Influences on Care Preferences of Older People with Advanced Illness: A Systematic Review and Thematic Synthesis

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OBJECTIVES: To determine and explore the influences on care preferences of older people with advanced illness and integrate our results into a model to guide practice and research.

DESIGN: Systematic review using Medline, Embase, PsychINFO, Web of Science, and OpenGrey databases from inception to February 2017 and reference and citation list searching. Included articles investigated influences on care preference using qualitative or quantitative methodology. Thematic synthesis of qualitative articles and narrative synthesis of quantitative articles were undertaken.

SETTING: Hospital and community care settings.

PARTICIPANTS: Older adults with advanced illness, including people with specific illnesses and markers of advanced disease, populations identified as in the last year of life, or individuals receiving palliative care (N = 15,164).

MEASUREMENTS: The QualSys criteria were used to assess study quality.

RESULTS: Of 12,142 search results, 57 articles were included. Family and care context, illness, and individual factors interact to influence care preferences. Support from and burden on family and loved ones were prominent influences on care preferences. Mechanisms by which preferences are influenced include the process of trading-off between competing priorities, making choices based on expected outcome, level of engagement, and individual ability to form and express preferences.

CONCLUSION: Family is particularly important as an influence on care preferences, which are influenced by complex interaction of family, individual, and illness factors. To support preferences, clinicians should consider older people with illnesses and their families together as a unit of care.

Worldwide demographic changes mean that more people are living with and dying from chronic illness and multimorbidity, and this number is expected to rise. The prevalence of chronic illness rises with age, so delivery of high-quality care in this growing population group is a priority.

Care of older people with chronic illness is often complex, and to deliver person-centered care in this population, a clear understanding of the person's preferences for care is needed. Person-centered care seeks to provide what is necessary to meet individuals’ physical, psychological, social, and spiritual needs by focusing on what is important to them, and care preferences can be defined as what people want from their care. They can be broadly classified as preferences for the context in which care is delivered; preferences for care relationships; preferences for involvement in care; and preferences for care outcomes, such as comfort versus extending life or place of death. Observational studies have determined the content of care preferences in this population.

Nevertheless, people do not simply have preferences, so we need to know more than the content of preferences to deliver responsive care. Preferences may become clear only as they are elicited or may vary in importance at different times. Equally, changes in contextual factors or experience may influence preferences. Influences on care preferences have been investigated in qualitative and quantitative studies, particularly factors associated with preference for home care or home death. One model lists illness-related factors, individual factors, and environmental factors associated with place of death. Qualitative studies have highlighted the importance of family support, personal experience and values as influences on preferences, but this evidence is mainly from small observational studies. This body of evidence has never been synthesized.

Key words: systematic review; patient preference; terminally ill; terminal care; palliative care

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To deliver care in line with personal preferences, it is important to understand how they are influenced at a broader level. Therefore, this review aimed to synthesize evidence regarding the influences on preferences for care of older people with advanced illness, producing a model of the influences on care preferences in this population.

**METHODS**

This systematic review followed the principles of the Preferred Reporting Items for Systematic Reviews and Meta-Analyses guidelines and was based on a pre-agreed protocol.

**Search strategy and selection of articles**

We searched online databases (Medline (Ovid), Embase, PsycINFO, Web of Science), grey literature repositories (Open Grey), and theses (ETHos database) from inception to February 2017. The search strategy used Medical Subject Headings and synonyms related to “preferences,” “advanced illness,” and “older people” and was adapted from search strategies used in previous systematic reviews. See Appendix S1 for the full search strategy. No limits were placed on language or date. We also hand-searched reference and citation lists of included articles. The search was updated in December 2017. See Table 1 for inclusion and exclusion criteria, and article selection process.

**Quality assessment**

Because this review includes studies of varied design, we used the validated QualSys guidelines, designed for this purpose, to assess the quality of included articles. QualSys incorporates separate checklists of criteria for qualitative and quantitative studies. Each criterion is rated on a scale from 0 to 2, and a total score is calculated. The research team agreed that a score of 75% or greater was required for high quality and 60% or greater for medium quality. No articles were excluded on quality, although we used QualSys scores, alongside quantity and agreement of evidence, to appraise the strength of evidence supporting the quantitative synthesis, based on existing criteria designed for this purpose.

**Data extraction and analysis**

Data were extracted from included articles using a bespoke pro forma that the research team agreed on in advance. Basic data from each study (e.g., authors, country, sample size) were extracted. For quantitative articles, findings regarding influences on care preferences were extracted. For qualitative studies, the entire Results or Findings section was extracted for thematic synthesis.

