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Patient experiences of integrated care within the United Kingdom: A systematic review

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Lea Davidson (1), Jason Scott (1) and Natalie Forster

Abstract

Introduction: Integrated care and patient experience are central to the coordination and delivery of high quality health and social care in the UK, but their joint application is poorly understood. This systematic review aimed to gain an understanding of patient experience within current integrated care services in the UK, and specifically, whether they reflect person-centred coordinated care (PCCC).

Methods: Following PRISMA, electronic databases (ProQuest, EBSCO and Cochrane Library) were searched from 2012 to 2019 for primary, peer-reviewed literature. Papers were included where patients' or carers' experiences of integrated care were reported. Papers were excluded where they focused on acute integrated care interventions, measured experience via satisfaction scores only, or findings lacked sufficient depth to answer the research question. Quality was assessed using Mixed Methods Appraisal Tool, and findings synthesised using a framework approach, incorporating the Rainbow Model of Integrated Care and Measuring Integrated Care Patient Framework.

Results: Sixteen studies were included. Person-centred and shared responsibility experiences were most often discussed. Experiences were not always described as positive and some patients experienced a lack of PCCC. Clinical, professional/organisational and functional integration processes were associated with experiencing domains of PCCC. **Discussion:** People with complex needs experience a lack of coordination across teams and wider community resources, and limited associations were made between integration processes and patient experience. Further research which gives context to individual experience, provides greater detail of integration processes and utilises validated patient experience measures of PCCC is required to understand the association between integration processes and domains of PCCC.

Keywords

Integrated health care system, healthcare delivery, United Kingdom, literature review

Introduction

People in developed countries increasingly have multiple chronic conditions and complex care needs, partly due to an ageing population,¹ which has led to increasingly specialised care that results in increased fragmentation of care.² This fragmentation occurs within and between providers, as well as at the interface between primary and secondary care or health and social care.³ There is therefore continued emphasis internationally upon improving quality of care through improved coordination and integration of services.⁴ Despite the benefits of integration often being cited as supporting the transition of care from hospital to community settings, improving self-care, reducing costs and reducing demand for acute care,⁵ there has been limited evidence

of its impact on patient outcomes within previous initiatives. 6

The processes of integration have been organised into the level of system at which they operate within the Rainbow Model of Integrated Care, which highlights person-centred care as a central component.⁷ Person-centeredness is a multifaceted concept, with three core values; considering the person's needs,

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Lea Davidson, Northumbria University, Faculty of Health and Life Sciences, Coach Lane Campus, Newcastle NE7 7XA, UK. Email: lea.davidson@nhs.net wants, perspectives and individual experiences; offering people the opportunity to participate in their care; enhancing the partnership and understanding within relationships with health professionals.⁸ However, it has been argued that these domains can sometimes lead to a tendency to define person-centred care through the behaviour of health professionals, such as care planning and shared decision-making skills,⁹ rather than through the lens of the patient and their experiences.

Amid increasing acknowledgement of the need to measure patient experience as a quality outcome,¹⁰ experience continues to be under-reported and poorly understood.¹¹ Patient experience data is not only an important measure of overall system performance,¹² it is also essential to improving quality, innovation in health care delivery redesign, and improving accountability.¹³ However, there is continued debate regarding the effective measurement of patient experience¹⁴ and the continued use of locally produced measures without proven validity or reliability.¹⁵ Measurement of patient experience of integrated care is also limited, with systematic reviews identifying that the potential effects of integrated care on patient and carer experiences are under-researched in UK literature¹¹ and internationally.¹⁶

The aim of this systematic review is to develop an understanding of current integrated care models in the UK and determine whether evaluations of patient experience reflect a person-centred coordinated care approach.

Methods

This systematic review adhered to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines for conduct and reporting.¹⁷ As the term integration is used broadly within health and social care, careful consideration was given as to whether studies reported a change in the coordination or integration of service delivery. This could include changes within a single service or multiple services. Detailed consideration was also given to whether the studies did in fact report on the experiences of patients and carers (i.e. gave a thick and comprehensive description of the receipt of integrated care), as opposed to more one-dimensional assessments of the satisfaction or preferences of patients and carers alone.

Search strategy

Searches were conducted between June and July 2019 in ProQuest, EBSCO, and the Cochrane Library. Search terms represent the key concepts of the review and the most common indexing terms found during an initial scoping of the literature. Satisfaction was included as a search term since it may be used interchangeably with experience. The following terms were used to search within each database;

1. experience* OR satisfaction OR perce* OR value* OR perspective* OR view* OR feedback (ABSTRACT)

AND

2. patient* OR client* OR user* OR consumer* OR carer* OR men OR women (ABSTRACT)

AND

 integrat* OR coordinat* OR co-ordinat* OR collabor* OR continui* OR joint OR multidisciplinary OR partner* OR "single point of access" (ABSTRACT)

AND

4. health OR social OR system* OR care OR team* OR service* OR housing (ABSTRACT)

AND

 "United Kingdom" OR "UK" OR England OR Scotland OR Wales OR "Northern Ireland" OR "Great Britain" (ANYWHERE)

Inclusion/exclusion criteria and study selection

Inclusion criteria were:

- Studies of all patients/clients/users and their carers (male and female) of health and social care services of any age that included patient or carer reported experience after the introduction of the Health and Social Care Act.¹⁸
- Studies that focused on integrated care (defined as changes in integration/coordination of service delivery in order to improve patient outcomes and experience) involving health, social and/or third sector organisations within the UK.
- Any study design, English language, empirical studies published in peer reviewed journals.

