 patients aged ≥65 years with hip fracture with no cognitive impairment able to read and write in English; n=6 informal carers; n=18 healthcare providers involved in the admission or discharge of the patient</td>
<td>Definition of 'continuity of care' provided</td>
</tr>
<tr>
<td>Vat et al. (2015)</td>
<td>Reasons for returning to the emergency department following discharge from an internal medicine unit: perspectives of patients and the liaison nurse clinician</td>
<td>Canada</td>
<td>To understand patients' reasons for returning to emergency department following hospitalisation</td>
<td>Qualitative descriptive approach with in-depth individual interviews</td>
<td>n=8 older patients with chronic illnesses</td>
<td>None provided</td>
</tr>
<tr>
<td>Wodskou et al. (2014)</td>
<td>A qualitative study of integrated care from the perspectives of patients with chronic obstructive pulmonary disease and their relatives</td>
<td>Denmark</td>
<td>To examine how patients with COPD and their relatives experience integrated care across care settings after the implementation of a COPD disease management programme</td>
<td>Qualitative design with focus groups and semistructured interviews</td>
<td>n=34 patients with COPD</td>
<td>None provided</td>
</tr>
</tbody>
</table>

GP, general practitioner; ICT, Information and Communication Technology; NA, not applicable.
of acknowledgement of the influence of the researcher on the research or vice versa, lack of a statement about the cultural or theoretical position of the researcher, and a lack of congruity between the stated philosophical or theoretical perspective and the research methodology.

For quantitative studies, the most common limitation was inadequate consideration of confounding factors (100%).

**Synthesis of findings**

Five themes were identified reflecting older adults’ experiences and views of integrated care: (1) access and availability; (2) involvement, initiative and follow-up; (3) communication and information; (4) referral and care transitions; and (5) coordination and cooperation.

**Access and availability**

Participants in 10 studies identified the ability to access healthcare providers and services as critical. Access was generally discussed in relation to: physical accessibility, convenient access to a known and trusted professional when needed, and access difficulties due to service eligibility restrictions, lack of formal home support and/or the unavailability of needed services or supports in different geographical locations. Three papers discussed difficulties associated with the physical accessibility of services. Problems related to physical access caused significant anxiety for participants, especially when these issues were compounded by unexpectedly long waiting times.

Access was discussed in relation to the continuity of relationships in 10 articles. Participants generally spoke of relationship continuity in terms of the establishment and maintenance of relationships between patients, their carers/families and a known and trusted health or social care professional, as well as positive health and psychosocial impacts of such relationships.

One report stated that patients ‘have clear preferences’ regarding seeing a familiar healthcare professional, such as a general practitioner (GP), giving this preference greater priority still when problems are chronic or distressing. Although patients valued having a long-standing relationship with a single provider, they were prepared to forgo seeing a familiar provider in favour of quick access.

Nine studies discussed the applicability of technologies as part of integrated care approaches for older populations. Participants viewed technology as having an important role, particularly when they experienced difficulty in accessing services, when care plans became increasingly complex and/or were updated repeatedly, and when additional information was needed urgently. Three studies described technical and experiential factors associated with the successful implementation of technologies. Participants in these studies identified user-friendliness as key factors influencing their views on specific technologies.

**Involvement, initiative and follow-up**

Participants wanted to be involved in decisions about their care and treatment in accordance with their needs, preferences and capacities at the time of the encounter. Although participants generally expected to be involved in decisions regarding their care, treatment and medicines, they often felt that care was not ‘centred’ on them. Some participants did not want, or were unable, to make their own healthcare decisions, stating that they preferred to leave decisions up to their family doctor, particularly those related to referral.

Others expressed that although they preferred their family doctor to consult with them, they ultimately wanted to be kept informed and given the opportunity to make their own decisions with the support of health professionals.

Participants felt that there was less scope for them to make decisions when consulting with specialists or when planning for discharge from hospital.

