A critical realist evaluation of advance care planning in care homes

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Funding information
This research received no specific grant from any funding agency in the public, commercial, or not-for-profit sectors.

Abstract
Aims: To evaluate care planning in advance of end-of-life care in care homes.
Design: A qualitative study.
Methods: Qualitative data were collected from January 2018–July 2019 (using focus groups and semi-structured interviews) from three care homes in the South West of England. The data were analysed using thematic analysis followed by Critical Realist Evaluation.
Results: Participants comprised of registered nurses (N = 4), care assistants (N = 8), bereaved relatives (N = 7), and domiciliary staff (N = 3). Although the importance of advance care planning was well recognized, the emotional labour of frequently engaging in discussions about death and dying was highlighted as a problem by some care home staff. It was evident that in some cases care home staff’s unmet emotional needs led them to rushing and avoiding discussions about death and dying with residents and relatives. A sparsity of mechanisms to support care home staff’s emotional needs was noted across all three care homes. Furthermore, a lack of training and knowledge appeared to inhibit care home staff’s ability to engage in meaningful care planning conversations with specific groups of residents such as those living with dementia. The lack of training was principally evident amongst non-registered care home staff and those with non-formal caring roles such as housekeeping.
Conclusion: There is a need for more focused education to support registered and non-registered care home staff to effectively engage in sensitive discussions about death and dying with residents. Furthermore, greater emotional support is necessary to help build workforce resilience and sustain change.
Impact: Knowledge generated from this study can be used to inform the design and development of future advance care planning interventions capable of supporting the delivery of high-quality end-of-life care in care homes.

KEYWORDS
dean and dying, long-term care, nursing home care, palliative care
End of Life Care (EoLC) in care homes is becoming the focus of growing attention due to the rapidly ageing population across developed countries (United Nations [UN], 2017). In Europe, care homes are increasingly becoming the most common places of death (Andreasen et al., 2019), and in the UK it is predicted that care homes will be the most common place of death by 2040 (Bone et al., 2018). In the UK, the generic term ‘care home’ describes both residential and nursing homes. Both of these types provide food and board, 24-h care cover and assistance where required with activities of daily living. In addition, nursing homes provide care by registered nurses (Laing & Buisson, 2016). Both provide EoLC.

Collectively, the care home sector in the UK provides care for approximately 418,000 people which represents 4% of the population aged over 65 and 15% aged over 85 (Laing & Buisson, 2016). Given age is synonymous with progressive chronic illnesses (World Health Organisation [WHO], 2017), more people are now living in care homes with multiple chronic conditions (multiple comorbidities), such as Dementia and frailty (Kingston et al., 2018).

It is therefore important that care homes staff are able to provide quality, person-centred care for residents approaching the end of their life often with multiple chronic conditions (Aasmul et al., 2018). Although the definition of end of life (EoL) differs between countries, is generally considered that people are probably to die in 12 months (National Health Service [NHS], 2018). Care provided during this period is termed EoLC which comforts and supports people who are dying from a progressive life-limiting condition (Fisher et al., 2000). Furthermore, EoLC should be person-centred which involves considering the persons wishes, values, beliefs and family circumstances; and incorporating these factors into their care (Fisher et al., 2000; Kitson et al., 2013). Advance care planning is the most common tool used by care home staff to support person centred EoLC (Spacey et al., 2020).

Although advance care planning practices can vary depending on the cultural and legal contexts of different countries (Sharp et al., 2013), efforts to standardize the definition have been made (Sudore et al., 2016). An international multidisciplinary panel described advance care planning as a process that supports individuals to share their personal values, life goals, and preferences about future care (Rietjens et al. 2017; Sudore et al., 2016). The advance planning process usually consists of a discussion between care home staff, the resident and/or their family, which is documented in an advance care plan (National Institute for Health & Care Excellence [NICE], 2019). Regardless of country, it has been well established that advance care planning can improve quality, reduce unnecessary admission to hospitals, and decrease stress and anxiety for both residents and their family members (Aasmul et al., 2018).