Exclusion criteria were:

- Integrated care which focused on acute care episodes only, e.g. emergency care models or surgical care pathways.
- Experience consisted of satisfaction score only.

• Studies which did not address the quality issues specific to this review with sufficient depth. Therefore, were unable to add value to the findings.

All articles identified in the database searches were downloaded into a bibliographical software package (EndNote X8, *Clarivate Analytics*) to facilitate the study selection process. Duplicates were deleted before articles were sifted for eligibility based on title and abstract by LD. The full text of any articles which could not be included or excluded by abstract alone were then reviewed by LD. All articles identified as eligible for inclusion, and 20% (n=23) of articles excluded on full text were reviewed by JS and NF, with discussion amongst all authors until agreement was reached. No reversal of decisions was required.

Data extraction

Data were extracted by LD using a data extraction form, which included country within United Kingdom, study design, aim and description of the intervention, target population, sampling process, participant characteristics, roles of involved providers, data collection and analysis (including validity/reliability/ethical issues), and summary of results. Separate forms were used to extract data on the integration processes involved and descriptions of patient centred coordinated care.

Quality assessment

The Mixed Methods Appraisal Tool (MMAT)¹⁹ was used to appraise the overall methodological quality of included studies. Along with reliability and validity, careful consideration was given as to how fit for purpose and relevant the studies were in addressing the review questions. Studies which did not meet a minimum score of 10 out of 12 were deemed to lack sufficient depth and relevance, and therefore excluded.

Data synthesis

Integrated care is complex, multicomponent and multilevel, involving different groups and organisations which results in varied and multiple outcomes.²⁰ The number of components within integrated care make linking them with emergent system properties such as patient-centred coordinated care very challenging.²¹ Therefore within this review, integrated care was viewed as a complex adaptive system; a living system of interacting parts capable of changing context for others through their actions.²² A framework synthesis approach was used due to the exploratory nature of the review. The Rainbow Model of Integrated Care and subsequent taxonomy^{7,23} were used to code the processes of integration data within the studies, whilst the Measuring Integrated Patient Care Framework²⁴ was selected to code data relating to patient experience. Table 1 provides an overview of the key concepts of the models used in the study.

Results

Sixteen studies were found to meet both the inclusion and quality appraisal criteria. The selection process is outlined in the PRISMA flowchart (Figure 1). Studies included within this synthesis are summarised in Table 2.

Study characteristics

Of the 16 studies included in the review, 11 were based in England,^{25–35} three in Scotland^{36–38} and one in Wales.³⁹ One study did not name the sites within the UK.⁴⁰ Only the English studies included services which had integrated across health and social care or between acute and primary care services. The Scottish and Welsh studies were focused on single conditions with limited or no organisational integration, whether formal or informal.⁵ Twelve studies took a qualitative approach,^{25,27,28,31–39} three studies took a mixed methods approach^{26,29,40} and one took a quantitative approach.³⁰

The focus of the studies reflected the potential of integrated care to address the needs of those with comorbidities.⁴¹ Seven studies looked at services which addressed the needs of those with, or caring for those at risk of complex co-morbid long-term conditions.^{25,27,29,31–33,35} One study involved a service providing both a disease-focused approach (Type-2 diabetes) and those at risk of complex co-morbidities,³⁰ which was also the only study from an Integrated Care Pilot site. The remaining eight studies focused on services for people with a single condition such as a mental illness,^{26,28,34,39} pregnancy,⁴⁰ chronic heart failure³⁶ and cancer.^{37,38}

Integration processes

Vertical and horizontal integration. Three studies described full vertical and horizontal integration across the micro, meso, macro and supportive levels,^{27,32,35} whilst three others described horizontal integration processes across all levels.^{29,31,38} All six studies involved the integration of health, social care and/or wider community services, reflecting the complex needs of the populations targeted within the studies.

Clinical integration (micro level). All the studies described clinical integration processes (micro level), with having a named key worker, care coordination and

| Model | Component/construct | Description |
|---|---|---|
| Rainbow Model of Integrated Care ⁷ | Clinical integration (micro) | Coordination of person-focused care in a single process across time, place and discipline |
| - | Service/professional integration (meso) | Inter-professional partnerships based on shared competences, roles, responsibilities and accountability to deliver a comprehensive continuum of care to a defined population |
| | Organisational integration (meso) | Inter-organisational relationships (e.g. contracting, strategic alliances, knowledge networks, mergers), including common governance mechanisms, to deliver comprehensive services to a defined population |
| | System integration (macro) | A horizontal and vertical integrated system, based on a coherent set of (informal and formal) rules and policies between care providers and external stakeholders for the benefit of people and populations |
| | Functional integration (micro, meso and macro) | Key support functions and activities (i.e. financial, management and information systems) structured around the primary process of service delivery to coordinate and support accountability and decision-making between organisations and professionals in order to add overall value to the system |
| | Normative integration (micro, meso and macro) | The development and maintenance of a common frame of reference (i.e. shared mission, vision, values and culture) between organisa- tions, professional groups and individuals |
| Measuring Integrated Patient Care Framework ²⁴ | Coordinated within care team | The individual providers (which may include physicians, nurses, other clinicians, support staff, and administrative personnel who routinely work together to provide medical care for a specified group of patients, hereafter the "care team") deliver consistent and informed patient care and administrative services for individual patients, regardless of the care team member providing them. |
| | Coordinated across care teams | All care teams that interact with patients, including specialists, hospital personnel, and pharmacies and deliver consistent and informed patient care and administrative services, regardless of the care team providing them. |
| | Coordinated between care teams and community resources | Care teams consider and coordinate support for patients by other teams offered in the community (e.g., Meals on Wheels). |
| | Continuous familiarity with patient over time | Clinical care team members are familiar with the patient's past medical condition and treatments; administrative care team members are familiar with patient's payment history and needs. |
| | Continuous proactive and responsive action between visits | Care team members reach out and respond to patients between visits; patients can access care and information 24/7. |
| | Patient centred | Care team members design care to meet patients' (also family members and other informal caregivers') needs and preferences; processes enhance patients' engagement in self-management. |
| | Shared responsibility | Both the patient and his or her family and care team members are responsible for the provision of care, maintenance of good health, and management of financial resources. |