The “three synthesis” approach proposed for reviews of mixed studies was used in the analysis. Thus two initial syntheses were undertaken, one for each for quantitative and qualitative studies, followed by a third integrative synthesis.

Thematic synthesis, supported by NVIVO Software version 10 (QSR International (UK) Ltd.), was used for qualitative synthesis. This involved reading, re-reading, and then coding all relevant text within the Results sections of included articles. We first coded inductively, allowing for themes to emerge from the data. Deductive coding, based on existing models, was then applied to ensure relevance to previous work. In particular, we used the Gomes model, which identifies illness, individual, and environmental influences on place of death, and the experience, health condition, and family components of the Hattori model. Inductive and deductive coding were combined to produce a final model that fit with the original data. Using the entire Results section for analysis meant that raw data and author interpretations were synthesized, allowing us to build on existing work.

For the quantitative synthesis, the variation in outcomes across included articles precluded meaningful meta-analysis. Therefore a descriptive synthesis of factors associated with care preferences was undertaken. Factors were grouped by preference category, and the direction of the effect was recorded. Associations were linked to the strength of evidence supporting them.

### Table 1. Inclusion and Exclusion Criteria

**Inclusion criteria**

- Study population aged ≥65, or median age ≥65 if mixed population (excluded if >25% of participants aged <60, or >10% aged <40).
- Study is of people with advanced illness, including people with specific illnesses and markers of advanced disease, populations identified in the last year of life, or populations identified as palliative care populations. See Appendix S2 for detailed definition.
- Study focuses primarily on what influences care preferences. We were interested in the broad areas of care context, relationships, and outcomes, rather than specific preferences about individual treatments or management options, and included preferences relating to any aspect of care, including physical, psychological, social, and spiritual. See Appendix S2 for detailed definition.

**Exclusion criteria**

- Review articles and editorials.
- Articles focusing only on concordance of preferences between patients and family or clinicians.
- Studies in healthy older people, on the basis that those with good health may have perspectives on care preferences different from perspectives of those with advanced illness.
- Studies of preferences for specific treatments including cardiopulmonary resuscitation, which have been synthesised elsewhere.

Articles were screened by title and abstract, and those unrelated to influences on care preferences were discarded. Full texts of remaining articles were assessed for inclusion by one researcher (SE), with 10% cross-checking by another researcher (AB). Disagreements were resolved by discussion within the multidisciplinary research team. When full text was unavailable, an attempt was made to contact the original authors. Attempts were also made to find follow-up articles from relevant conference abstracts.
In the integration stage, findings from the quantitative synthesis were mapped onto the qualitative model to produce an integrated model of the influences on care preferences, using existing models as a starting point. Triangulation of qualitative and quantitative data was used to identify areas where evidence from different methodology concurred and areas where there was disagreement.

RESULTS

The search retrieved 8,723 records once duplicates were removed; 391 full-text articles were reviewed, and of these, 26 qualitative papers, 30 quantitative papers, and 1 mixed-methods analysis were included (Figure 1). Included papers were published between 1995 and 2017. In total, 14,474, unique participants were included in the quantitative papers and 857 in the qualitative papers. Twenty-five (44%) studies were conducted in North America, 19 (33%) in Europe, 8 (14%) in Asia, and 3 (5%) in Australasia, and 2 (4%) collected data internationally.

The 27 qualitative papers (Table 2) investigated influences on preferences for care context (place of care), care involvement (communication and decision-making), and care outcomes (quality of life, place of death). Thirteen papers were of high quality, 13 of medium quality, and 1 of low quality. All studies used interviews, ranging from structured interviews to unstructured in-depth interviews. The 30 quantitative papers (see Appendix S3 for details) varied in sample size from 38 to 2,452, and methodology ranged from in-person questionnaires to national postal surveys. Sixteen were high quality, 11 medium quality, and 3 low quality.

Based on qualitative, quantitative, and integration syntheses, we produced a model of the influences on preferences for care, which extends the Gomes model of influences on place of death. Factors influencing care preferences relating to family and care context, the individual, and the illness were identified and explored. We also identified mechanisms by which preferences are influenced (Figure 2). The synthesis indicates that it is the norm in this population for preferences to be incompletely expressed, and a clear decision-making process regarding preferences is rare. Instead, preferences emerge from a complex interaction of the areas and mechanisms identified in this model, as described below.

Three areas describe the influences on care preferences in this population: family and care context, the individual, and the illness. How these are derived from the included qualitative papers and aligned to the model is outlined below.