Thus, the care home staff delivering EoLC need to plan, gather and record information about a person’s wishes and preferences and take them into account to provide the best care possible for that person (Cloninger, 2011). Despite the importance of advance care planning in care homes, research has highlighted a lack of insight from the care home staff engaging in advance care planning, as well as the perceptions of family members (Spacey et al., 2018).

1 | THE STUDY

1.1 | Background

Given the increasing numbers of residents dying in care homes, and the complexities associated with advance care planning a plethora of interventions have been designed to support care home staff to engage in advance care planning discussions with residents and their family members (Braun & Zir, 2005; Cronfalk et al., 2015; Mayrhofer et al., 2016; McGlade et al., 2017; O’Sullivan et al., 2016; Temkin-Greener et al., 2017). The main focus of these interventions tends to be on educating staff on the importance of advance care planning, how to approach such conversations with residents and the legal and cultural factors associated with care planning (Sharp et al., 2013). For example, some residents may not want to discuss death and dying due to cultural or religious beliefs. Education also informs care home staff of The Mental Capacity Act (2005), and in cases where a resident lacks mental capacity, a legally appointed decision maker can engage in the care planning process on their behalf.

Both registered and non-registered care home staff are involved in advance care planning and are responsible for delivering patient-centred care for people at the end of their life (Leaman 2011). Despite this, existing studies have not explored to perceptions of both registered and non-registered care home staff engaging in care planning (Spacey et al., 2020). The UK care home workforce is made up of non-registered care assistants (76%) and registered nursing staff (12%); the remaining workforce is made up of ancillary staff such as housekeeping and administrative staff (Griffiths et al., 2019). Despite these complexities, current research tends to be outcome rather than process focused with the number of advance care plans in place for residents receiving EoLC being the most commonly reported marker of the quality care (Garden et al., 2016; Kinley et al., 2014). A recent systematic literature review evaluating advanced care planning interventions found that these outcomes were often short lived and that the numbers of advance care plans dropped post-intervention suggesting limited sustainability (Spacey et al., 2020). The reasons for this are currently unclear due to a sparsity of research exploring the underlying processes behind advance care planning in care homes. Specifically, current evidence about the rationale behind intervention design, the effect of context, and the response of stakeholders is sparse. This means that the impact of these factors on outcomes has thus far not been adequately established.

2 | THE STUDY

2.1 | Aims

This study aims to evaluate care planning in advance of end-of-life care in care homes. The following research questions were set to achieve this aim:
• What mechanisms currently exist to support care home staff to effectively engage in advance care planning? (intervention mechanisms)
• Can the context of care homes influence advance care planning? (contextual mechanisms)
• How do stakeholders tend to respond to advance care planning? (human agency)

2.2 | Design

This study reports on the results developed over the course of a two-phased study that adopted a Critical Realist design (Porter, 2015), which assumes that the outcomes of interventions result from the interaction of a plurality of causal mechanisms. Mechanisms are embedded in the intervention itself (intervention mechanisms); in the social and organizational context (contextual mechanisms); all of which are filtered through individuals’ interpretations and responses (Human agency; Porter, 2015). Thus, each of the above causal categories will be used to explore advance care planning in care homes. Table 1 details each of the causal categories.

Qualitative data was collected from January 2018–July 2019 using serial semi-structured interviews and focus groups to explore advance care planning in three care homes in the South West of England. Participants included bereaved relatives (N = 7), registered nurses (including managers; N = 4), care assistants (N = 8) and domiciliary/administrative staff (N = 3) who were all interviewed twice. All participants had been involved in advance care planning. The qualitative data were transcribed verbatim, analysed and results were reported in accordance with the Consolidated criteria for reporting Qualitative Research guidelines (Tong et al., 2007).

2.3 | Setting

The Care Quality Commission’s (CQC) electronic database was used to search for care homes in one county in the South West of England to include both residential and nursing homes. The CQC is an independent regulator of health and social care services in England. Purposive sampling was also used for the selection of data collection sites (Tongco, 2007) to ensure variation in the care homes settings.