Table 1. Models used to develop the coding framework

self-management support the most frequently cited processes. This finding is indicative of recommendations to focus on reducing fragmentation of services, without the requirement for integration at the organisational or system level.^{5,42} See Table 3 for details of the clinical integration processes described within each study.

Service/professional integration (meso level). The data once again reflected the recommended focus on clinical and service integration, with all but three of the studies^{28,33,40} describing service/professional integration processes. The processes ranged from simply managing and following up referrals³⁸ to multidisciplinary teams making decisions and planning care together, of which

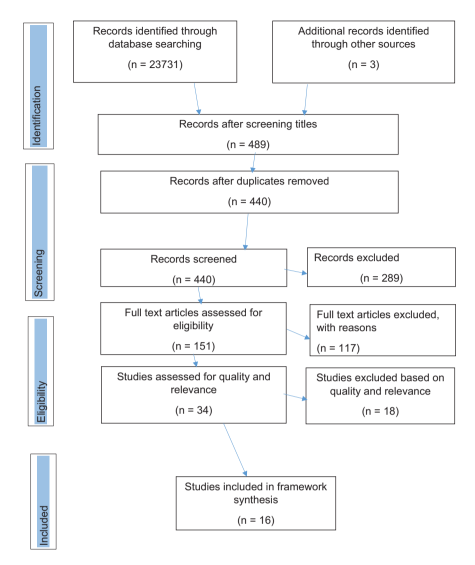


Figure I. PRISMA flow diagram.

these were the most commonly described processes.^{25–} ^{27,29,30,32,34,39} In Fairbrother et al.³⁶ service integration simply consisted of an Unscheduled Care Service taking over the weekend telemonitoring of people with chronic heart failure. Moule et al.³¹ described an integrated team consisting of one professional (social worker), unqualified practitioners and a customer service officer with a team manager, working with trained assessors from other disciplines and teams to perform assessments, sign off and provide payments. Payne et al.³² also described one of the hospices working as a hub for specialist palliative care services, supplementing primary care services and one consultant working across the hospital and hospice, while another hospice worked with specialist community heart failure nurses and community matrons to provide targeted services. In Wye et al.³⁵ professional integration focused on the role of a Discharge-In-Reach nurse who provided

education and information to other professionals about end of life care and available services.

Organisational integration (meso level). Organisational integration was generally described in less detail if at all. In some studies organisational integration consisted of agreements to work across organisational boundaries providing support²⁷ and education.^{28,31,32,38} In Hamilton et al.²⁸ assessors were trained across health, social care and voluntary organisations to perform assessments on behalf of the local authority. In other studies the multidisciplinary teams and/or key workers worked across organisational boundaries to deliver planned care.^{30–32,34,35} Payne et al.³² also described formal collaboration between hospices and other organisations with the use of similar paper-based records across one local care system. More formal arrangements such as co-location and/or management

| Table 2. Charac | Table 2. Characteristics and summary of studies included | | in the review, grouped by intervention type. | | |
|---|---|---|---|--|--|
| Author(s) | Study design | Target population | Intervention | Data collection and analysis | Summary of results |
| Interventions focuss Boudioni et al. 2015 ²⁵ | Interventions focused on people with or at risk of at least one long-term or age-related condition (n = 7) Boudioni et al. Experience-based code- Patients (65 years or over) Partnership betwo 2015 ²⁵ sign (EBCD) using visual in the community, authority (LA) thematic approach. mainly with long-term mary care trus or complex range of PCT hosted a t age-related conditions 8 integrated ca in a London borough. workers, manag the LA and co- within the com- | I least one long-term or age-rela Patients (65 years or over) in the community, mainly with long-term or complex range of age-related conditions in a London borough. | ted condition $(n = 7)$ Partnership between local authority (LA) and pri- mary care trust (PCT). PCT hosted a team of 8 integrated care social workers, managed by the LA and co-located within the community matron team. | Video recorded interviews (n = 7). Videos analysed by researcher with 2 independent service users trained in qualitative analysis. | Two main themes; I. Overwhelming feelings and emotions across their journey. 2. The emotional journey and experience of powerlessness. Receiving extra integrated care reinstated feelings of control and power for most participants. Through integrated care, most participants obtained knowledge and skills required to manage condition and |
| Gowing et al. 2016 ²⁷ | Qualitative | Patients at high risk of unplanned hospital admission in Northumberland. | Northumberland High Risk Patient Programme (NHRPP). Case finding of frail patients using a multidisciplinary team- led community case management pro- gramme and high-risk register. | Semi structured inter- views (n = 23) in own home by independent academic GP trainee. Thematic analysis. Coding framework developed by team with diverse backgrounds. | professionals. Four main themes; I. Awareness and understanding 2. Confidence in primary healthcare team. 3. Limitations of home care. 4. Active role of being a patient. Positive experience in general in terms of needs being met and having access to team members when needed. NHRPP experienced differently by different patients, especially those without informal carers, living alone without informal carers, living alone with deteriorating health. Further integration is needed to meet complex health, psychological and social needs of frail older people, as well as reduce unplanned |
| | | | | | readmissions. (continued) |