The importance of initiative and follow-up was discussed in seven articles. Participants expected providers to demonstrate initiative by being knowledgeable about their condition and the patient ‘as a person’, considering the applicability of diagnostic investigation, regularly reviewing patients’ care and treatment and opening up discussions about referral and/or patients’ home care needs.

Participants expected providers to take responsibility for following up on previously initiated actions. Indicators of lack of follow-up included missing test results, sudden termination of home care and support without a needs reassessment and serious health conditions remaining untreated for a significant period of time following initial diagnosis.

Participants’ views on the involvement and needs of carers and families varied. Participants appreciated when carers accompanied them to appointments with healthcare professionals because it assisted them with comprehension, remembering care instructions, scheduling future appointments and providing personal health information when necessary. This was sometimes difficult, however, as carers were generally ‘further removed’ from the dissemination of information. As a result, confusion often existed between professionals and carers about individual roles and responsibilities, resulting in ‘blurred boundaries’ and ambiguity in information sharing and flow. Participants considered it important to attend to carers’ informational and emotional needs, for example, by providing them with instructions about disease management and assessing their stress tolerance.

**Communication and information**

Older adults expected highly developed communication skills and clear, comprehensive information from all providers with whom they interacted regardless of their condition(s), the care setting or the provider’s qualification.
Participants expected providers to demonstrate these traits/competencies by: listening carefully to patients’ perspectives and preferences; informing patients of the relative advantages and disadvantages of referrals and treatments; providing personalised care (ie, not treating patients 'like a number'46 49); using appropriate and accessible language; taking patients’ concerns seriously; and responding empathetically to patients’ emotions. These actions were reported to enable open communication and shared decision-making. Some participants preferred to receive information with and correct any misinformation. Participants appreciated when providers informed them of their hospital treatment and be informed in advance where they were going and the name of their primary contact person; assured that information was shared in a consistent manner when information was provided at important junctures in their care journey, and was followed up promptly and consistently. Having a care plan on record, and having knowledge of its contents and updates, was associated with more positive patient experiences, particularly in terms of feeling involved in healthcare decision-making and care processes. Pati\ents disliked having to ‘repeat their story’ to multiple providers—this was cited as a main reason for wanting to see a familiar healthcare provider. Participants appreciated when providers informed them of other available services to which they might be entitled, as well as information about how to manage financially. Participants additionally desired information that could help them comprehend and prepare for the impacts of their health conditions on other aspects of their life.

Participants’ preferences regarding sources of information were inconsistent. Some participants preferred contact with specialists due to their expertise and prior negative experiences of family doctors providing insufficient information to address their health concerns. A majority of participants, however, preferred to receive information from a primary care provider, such as a GP, in the first instance. Nurses were identified as a valued information source due to their perceived capacity to provide extensive and comprehensible information. Supplementary (usually written) information was appreciated, and was seen as particularly useful when the presence of multimorbidity increased the complexity of developing, understanding and executing care plans. Participants felt it was important to be able to view their health records at any time in order to determine who to share this information with and correct any misinformation.

Problems related to information sharing and transfer were reported in seven articles. Participants felt that different providers across the primary–secondary care interface often had conflicting information about, and opinions of, their care. Missing or conflicting information caused a great deal of uncertainty and confusion for patients and their caregivers. Relatedly, incomplete transfer and availability of relevant information to other healthcare providers was identified as a common inter-organisational and intraorganisational barrier that could lead to fragmented care, confusion or dissatisfaction. In hospital, some participants experienced conflicting information about discharge and were unaware which ward staff (if any) were planning their discharge, sometimes due to the absence of a written discharge plan.

Referral and care transitions

Transitions between services and care settings were generally seen as significant points at which older patients were particularly susceptible to lapses or losses of continuity. Participants expressed satisfaction with the information they received from providers, particularly when information was provided at important junctures in their care journey, and was followed up promptly and consistently. Having a care plan on record, and having knowledge of its contents and updates, was associated with more positive patient experiences, particularly in terms of feeling involved in healthcare decision-making and care processes. Patients disliked having to ‘repeat their story’ to multiple providers—this was cited as a main reason for wanting to see a familiar healthcare provider. Participants appreciated when providers informed them of other available services to which they might be entitled, as well as information about how to manage financially. Participants additionally desired information that could help them comprehend and prepare for the impacts of their health conditions on other aspects of their life.