Moreover, the local Care Commissioning Group (which are government run organizations responsible for commissioning most hospital and community services in England) provided advice throughout the process which was used to develop the following purposive sampling variables and eligibility criteria (Table 2):

| Outcome | Definition |
|---------|------------|
| Human agency | People’s experiences, interpretations and responses to the intervention and contextual mechanisms |
| Intervention mechanisms | The mechanisms contained in interventions which are designed to change the behaviour of those at whom they are targeted |
| Contextual mechanisms | The resources and restrictions embedded in the social and organizational context may inhibit or promote the effectiveness of intervention mechanisms |

Using the purposive sampling method and the eligibility criteria, five care homes were selected and approached. The first three care homes that agreed to participate that met the eligibility criteria were invited to take part in the study. Three variations of care homes were chosen because they represent the largest proportion of care homes types in the UK (Competition & Markets Authority [CMA], 2017; Laing & Buisson, 2016; see Table 3).

2.4 | Sample/Participants

The selection of participants was informed by two systematic literature reviews carried out by the authors (Spacey et al., 2018, 2020), which revealed that few studies reported on experiences of advance care planning from the perspectives of bereaved relatives, particularly those from a residential setting. Additionally, whilst current literature explored the experiences of registered care home staff, the perspectives on non-registered care home staff, especially those with non-formal caring roles were often overlooked (Spacey et al., 2020). After discussion with the participating care homes and the ethics committee, it was deemed inappropriate to include residents who were receiving EoLC in this study. Specifically, the researcher would not be aware of residents’ level of knowledge as to whether they knew they were receiving EoLC, so it was deemed insensitive and unethical to ask them about EoLC. However, the inclusion of bereaved relatives added a broader service user’s perspective. All included participants and their allocated pseudonyms are detailed in Table 4.

To aid staff recruitment, researchers attended staff meetings to inform the staff about the study and hand out invitation letters to potential participants. This enabled those who expressed an interest to contact the researcher independently of the care home manager. Bereaved relatives were contacted through managers using an invitation letter which included the researcher’s contact details. When the potential participants expressed an interest in taking part in the study, they were provided with participant
information sheets, and a consent form. No participants refused to participate.

### 2.5 Data collection

There were two phases of data collection meaning participants were interviewed twice. The questions used in phase one for the focus groups and semi-structured interviews were developed from two systematic reviews of the literature (Spacey et al., 2018, 2020). Phase two involved presenting the initial findings to the participants (from phase one) using focus groups and semi-structured interviews. Murray et al. (2009) found that the use of repeating or serial interviews helped to uncover the complexities of individual situations by allowing narratives to develop.

Both sets of focus groups with registered nurses and care assistants lasted 1 h and were conducted by two researchers, one acting...
as facilitator, the other as moderator. The semi-structured interviews (with bereaved relatives and care home managers) lasted up to 45 min. All the researchers participated in data collection, and had experience of qualitative interviewing and had no prior relationship with the participants.

Staff interviews and focus groups were conducted in private rooms in the participating care homes, and the interviews with bereaved relatives were carried out in their own homes or the University. All interviews were audio-recorded, and the moderator took notes during the focus groups. The participants were informed that we were researchers conducting a study exploring EoLC in care homes.

2.6 | Ethical considerations

Ethics approval was granted on the 15/01/2018, by the Health Research Authority, a UK government arm of the National Health Service providing governance of health research (Regional Ethics Committee reference 27/LO/1828).

2.7 | Data analysis

Analysis was carried out at the same time as data collection for each phase, and was discussed during fortnightly meetings of the research team; this helped determine sample size and data saturation. Throughout the process, analysis was triangulated by the four members of the team. A different mode of analysis was used for each of the two phases of focus groups and semi-structured interviews. The primary purpose of the first phase was to attain descriptive data about the use of advance care plans in care homes. The data gathered were thematically analysed (Braun & Clarke, 2006) by coding to identify recurring patterns. These patterns were arranged into themes and subthemes that augmented, refuted or refined the theoretical insights about intervention mechanisms, contextual mechanisms and human agency.

3.1 | Intervention mechanisms

This section examines the generative mechanisms embedded in interventions that were hypothesized to promote behaviour conducive to supporting more effective advance care planning. These consisted predominantly of educational initiatives.