Family and care context

The strongest influences on preferences came from concerns of and about family, and the wider care environment. We found that trusting care relationships influence preferences...
for communication and that the level of support available from health professionals and family is an important determinant of preferences for care. The wish to avoid being a burden often influenced preferences, and the views of and concerns about family were of paramount importance:

Just to keep my daughter as calm as possible—that’s the main thing. I don’t want to upset her any more than I have to. What can you do? Your parents die, that’s going to happen, so … there is nothing to be done about it, but I want to, want her to be as calm as possible. (82-year-old female) Gardner et al.43

Frequently, concerns about family were more prominent than individuals’ own concerns about care preferences. Although patients’ wishes may not be their preference, such was their concern about their carer and not wanting to become a burden; they were not prepared to consider any other option for place of end-of-life care. McCall et al.53

The Individual

Characteristics of individuals such as age influenced preferences in some cases. Past experience of care, especially caring for others at the end of life, influenced preferences for one’s own care, whether relevant to one’s situation or not.
One of the most universal and influential events in the lives of the residents was the death of a family member or close friend. This influence was attributed to grief, and by facts about end-of-life decisions and death learned from the experience. In many cases, the situation of the loved one was not comparable with the resident’s current situation. Nevertheless, many residents applied the experience of the deceased as if it were identical to their own situation. The opinions generated by these experiences were so strong that they were not overridden by factual information about the residents’ own health. Lambert et al.23

Perceptions of care, often based on experience, were seen to shape preferences, sometimes very strongly.

Beliefs about what medical care was possible ... shaped preferences so fundamentally that a number of respondents could not form a preference distinct from these beliefs. Fried et al. (1998).42

Personal values, which are often long-held views, were another strong influence on preferences, for example a value of maintaining comfort may result in prioritization of quality over quantity of life.

The desire to “die easily, without suffering” was the most prevalent underlying wish, and one which never changed. Hattori et al.23

A long-held desire to maintain independence may influence preference for place of care.

I want to be self-sufficient. I don’t want to be sick. Until I’m overwhelmed, I want to be able to deal with it [my illness] on my terms (CA1030).54

The Illness

The illness context strongly influenced preferences, particularly for those who were aware of their disease status and accepted their situation.

But you know it has to end sometime when you are 80 and are also terminally ill [cancer]. Then you know that you have to accept it. [interview 25] Piers et al.55

Concerns about future changes in health were another area influencing care preferences.

A number of participants wished to be placed in a nursing facility when the time came, assuming their physical functions would deteriorate and they would require nursing care in the future. Hattori et al.23

MECHANISMS

In the qualitative synthesis, we identified mechanisms by which these areas interact, namely, expectations, trade-offs, engagement, and ability to form and express preferences.

Expectations

Personal experience, perceptions, illness context, and level of knowledge combined to produce expectations about the future, which guided care preferences, especially with regard to place of care.

Preferences for site of care depended on the anticipated outcome of the illness episode. Fried et al. 1998.42

Maintaining and maximising personal dignity given the prospect of loss of control over bodily functions, of extreme pain, or of ‘going mad,’ was bound up with a preference to place their final care in the hands of professionals in a hospice setting. Thomas et al.59

Trade-Offs

Personal values, illness context, and the wider care environment may be traded off in formulation of preferences. Presence and support of family exerted a strong influence on preferences, and the views of loved ones may supersede personal views and perceptions.

I would prefer to be at home. Um, but then again, by the same token, if hubby doesn’t think he can cope, which, it may come to that point, where he can’t cope, with the whole physical thing, the mental thing of me being at home, then I will willingly come in [again]. (Female, married, 51–60). Broom et al.58

Engagement

Preferences were not always actively expressed, and disengagement, or “living in the moment,” was common. Leaving decisions to chance or delegating to others may result.

I never thought about that. I think it should be left to the doctor. Fleming et al.40

Conversely, some seek to keep control and maintain independence and may express stronger preferences. Engagement also depends on temporal focus, acceptance of the illness context and personal self-efficacy.

I am my own boss, am I not? My children don’t control me. I decide what I want. Whether they like it or not, yes it’s my decision. It’s my body. That’s my opinion and no one can help me with that, no. [interview 30]. Piers et al.55

Ability to form and express preferences

Illness, cognitive impairment, or disengagement commonly resulted in low self-efficacy, such that preferences were not formed or expressed. Expectations, trade-offs, and engagement were less relevant to this group, who were unable to form preferences using these mechanisms.

The data suggested that perceptions of self-efficacy may influence not only the likelihood of setting explicit goals but also the nature of goals themselves. Low self-efficacy resulted in either the setting of no or less challenging goals. Bradley et al.24

The quantitative synthesis also identified factors associated with care preferences, which we used to confirm
DISCUSSION

This systematic review and thematic synthesis brings together, for the first time, qualitative and quantitative evidence regarding influences on care preferences in older people with advanced illness. Family and care context, particularly support from family, are dominant influences on care preferences in this population. These factors interact with individual—sociodemographic characteristics, personal experience, and values—and illness-related factors to influence care preferences. We identify four mechanisms affecting how these areas interact, conscious or unconscious trade-offs, consideration of what is expected to occur, engagement with preferences, and ability to form or express preferences.