| Table 2. Continued | panu | | | | |
|------------------------------------|---|---|---|---|---|
| Author(s) | Study design | Target population | Intervention | Data collection and analysis | Summary of results |
| Hu 2014 ²⁹ | Mixed method post- implementation evaluation. | Older people requiring a defined range of com- munity services across Cambridgeshire. | Locality integrated multi- disciplinary teams across four PCT areas. | Face to face/telephone administered question- naire ($n = 100$) and semi-structured inter- views ($n = 27$). Data analysed using SPSS and thematic approach by one researcher. Coding checked by supervisor. | Ten domains. Satisfaction with services; Waiting times for assessment and receiving services; Awareness of the Cambridgeshire Direct Contact Centre; Improvement in physical functioning of some users of occu- pational equipment; Promotion of faster recovery from illness; Meeting basic physical needs; Living in a clean and tidy environment; Control over everyday life; Being treated as an individual with respect; Having a say and control over services. Positive outcomes for some but not majority. Low level of satisfaction with social care associated with impact of agency work, aim of achieving maintenance/prevention outcomes and staff behaviour/atti- tudes. No evidence of more positive outcomes than other integration models, with or without structural |
| Moule et al. 2014 ³¹ | Qualitative | Carers of people eligible for and receiving provi- sion from adult social care teams in Bristol. | Joint carers strategy sup- ported by Bristol City Council and Bristol City Clinical Commissioning Group. Integrated carers' teams work directly with carers to assess for one off direct payments to address outcomes identified. Also those funded by Continuing Health Care who do not have access to social workers. | Telephone interviews ($n = 39$) and home interview ($n = 1$). Interview schedule developed with input from 2 carer members of research team and project advisory board (carers and staff from both organisations). Thematic analysis using recognised framework. | changes. Four themes; 1. Administration 2. Circumstances of the carers 3. The break 4. Effects of the break Assessment mostly positive experience but challenging to reach isolated group. Integrated approach to carer breaks should be maintained. |

(continued)

| Table 2. Continued | nued | | | | |
|--|---|---|---|---|---|
| Author(s) | Study design | Target population | Intervention | Data collection and analysis | Summary of results |
| Payne et al. 201 <i>7</i> ³² | Qualitative longitudinal organisational case studies. | Patients and families with cancer and chronic conditions requiring palliative care support in Northern England. | Integration between hos- pices and local health care providers to pro- vide direct multidisci- plinary patient care to one or more diagnostic groups (advanced air- ways disease/heart fail- ure/cancer). | Semi-structured inter- views ($n = 47$) 3 months apart, mostly at home. Interviewed by researcher: Secondary thematic analysis of data from Integrated Palliative Care in cancer and chronic conditions (InSuP-C) study. Coding checked by another researcher. | All valued continuity of care, integrated working and named professional as point of contact. Key area – information sharing. Highlights needs of patients with non- cancer diagnosis and multi-morbid- ities. Greater integration by hospices required to work with, rather than replace local providers. Priorities for integration identified as – single information system, skilled named professional to coordinate care and |
| Randall et al. 2014 ³³ | Qualitative/Pragmatic | Patients with complex comorbid long-term conditions in a UK inner city PCT. | Case management by Community Matrons (CM), including care coordination, effective disease management and self-management support. | Semi-structured inter- views (n = 21). Domains of case man- agement used to guide framework analysis. | form meaningful relationships. Four themes; 1. Visibility 2. Interpersonal relationships 3. Leadership 4. System/Professional boundaries Greatest impact – relating to person as a whole rather than dealing with single disease. Coordination role most valuable. Consistent approval of CM service from patients and carers. Reducing levels of hospital admission major challenge due to fragmented services and limited vision of working in partnership held by other |
| Wye et al. 2014 ³⁵ | Realist evaluation | All palliative patients across North Somerset and Somerset. | Marie Curies Cancer Care's Delivering Choice Programme (DCP) working in part- nership with local pro- viders and commissioners to develop 24-hour serv- ices to meet local need. | Face to face and tele- phone semi-structured interviews (n = 43). Framework analysis. Local stakeholders attended hypothesis generation workshops. | professions. Highly skilled, experienced, customer- focused palliative professionals are important. Willingness to have diffi- cult conversations and access to community resources to support home care led to confidence and reassurance to patients. Inconsistency of referrals limited suc- cess and kept focus on those with cancer. Coordination Centres within |
| | | | | | (continued) |