3.1.1 | Education to support advance care planning

Education mechanisms were identified during both phases of this study and recognized as being key to providing care home staff with the skills and knowledge to effectively and sustainably carry out advance care planning discussions. Analysis suggested that shadowing (where colleagues would observe each other’s practice) was used by experienced care home staff (5 or more years’ experience) to train less experienced staff to engage in advance care planning discussions. Shadowing was used by both registered nurses and non-registered care assistants, enabling them to learn from each other how to approach and sensitively build up to discussions about death and dying.

Care assistant 2 C1: They can observe how I start the conversation and build up to some of the sensitive topics...I think this has to be experienced...seeing how others do it.

Registered nurse 1 C3: ...those who are new with less hands-on experience watch how I style certain questions...I talk them through it all afterwards as well...

Analysis of the data across all three care homes in this study suggested that shadowing was the most common education mechanism used to improve practice related to advance care planning and person-centred care. The data implied that care assistants found methods such as shadowing to be effective due to the ‘practical’ nature of their job rather than having more formal education.

Care assistant 1 C1: It’s very practical you just have to get it done we learn from watching each other on the job that’s how I learnt anyway... when we get new staff, we put them with the older staff so they can pick things up and learn.
However, shadowing was not the only educational method used to support advance care planning. Analysis of the transcripts from care home three uncovered a range of education mechanisms used by the care home staff from workshops to video sessions to accompany shadowing as part of the Gold Standard Framework for care homes (GSFCH) programme. The GSFCH is a framework used to support the early identification of residents approaching the end of their life, and to help them plan ahead (The Gold Standards Framework [GSF], 2018).

Registered nurse 1 C3: Training is provided through workshops and paperwork like the portfolio that come with the framework. We have video sessions that we watch throughout the year too...

Despite only one care home using these additional education strategies, several benefits were evident which appeared to be unique to the workshops and videos. Specifically, educational workshops tended to include more diverse groups of care home staff compared with shadowing. For example, it was evident that staff with non-formal caring roles such as housekeeping engaged in workshops and videos on advance care planning. This is an important finding as bereaved relatives valued staff with non-formal caring roles being part of their loved one’s care in terms of advance care planning. For example, bereaved relatives expressed developing good relationships with housekeeping staff helped them feel more comfortable and able to share information with them pertaining to their loved one’s care.

BR5 C2: I also dealt a lot with cleaning staff when they would come in the room, they made conversation with us, my aunt took a liking to one of them....

The section has highlighted some of the different education mechanisms employed in care homes to support staff engagement in advance care planning discussions. Data collected from bereaved relatives suggests that there may be some benefits to including a more diverse range of care home staff in advance care planning training.

3.2 | Contextual mechanisms

This section will explore the resources and restrictions embedded in the social and organizational context which were hypothesized as either promoting or inhibiting the effectiveness of advance care planning. Two significant contextual mechanisms were identified: financial variability between care homes and the diversity of residential populations.

3.2.1 | Financial variability in the care home sector

Analysis of the initial interview transcripts identified financial and organizational barriers to the implementation of education and training to support staff to engage in advance care planning. It was apparent that the type and size of care home influenced their financial and organizational ability to deliver and sustain intervention mechanisms. Specifically, care home one was a small residential home with 32 beds, which delivered EoLC a few times a year, while care home three was a 75-bed nursing home where EoLC was delivered far more frequently.

Care assistant 2 C1: We do not deliver much end of life care here, so we only have a few deaths a year. Usually quite a few in a short amount of time, then you’ll go through a period where lots die. It’s always Christmas before or after. So, when a death comes it can be hard hitting, we’re not as used to it as some other places may be.

It was acknowledged that the GSFCH provided useful education and training to help a more diverse range of care home staff engage in advance care planning. However, managers from care homes one and two spoke about the time commitments with setting up such training interventions, compounded by the implementation costs which were not viable in their smaller homes.

Manager C1: Lots of homes are signed up for the Gold Standards Framework then you have to update it and keep paying... we are a residential small home so would rather not be paying for training that may not work for us and that may be phased out soon. We cannot afford to waste money like that.

This data suggests that financial and organizational contexts are preventing these homes from benefiting from many of the supportive mechanisms that come with the GSFCH, that care home three was able to use. The manager from care home three highlighted that they had more financial freedom compared with other homes which allowed for flexibility in terms of training and education. However, the manager from care home three also highlighted that providing EoLC is an important priority that should not come down to financial resources.