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the hospital setting or when waiting for diagnostic investigation,\textsuperscript{20} \textsuperscript{22} \textsuperscript{49} but expressed concern about ‘being rushed’, particularly during periods of recovery or when facing significant decisions about their care and/or treatment.\textsuperscript{22} Participants from two studies recounted negative experiences of poor care coordination between providers and services, with some explaining that they had waited up to a year before seeing a specialist following referral from their GP.\textsuperscript{25} \textsuperscript{34} Lengthy wait periods caused feelings of uncertainty, frustration and abandonment, particularly if symptoms were perceived as worsening and/or beyond patients’ self-management abilities.\textsuperscript{21} \textsuperscript{22} 

**Coordination and cooperation**

Twelve studies discussed the importance of cooperation, coordination and communication across organisations, between service providers and across types of care.\textsuperscript{20} \textsuperscript{22} \textsuperscript{25} \textsuperscript{27} \textsuperscript{30} \textsuperscript{34} \textsuperscript{38} \textsuperscript{45} \textsuperscript{46} \textsuperscript{48} \textsuperscript{49} Although participants generally had minimal interest in institutional/organisational priorities regarding integrated care, they wanted professionals and services to ‘work together as a team around the patient’.\textsuperscript{49} Divisions between primary, secondary and community care were regarded as relatively meaningless compared with patients’ overarching desire for high-quality care and continuity, regardless of the source and/or setting.\textsuperscript{49} Participants were generally satisfied with their care when: they were referred to services without difficulty\textsuperscript{34}; interprofessional communication was perceptible and shared with the patient,\textsuperscript{49} and discharge processes were perceived as consultative and coordinated.\textsuperscript{20} However, the fragmented nature of the healthcare system became apparent to patients when they needed to contact multiple providers to coordinate care, errors occurred between care transitions, or when providers disagreed on necessary care and services.\textsuperscript{34} Coordination and cooperation was also discussed in relation to resources, patient rights and entitlements regarding support and financing across organisations and care settings.\textsuperscript{19}

In cases where little teamwork was perceived, participants tended to experience inappropriate, inefficient or inconsistent referrals, reduced motivation to comply with treatment and ineffective or inappropriate responses to emergent needs or unanticipated problems.\textsuperscript{25} \textsuperscript{26} \textsuperscript{34} \textsuperscript{45} Participants viewed serious communication breakdowns (eg, failure to record drug allergies) as unacceptable and damaging to their health.\textsuperscript{25} Patients living with severe illness and/or those with multiple health problems were seen as suffering most from insufficient interprofessional cooperation.\textsuperscript{34}

Four studies identified a possible tension between increased specialty care use and primary care providers’ capacity to coordinate care.\textsuperscript{22} \textsuperscript{28} \textsuperscript{34} \textsuperscript{38} Some participants recounted that although communication between their primary care provider and specialist was evident around the time of referral and the initial specialist appointment, they were unsure whether ongoing communication and follow-up was occurring.\textsuperscript{22} \textsuperscript{28} Participants perceived that visits to specialist care added further complexity to the care delivery process and presented opportunities for gaps in care coordination to occur.\textsuperscript{22}

The appointment of a care coordinator, also referred to as a case manager, was proposed as a solution to poor access, follow-up and coordination between organisations, sites, or providers in six studies.\textsuperscript{34} \textsuperscript{36} \textsuperscript{37} \textsuperscript{44} \textsuperscript{47} \textsuperscript{49} Functions of care coordinators described by participants included: acting as a primary contact person and ‘main person’ responsible for patients’ care, particularly aged and frail patients with serious and chronic conditions; coordinating care and social services across different agencies within the healthcare system; regularly contacting patients and their families; offering guidance about symptom management, and providing assistance with instrumental activities of daily life.\textsuperscript{34} \textsuperscript{47} \textsuperscript{49}