8

| Author(s) | Study design | Target population | Intervention | Data collection and analysis | Summary of results |
|---|--|--|--|--|---|
| | | | | | social services on local authority premises and Discharge In Reach Nurses had most success accessing all eligible patients. 24 hour integrated palliative care serv- ices have an overview to steer fami- lies and patients through the system. |
| Interventions focu Darwin et al. 2017 ⁴⁰ | Interventions focused on those with a single condition/diagnosis $(n = 8)$ Darwin et al. Mixed method multisite Pregnant women from 2017 ⁴⁰ evaluation informed by low income commu realistic evaluation. ties, two sites restr ed to women from ethnic minority gro and third site has a l ethnic minority population. | ddition/diagnosis $(n = 8)$ Pregnant women from low income communi- ties, two sites restrict- ed to women from ethnic minority groups and third site has a large ethnic minority population. | Predominantly voluntary service organisations train volunteers from local community (doulas) to offer infor- mation, emotional and physical support. Work closely with existing services, facilitating communication and signposting to other services. | Questionnaires ($n = 136$) and semi structured interviews ($n = 12$). Discussions with key informants to develop topics and a priori hypotheses. Completed with assistance of researcher or inter- preter or self-complet- ed by post. Quantitative data analysed using | Most reported positive impacts on emotional wellbeing; combating feel- ings of depression, fears allayed, increased confidence and self- esteem. Continuing support beyond 6 weeks should be considered. Negative experiences most commonly associated with lack of continuity. Yolunteer doula services have potential to overcome barriers and facilitate access to improve outcomes of pregnant women with complex social |
| | | | | descriptive statistics and chi-squared with Yates' continuity cor- rection using SPSS ver- sion 20. Qualitative data analysed using content analysis. | factors. |
| Deslandes et al. 2015 ³⁹ | Qualitative case study | Patients with a diagnosis of mental illness attending a secondary care psychiatrist clinic in Wales. | A pharmacist supplemen- tary prescriber working in partnership with five psychiatrists and patients to provide dose titration and medicine information. | Semi-structured inter- views and self-complet- ed diaries (n = 11) over 6 weeks. Code and retrieve analysis of interviews. Thematic content analysis of dia- ries. All transcribing and analysis by main researcher. | Interview themes; I. Pharmacist-patient relationship. 2. Comparison to other professionals. 3. Time allowed for consultation. Diary themes; I. Patient satisfaction. 2. Consistency of care. 3. Pharmacist knowledge. 5. Mental health patient. Positive experience associated with increased access to and continuity of |
| | | | | | healthcare professional and more active role for patients in decisions. |

Table 2. Continued

(continued)

| Table 2. Continued | ned | | | | |
|--|---|---|---|--|--|
| Author(s) | Study design | Target population | Intervention | Data collection and analysis | Summary of results |
| Fairbrother et al. 2013 ³⁶ | Qualitative | All patients with chronic heart failure(CHF) in the community in Lothian, Scotland. | Telemonitoring of patients at home by GP or CHF nurse. | Semi-structured inter- views (n = 18) by 2 independent graduate researchers. Framework analysis. | Five themes; I. Information, support and reassurance. ance. 2. Compliance and dependence. 3 Changes and challenges. 4. Determining the criteria for patient applicability to telemonitoring. 5. Continuity of care. High level of satisfaction. Telemonitoring enhanced patients knowledge and understanding of condition and useful in management of CHF. |
| Firn et al. 2018 ²⁶ | Mixed method uncon- trolled observational follow-up design. | Mental health patients in suburban West London Borough. | A hybrid model that incorporates practice and principles of Assertive Outreach (AO) into Community Mental Health Teams (CMHT). | Validated Team Attachment Questionnaire (TAQ) 12 months pre (n = 37) and post (n = 33) ser- vice change. Appreciative Inquiry method to free text responses.Change in measures of patient experience reported using mean change and 95% confidence inter- vals, paired t-tests to test significance. | TAQ scored highly in both services. Four themes; I. Continuity. I. Continuity. I. I. Continuity. I. Indifference to change. Reduction in contact and support. AO may have over provided care, nurtured dependency and continued beyond current need. Unnecessary dependence may explain concerns about reduced contact after integration. |
| Hamilton et al. 2016 ²⁸ | Qualitative | People eligible for publicly funded social care sup- port within a mental health service. Across three local authority areas (rural South England, urban North England and London borough). | Social workers, occupa- tional therapists and community psychiatric nurses work in part- nership with client to assess personal budget needs on behalf of local authority. | In-depth semi-structured interviews (n = 52). Analysed using Interpretive Framework Approach in collaboration with advisory board of ten service user researchers. | One group of themes relating to power, choice and control. Generic local authority systems not well suited to people with mental health conditions whose level of need and capacity may fluctuate and may find process stressful. |
| | | | | | (continued) |