The great importance of family support is evident in qualitative and quantitative data. “Living with someone” was the only factor associated with a care preference with high strength evidence. In the qualitative synthesis, concerns about family and concerns of family were strong enough to cause many people to change their preferences. The nuances of family relationships are therefore important in determining preferences. This supports the concept in the palliative care literature that the unit of care should be the person and his or her family, not the person alone.62 Previous work has demonstrated the importance of family in decision-making,63,64 but has not demonstrated the influence of family support on a wide range of preferences.

Our findings highlight the overarching importance of family and perceived family support when discussing preferences in practice, and it is clear that family should be part of these conversations. Because concerns about family may result in people expressing preferences different from their underlying wishes, clinicians cannot necessarily take stated preferences at face value. Instead, they should consider how they can best provide support to patients and their families to allow patients to achieve their preferences, particularly when there is conflict in views between individuals and those close to them.

We also identify potential areas clinicians could focus on to support patient preferences. It may be impossible to change illness or individual factors, but it is potentially possible to make positive changes to care and to support family involvement.65 An individual whose preference for place of care is home but is concerned about the effect on his or her family may trade off and change his or her preference to institutional care. Discussion about how the family may be better supported might allow the individual to stay in their preferred place of care.

The mechanisms we identify in this synthesis extend existing theory. Economic theories of preference, such as expectancy value theory, consider preferences as a process of trading off expected outcome and the value one would place on that outcome.66 Expectations are highlighted as a mechanism for “response shift,”16 and cognitive theories of preference also consider the process of trading off to be important,67 although these theories do not consider the importance of individual engagement and the ability to form and express preferences, which came across strongly in our synthesis. Older people, especially those with cognitive impairment, may choose to cede control or may not wish to consider preferences overtly. Others may be cognitively unable to express preferences.44 In this situation, the importance of the care context and family support grows.8 Our model therefore highlights the importance of broader, nonrational processes influencing preferences,6,68 as well as the trade-offs and expectation-based mechanisms proposed in economic and cognitive models.
This review has found considerable research into care preferences in this population, but many studies are small and not of high quality, with inconsistent reporting of nonsignificant findings. Some areas that might be expected to influence preferences—ethnicity, culture, religion—were not prominent in this synthesis, although there was some evidence that religiosity is associated with preference for place of care. This probably represents absence of evidence, rather than evidence that these areas do not influence preferences, and further study is needed.

Most included quantitative studies are cross-sectional, which limits the inferences that can be drawn from associations. The influence of time on preferences is evident in some of the qualitative data, but it is unclear how the various influences on preference interact to affect the stability of preferences over time. Studies that collect data serially show that a large minority of participants change their preferences, so it is important to know how changes in illness, individual, family, and environmental factors over time may affect preference stability. Further prospective research is needed to describe and explore influences on the stability of care preferences in this population.

A strength of this review is the systematic methodology, resulting in a robust model based on international qualitative and quantitative data. The three synthesis approach allowed triangulation between methodologies, which increased the robustness of our model. Inclusion of articles investigating a range of care preferences increased the scope of the model and its applicability. Although the evidence in this area is spread across fields and is challenging to identify, our use of a systematic, evidence-based search strategy without limitations on language or publication date means that we are confident we have identified the relevant evidence, although it is possible that some studies were missed. We found that quantitative studies inconsistently reported nonsignificant findings. Our synthesis therefore focused primarily on significant findings, which may have introduced a bias against negative findings, although including the nonsignificant findings that were reported did not affect our final model. Focusing on a more specific set of care preferences might have resulted in a more precise, albeit less broadly applicable synthesis. Additionally, thematic analysis of published articles inevitably limits the conclusions that can be drawn from the data because there is no access to the original data set.

In conclusion, this systematic review shows the importance of family and care context, particularly family support, as influences on care preferences. These factors combine with individual and illness-related factors to influence preferences through mechanisms of trade-offs, expectations, engagement, and the ability to form and express preferences. Clinicians must take these factors and mechanisms into account when considering preferences. Influences on the stability of care preferences are unclear, and further investigation with prospective longitudinal research is needed.

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SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article:

Appendix S1. Full search strategy for each database.
Appendix S2. Definitions.
Appendix S3. Details of quantitative papers.
Appendix S4. Full quantitative synthesis.
Appendix S5. Qualitative coding framework.
Appendix S6. Strength of evidence assessment details.
Appendix S7. Additional references.

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