**Discussion**

This review highlights that older adults typically define their perspectives towards integrated care with respect to the relational, informational and organisational aspects of care. These aspects of care and types of continuity were considered to be important drivers and benchmarks of person-centred integrated care, and were central features of patients’ diverse narratives. Our findings concur with previous research exploring the semantic misalignments between patient and medical narratives and understandings of person-centred care coordination.\textsuperscript{9} \textsuperscript{51}

**Enablers and barriers from the perspective of older patients**

Based on our synthesis of patient narratives, several enablers and barriers were identified. Key enablers included: access arrangements that reflect patient needs and preferences regarding which services to access, the speed of access and the methods of access,\textsuperscript{21} \textsuperscript{25} \textsuperscript{27} \textsuperscript{29} \textsuperscript{34} \textsuperscript{41} \textsuperscript{44} \textsuperscript{49} appropriate user-friendly technologies,\textsuperscript{28} \textsuperscript{41} clear communication coupled with appropriate information,\textsuperscript{22} \textsuperscript{24} \textsuperscript{26} \textsuperscript{28} \textsuperscript{30} \textsuperscript{34} \textsuperscript{54} regular contact with a familiar, trusted healthcare provider,\textsuperscript{22} \textsuperscript{32} individualised care planning with appropriate patient involvement in healthcare decision-making\textsuperscript{28} \textsuperscript{34} and systems to reduce gaps in information and to enable regular follow-up,\textsuperscript{22} \textsuperscript{28} \textsuperscript{34} \textsuperscript{36}

Barriers to integrated care as defined by older adults included: unavailability of needed providers and services in certain areas/jurisdictions,\textsuperscript{28} \textsuperscript{34} \textsuperscript{45} lack of opportunities to clarify patients’ needs, priorities and preferences,\textsuperscript{22} \textsuperscript{27} \textsuperscript{34} including those related to patients’ information and/or communication,\textsuperscript{22} \textsuperscript{25} \textsuperscript{28} \textsuperscript{34} \textsuperscript{47} \textsuperscript{49} conflicting information, clinical advice, treatments and/or management,\textsuperscript{22} \textsuperscript{25} \textsuperscript{34} \textsuperscript{48} \textsuperscript{49} lengthy wait times,\textsuperscript{20} \textsuperscript{22} \textsuperscript{25} \textsuperscript{33} \textsuperscript{34} \textsuperscript{47} \textsuperscript{49} limited interprofessional or multidisciplinary teamwork,\textsuperscript{25} \textsuperscript{26} \textsuperscript{34} \textsuperscript{45} \textsuperscript{47} \textsuperscript{49} and relational and informational discontinuity at the primary-secondary care interface.\textsuperscript{20} \textsuperscript{22} \textsuperscript{28} \textsuperscript{32} \textsuperscript{34} \textsuperscript{39} \textsuperscript{39} \textsuperscript{47} \textsuperscript{49}

Defining the meaning of integrated care from the perspectives of different stakeholders, and reconciling those perspectives, poses considerable policy and research challenges.\textsuperscript{92} Structural and organisational-based
definitions are well founded insofar as they are driven by the need to overcome sectoral fragmentation in healthcare systems. However, such concerns are likely to be less important to patients than their smooth, seamless and supported journey through the care system. Organisationally based definitions can overshadow person-centred understandings of care integration, which are needed to guide the delivery of appropriate, coordinated and responsive care. Person-centred definitions based on patient experiences capture a fundamental principal of integrated care, and offer a cogent logic as to its key objectives and success measures.