| Table 2. Continued | ned | | | | |
|--|---|---|---|---|---|
| Author(s) | Study design | Target population | Intervention | Data collection and analysis | Summary of results |
| Primeau et al. 2017 ³⁷ | Qualitative component of a randomised con- trolled trial (RCT). | Men with metastatic prostate cancer in Tayside, Scotland. | Multi-modal supportive care intervention (Thriver -Care) which includes: a holistic needs assessment with a prostate cancer spe- cialist nurse (PCSN), individualised self-man- agement care plans, group-based seminar and educational materi- als. Compared with standard care. | Semi-structured interviews at home or views at home or cancer centre ($n = 26$). Framework analysis. Data checked by peers to verify themes. | Overall high satisfaction with interven- tion. Perceived empowerment in their self-management. Important themes; being listened to by some- one who could facilitate emotional expression; being provided individu- ally tailored information; receiving practical help and evidence-based advice for managing consequences of cancer and it's treatment. Experienced less unmet needs than standard care group. |
| Thomson et al. 2018 ³⁴ | Qualitative component of larger randomised con- trolled trial. | People diagnosed with major depressive disor- der across Nottinghamshire, Cambridgeshire and Derbyshire. | A collaborative and inte- grated psychobiosocial model. Joint assess- ments by psychiatrists and psychological therapists, with devel- opment of structured management plans. | Semi-structured interviews ($n = 21$) with experienced qualitative interviewer at 12-18 months after baseline. Inductive thematic analysis using grounded approach. Second researcher coded and themes checked by team. | Four themes; I. Specific treatment components. 2. Individual qualities of clinicians. 3. Collaborative team approach. 4. Accessibility of treatment. Positive experiences centred on trust- ing relationships with therapists, stable and collaborative team, fre- quency and accessibility of sessions and ability to top-up/reinforce tech- niques learned over a longer period. Transition to usual care difficult for some. |
| Young and Snowden 2019 ³⁸ | Qualitative component of longitudinal mixed methods study. | People with a new diag- nosis of cancer in Glasgow City Council area. | An integrated community- based supportive cancer service run by a joint health and social care board. | Semi-structured inter- views (n = 20), mostly at home. Conducted by one researcher. Thematic analysis by two researchers. | iemes; e person, one place utes to unexpected sty net. I for majority of pa impact associated w to discuss their col one separate to clin one separate to clin one separate to clin ent/personalised gui ;e. |
| | | | | | (continued) |

| Author(s) | Study design | Target population | Intervention | Data collection and analysis | Summary of results |
|---------------------------------------|--|--|--|---|---|
| Interventions whi Mastellos et al. | Interventions which focused on people with a single diagnosis and those with age related conditions $(n = 1)$ Mastellos et al. A cross-sectional survey Patients registered with Practices were offered | ingle diagnosis and those with Patients registered with | age related conditions $(n = I)$ Practices were offered | St | Positive experience with Integrated |
| 201430 | design. | the Integrated Care Pilot in North West | incentives to develop specific, bespoke care | Likert-scale question- naire (n $=$ 405). Paper- | Care Pilot. Benefits for those aware of having a |
| | | London who had a | plans in order to iden- | based and self-complet- | care plan; increased involvement in |
| | | diagnosis of diabetes | tify those at risk of | ed. | decision-making, improved patient- |
| | | and /or were aged over | hospital admission and | Analysed using software | provider relationship, better organi- |
| | | 75 years. | coordinate care across | and descriptive statis- | sation and access to care, enhanced |
| | | | services in primary, | tics used. Case analysis | communication amongst healthcare |
| | | | secondary and commu- | used for each variable | providers. |
| | | | nity care. | to handle missing data. | Incentive structures may be open to |
| | | | | | abuse. |
| | | | | | Provides empirical evidence integrated |
| | | | | | care has potential to improve patient |

experience.

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of health and social care professionals under one organisation were found in three studies.^{25,29,35} In Boudioni et al.²⁵ health (community matrons) and social care (social workers) professionals were co-located within the Primary Care Trust. However, social workers continued to be managed by the local authority. In Hu²⁹ professionals across health and social care for older people were employed directly by one Older People's Service.

System integration (macro level). Less than half of the studies described integration at this level, reflecting well-documented barriers which continue to exist regarding funding and lack of shared objectives.^{43,44} Seven studies provided limited description of the following system integration processes; national policy or funding;^{27,35,38} joint commissioning and a pooled budget;²⁹ legislation²⁸ and service level agreements.³²

Normative and functional integration (supportive). Supportive processes are cited as important to achieving integrated care at all levels,^{23,43} however only nine studies described any. The use of technology was the most frequently cited support process.^{27,30,32,35} Other processes focused on creating shared culture,^{29,40} learning^{29,31,32,38} and governance.^{29,31,34,35,38}

Experiences of person-centred coordinated care (PCCC). The experiences captured within the studies are summarised in Table 4. A positive experience is defined as when the patient or carer has described a beneficial or positive effect when experiencing a particular domain of care. Conversely, when experience of a domain has had a detrimental effect to the person it is recorded as a negative experience. Some of the comments and in particular the quantitative data only confirms the domain was experienced, without any indication of positive or negative views, these are recorded as neutral experiences. An opposing experience was recorded when a person described an experience which was the opposite of a particular domain e.g. they were not supported to share decision making or responsibility for their care. Only one study described patient experience of all domains of PCCC.34

Coordinated care within the team

This domain received limited descriptions of patient experience, which were mainly neutral.^{25,27–31,34} Much of the coordination referred to within this domain relied on communication between team members and administrative processes, which are not always evident to patients and carers.

Table 2. Continued

| Study | Care coordination | Care planning | Key worker | Remote monitoring | Self-management support | Supplementary prescribing | Total number of processes described |
|---------------------------------|----------------------|------------------|---------------|----------------------|----------------------------|---------------------------|---|
| Boudioni ²⁵ | • | • | • | | | • | 4 |
| Darwin ⁴⁰ | | | • | | | | 1 |
| Deslandes ³⁹ | | | | | • | • | 2 |
| Fairbrother ³⁶ | | | • | • | • | | 3 |
| Firn ²⁶ | • | • | • | | • | | 4 |
| Gowing ²⁷ | • | • | • | | • | | 4 |
| Hamilton ²⁸ | | • | | | | | I |
| Hu ²⁹ | • | | • | | • | | 3 |
| Mastellos ³⁰ | • | • | | | • | | 3 |
| Moule et al. ³¹ | | • | | | | | I |
| Payne et al. ³² | • | | • | | | | 2 |
| Primeau ³⁷ | | • | • | | • | | 3 |
| Randall ³³ | • | • | • | | • | • | 5 |
| Thomson ³⁴ | • | • | | | • | | 3 |
| Wye ³⁵ | • | | • | | • | | 3 |
| Young and Snowden ³⁸ | • | • | • | | | | 3 |
| Total number of studies | 10 (63%) | 10 (63%) | (69%) | l (6%) | 10 (63%) | 3 (19%) | 45 (47% of maximum number of processes available) |

Table 3. Summary of clinical integration processes at the micro level.

Coordinated care across teams

All studies except for three^{36,38,39} included experiences of this domain. Only two studies included a negative experience.^{26,29} For people with complex care needs the experience was generally positive.²⁷ In Wye et al.³⁵ a coordination centre put a care package in place to support a dying person at home. However, two studies described a lack of coordinated care across teams,^{31,32} which was described as people being "too scared to cross each other's territories".³²

Coordination of care across the team and between wider community services

Experience of coordination across the team and between wider community services received only negative descriptions in Firn et al.²⁶ and Hu,²⁹ while in Young and Snowden³⁸ they were all positive. The majority of negative experiences related to social care services and the often frustrating limitations in terms of time and quality of care provided. For example, in Hu²⁹ ten patients reported their basic physical needs had not been met due to a decline in social care quality. Positive experiences relating to social care involved carers feeling supported through the provision of overnight care²⁷ and home care packages to support dving at home.³⁵ Opposing experiences were described by palliative care patients regarding lack of

communication and information sharing between health and social care providers.^{32,35}

Continuous familiarity with a service over time

While there were mixed experiences of this domain, it was only described negatively in two studies where patients felt unsupported when a transition to another service or professional was required.^{32,34} Some patients found it to be an important factor in establishing recovery, which points to a need for personalisation and flexibility in services.³⁴ While in Hu²⁹ social care experiences demonstrated a lack of continuous familiarity. For older people, those with complex conditions and those with a single condition, the value of access to a key worker who provided continuity featured in many positive experiences.^{25,28,32,33,37-40}

Continuous proactive and responsive action between visits

The experience of this domain received mainly positive descriptions.^{25,27,30,32–39} The experiences reflected a sense of reassurance from being actively followed up and being able to contact someone when needed, particularly older people with complex needs.²⁷ Where an opposing experience had been described it related to patients feeling they were still having to follow-up and coordinate care themselves.^{25,32}

| | Coordinated care within | Coordinated | Coordinated across team and community | Continuous familiarity | Continuous proactive and responsive action | | |
|---------------------------------|----------------------------|----------------------------|---|---------------------------|--|----------------------------|-------------------------|
| Study | team | care across team | services | over time | between visits | Patient centred | Shared responsibility |
| Boudioni ²⁵ | + | + / | | + | + × | + /×× | + / × |
| Darwin ⁴⁰ | | | | ×+/++/ | | $+ \times \times + + /$ | + |
| Deslandes ³⁹ | | | | ++ | + | + | ++ |
| Fairbrother ³⁶ | | | | | + | ++ | + × |
| Firn ²⁶ | | - / | I | ++++ | | + + + | |
| Gowing ²⁷ | 111 | + / | + -++++++++++++++++++++++++++++++++++++ | | ++++ | + / + / + + + | + |
| Hamilton ²⁸ | + | | | | | $+$ \times $ +$ \times | $- \times - \times + +$ |
| Hu ²⁹ | $+ \times /$ | ++ | | ×× | | $+I \times \times$ | + / × |
| Mastellos ³⁰ | / | | | / | 1 | × / /+/ / | //× |
| Moule ³¹ | / × | × | | | | /++-/++/ | |
| Payne ³² | | | | | | | |
| Hospice A | | $\times + \times + \times$ | I | + + × / + | | + | |
| Hospice B | | × | + | + | | + | |
| Hospice C | | /++/ | × | + | | | |
| Hospice D | | //+×× | + | / | $+ \times /$ | × + | |
| Primeau ³⁷ | | | | / | | /+++ | 11 |
| Randall ³³ | | ++ | | / | ++ | + | + |
| Thomson ³⁴ | / | + | 1 | + + | + | + /++ | - / /+/ |
| Wye ³⁵ | | ++++ | + × | | ++++ | + | |
| Young and Snowden ³⁸ | 38 | | + + + + | ++ | ++++++++ | × ×// | |

14

Patient/person-centred care

Patients and carers described the experience of personcentred care frequently and in all studies, which links to the focus on micro level integration within the studies. There were positive experiences across all studies and negative experiences in only four studies, related to lack of flexibility and personalisation in practitioners approaches.^{26-28,31} For those with single conditions positive experiences centred on increased knowledge³⁷ and feeling listened to.³⁹ However, nine studies included experiences which demonstrated a lack of person centred care, ^{27–32,38,40} in particular when policy dictated when and for how long a service was available rather than individual need. For example, 33.1% of women receiving a doula service felt support ended too soon and at a difficult time.⁴⁰ In Mastellos et al.,³⁰ despite 78.8% of patients understanding how their care plan worked, only 36.4% were involved in creating the care plan.