Implications for implementation and knowledge translation

Our findings suggest a need to better prepare and support providers across the healthcare system to deliver coordinated, efficient and appropriate person-centred care within a complex milieu. The successful implementation of integrated care into practice requires providers to internalise the value of the principles of integrated care while being open to new ways of working within non-traditional models of care delivery. Implementing sustainable integrated care systems for older adults will require multi-pronged, transformative action at the clinical (e.g., facilitating shared decision-making and goal setting) service (e.g., supporting the coordination of services delivered by multidisciplinary providers) and healthcare system (e.g., strengthening governance, accountability systems and financing mechanisms enabling equitable access to services) levels. These actions should coincide with efforts to clarify or respecify discrete tasks, roles, and responsibilities, decision-making processes, and relevant clinical and patient-reported outcome and experience measures.

Care coordination with case management was identified as a critical factor in facilitating communication among providers, assisting patients to implement their care plans and enhancing access across different parts of the healthcare system. Further research into the precise role and value of care coordinators and facilitating technologies, among other elements, in integrated care models is needed.

Limitations

This is the first review to provide a comprehensive synthesis and a quality appraisal of the literature on older adults’ perspectives in relation to integrated care. The limitations of this review mainly relate to: (1) the lack of consistency in concepts, definitions and terminology used to describe ‘integrated care’—a majority of the studies did not provide a definition of integrated care and/or used conceptually similar terms pertinent to integrated care without offering a definition or rationale; and (2) the lack of a comparison of subgroups of older patients. We acknowledge that other terms pertaining to integrated care concepts and/or activities could have been included in the search strategy. However, due to the complexity of the field and the varying definitions of integrated care used in the literature, our initial strategy aimed to identify citations focusing explicitly on older patients’ experiences of their care journey within various integrated care models.

It is possible that the inclusiveness of this review, which included studies undertaken on different client groups and service areas within various healthcare settings, may have compromised the specificity of the findings. Our intention was not to address a specific question with narrow parameters about patients’ experiences to retrospectively derive new models of care that fit with the WHO definition. Nor was our intention to refer to older persons’ perceptions and experiences to assess the effectiveness of different integration models from an organisational health system perspective. Such an approach would be inappropriate as most aspects of integrated care management and governance occur ‘behind the scenes’ and are generally ‘invisible’ to patients. As we have seen, patient narratives are more likely to reflect the visible and tangible aspects of service delivery, and are mainly limited to concerns regarding quality and safety, relationships, and (dysfunctional) coordination. Rather, we aimed to synthesise the available literature on patient experiences as an informed stating point to inform the design, implementation and evaluation of locally relevant approaches that are: (1) underpinned by a patient-centred and system-wide view of care integration; and (2) informed by the journeys of older patients as they move between providers and services and across organisational and/or programme boundaries. Given the need for healthcare systems to embrace the flexibility, contingency and complexity that characterises integrated care, a broad and inclusive approach to understanding patients’ views and experiences is justified. Recognising that a ‘one-size-fits-all’ approach is unlikely to be appropriate when it comes to integrated care delivery, future studies could consider coproduction solutions and the application of interpretative approaches to examine what strategies work for which group of patients and under what circumstances, that is, identifying whom, when, how and why.

CONCLUSION

This review highlights that older adults define their experiences of integrated care in relation to: accessibility—timely access to needed services, age-friendly infrastructure, equitable financing and accessible information; care—feeling respected, heard, involved, informed and cared for; and coordination—uninterrupted care delivered smoothly across settings and services, with clear roles, responsibilities and points of contact. These patient-reported concerns are not adequately represented in current operational definitions that focus primarily on integrated care from an organisational and management perspective. The review draws attention to the humanistic and experiential nature of integrated care experiences and suggests that different patient-centric indices may be needed to assess the quality of integrated

care and to measure the key outcomes of importance to older patients and their carers. Future work on this topic is warranted and aligns with contemporary research and policy efforts focusing on developing integrated care programmes that improve patient care experiences while reconciling the inherent complexities and tensions involved.

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Data are available upon reasonable request. The data analysis plan outlined in the study protocol will be made available by the corresponding author immediately following publication.

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