Shared responsibility

The experience of shared responsibility was the second most described within the studies and reflects the frequency of self-management support in these studies.^{25,27–30,33,34,36,39,40} However, three of these studies included negative experiences.^{27,28,34} In Hamilton et al.,²⁸ people with mental health conditions felt overwhelmed by the expectation of shared decision-making regarding their personal budgets or felt pressured because they did not want to challenge the healthcare professional. Older people with complex needs also reported mixed experiences; some welcomed self-help as a good thing²⁷ and felt able to have a say in decisions, whilst others felt they did not have this right.²⁹

Associations between integration processes and person-centred coordinated care

Associations between integration processes and experience of person-centred coordinated care were very limited in the included studies. Boudioni et al.²⁵ associated feelings of power and control, improving selfmanagement skills and becoming active partners with professionals, with the overall service rather than a particular process. Hamilton et al.²⁸ acknowledged the experience of taking control was influenced by organisational culture and processes, as well as practitioner skills and attitudes. Positive experiences were associated with increased access to and continuity of healthcare professionals, along with a more active role in decisions,³⁹ individualised self-management interventions,³⁷ ability to accommodate preferences,^{31,37} trusting relationships and a collaborative team³⁴ and increased access to community resources for home care.³⁵ Negative experiences were associated with a lack of shared organisational culture such as social care's focus on maintenance and prevention, as well as staff behaviour and attitudes.²⁹

Discussion

Sixteen studies were identified which explored patient experience of integrated care within the UK since the introduction of the Health and Social Care Act.¹⁸ Included studies examined varied types and levels of integration; some focused on single services, some explored changes to multiple services and others addressed the use of personal budgets to allow choice in services received. The breadth of integration processes examined, and the joint focus on integration and patient experience forms a key strength of this review.

Limited associations were made between integration processes and patient experience despite patient experience being a central component of integrated care.⁷ Eight studies included in this review identified positive experiences of integrated care, with the majority of experience data focused on person centred care and shared responsibility. A negative experience of patient centred coordinated care occurred when integration did not allow for the flexibility and responsiveness required to meet individual needs⁴⁵ or when services try and fit people into a particular model that may not be appropriate.⁴⁰

Despite evidence that integrating health and social care improves access to care for people complex and palliative care needs,⁴⁶ people with complex needs continued to experience a lack of coordination across teams and wider community resources.32,35 Findings such as this strengthen the need to understand which integration processes are associated with patient centred coordinated care.²⁴ To facilitate this, research into integrated care would benefit from the use of a common framework or language regarding integration processes to improve shared understanding of links between implementation and patient experience.47 Comparing individual patient experiences is challenging⁴⁸ and was found to be more achievable within studies where context and social factors were taken into account.38,40

Overall, the experiences of patients in this review support the view that person centred care is a core element of integrated care.^{9,24} However, the argument that person centred care is often defined by the behaviour of health professionals⁹ remains evident. While definitions of integrated care remain unclear, this review supports the findings of previous research⁴⁹ that patients are clear on the concept of patientcentred care coordination as their experiences reflect the importance of coordination (within, across and with wider community resources), continuity, information sharing and engagement. As the patient can be the only person to have a holistic perspective of their care, particularly across multiple services, capturing data on their experiences is an important component towards understanding complexity, though patient experience data alone is insufficient for determining whether models of integration can be deemed successful.⁵⁰ The insights gained from this review affirm the potential of integrated care to provide PCCC but also reveal the potential for negative or opposing patient experiences when integration fails to go beyond the clinical level.

A potential lesson for other developed countries is the failure to treat healthcare systems as complexadaptive, which has led to poor professional and organisational alignment, lack of funding incentives for collaboration and a continued top-down approach to management.⁴ Integrated care requires the facilitation of strong relationships between providers and organisations over time,⁴ which supports emergent collective insight and distributed control. However, confusion exists regarding the complex interactions between integration and integrated care which can occur at different levels of the health system.⁵¹ In the UK, the Covid-19 pandemic has highlighted the need for a legislative framework which encourages provider collaboration and flexibility of service provision in order to effectively provide integrated care.

A limitation of this systematic review is that the data extraction process was completed by one researcher. However, the process was repeated to increase intrarater reliability. A clear quality appraisal process enabled the identification of the most relevant studies for the review which also improved the overall validity of the findings. The exclusion of any non-empirical unpublished studies and low-quality studies does however mean this could not be considered a comprehensive review of all available data, but instead of only the highest quality data.

In conclusion, the findings of this review highlight the importance of measuring patient experience of integrated care, which could facilitate a shared understanding of experiences across health and social care. However, in order to compare experiences across such broad health, social and third sector services, far more focus on integration processes is required than were found within this review. The complexity of integration processes, which take many different forms and occur simultaneously at micro, meso and macro levels, pose key challenges in regard to conceptual clarity and practical application. As such, there is a need for greater consistency in the definition of this construct in order to facilitate its understanding and implementation. Likewise, future work should consider the use of validated measures of patient experience of PCCC in addition to in-depth qualitative methods. The review demonstrated the need for further work highlighting the processes which support integration more explicitly, in order to inform its operationalisation. Evidence on the relationship between integration and patient experience was shown to be scant, with this forming a key avenue for future research. In particular, the impact of organisational and system level integration for patients is currently very poorly understood, relative to more micro integration processes. The combined application of the Rainbow Model of Integrated Care and the Measuring Integrated Care Patient Framework²⁴ within this paper provides a foundation for other work examining the links between integration processes and patient experience.

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