Manager C3: We’ve got that flexibility and I just wish other homes had that. There is a financial thing in that. But you can’t put a price on end of life care; the memories are everything to a loved one, if you don't get it right you’ve had it.

3.2.2 | The diverse resident population

During phase two of the study care home staff expanded on the diverse resident population, highlighting some of their perceptions in regard to caring for residents living with dementia. Our findings indicate that some care home staff assumed residents with dementia would get too upset or would not understand if they talked about
death and dying. Therefore, to avoid upset and confusion, some care home staff would avoid sensitive discussions with residents living with dementia and discuss with the next of kin.

Care assistant 1 C2: ...for residents who have dementia we go to their families directly because it can be very upsetting, and it can confuse the resident even more if we start asking too much questions. It’s about supporting them to enjoy the time they have left.

The perception that talking about death and dying would confuse or upset residents living with the dementia was not only apparent in the quotes from care assistants. Managers spoke about the increasingly demanding and complex EoLC needs of residents. The manager from care home one spoke about how residents requiring EoLC frequently have multiple conditions such as frailty and dementia. The complex ‘high needs’ of residents living with dementia as well as their mental capacity was used to justify this behaviour.

Manager C1: Over the past few months a lot of our residents have become high needs and we haven’t got the capacity. So mostly it’s about discussing it with the next of kin and not the resident. You’ve got to think what is right for that person, so if they don’t understand what you’re talking about, why you going to talk to them about dying that’s not nice.

These findings imply that some care home staff’s perceptions prevented residents with dementia from being involved and contributing to their own care. Rather than attempt to engage or assume capacity, these staff assumed a lack of capacity and went straight to the family, excluding the resident living with dementia. Similar findings in regard to discussing death and dying as part of advance care planning were evident when analysing the bereaved relatives’ experiences. One bereaved relative explained that the care home staff went directly to her as the next of kin to ask about her husband’s preferences who was living with dementia.

BR2 C1: They wanted to know what my wishes were you know for him I didn’t mention any of this to him and I don’t think he knew I let them know.

3.3 | Human agency

Human agency represents how stakeholders interpret and respond to the identified intervention and contextual mechanisms. The most significant factor identified was the emotional difficulty that staff experienced in initiating discussions about death and dying.

3.3.1 Emotion and reluctance to discuss death and dying

It was evident that some care home staff found it emotionally difficult delivering EoLC to residents, especially towards the end stages of their life. The emotional labour of delivering EoLC was expressed by a range of care home staff from housekeeping, to care assistants to registered nurses and managers with their years of experience ranging from 2–29 years.

Care assistant 2 C1: In some ways you become their family because you’re spending more time with them than their family. Often families stay away as they don’t want to see their loved one dying so often you take their place. You try being their family for them. So, losing a resident like that can be very emotional yes.

Housekeeping C3: We get very emotional because we get attached to them, you can’t help that you’re human.

It was apparent staff’s emotional attachment to residents impacted on their ability to discuss death and dying as part of advanced care planning. Analysis implies that staff’s close attachments with residents led to them feeling uncomfortable discussing sensitive and potentially upsetting topics, such as death and dying. Despite the reluctance to discuss death and dying it was well understood by all care home staff that advance care planning was part of their ‘professional job’, thus it was not avoided. However, it was evident throughout the transcripts that discussions about death and dying as part of advance care planning were sometimes rushed and overlooked.

Manager C2: We often shy away because we feel uncomfortable… But we really don’t talk about that...we really don’t explore that. So, it makes us come across as a bit stilted and a bit stiff.

Care assistant 3 C1: You just have to try to put it to the back of your mind. Although it’s upsetting it is your professional job at the end of the day, you can’t let it affect your work and how you are.

However, these quotes suggest that it is more than just care home staff’s relationships with residents. This data uncovers a defence mechanism used by some care home staff which may be detrimental to residents and relatives. Specifically, the quotes imply that care home staff are avoiding discussions about death and dying, not only to prevent upset to residents, but to also protect themselves from upset.

This reluctance was noticed by bereaved relatives during phase 2 of the study. Specifically, they expressed how some staff would attempt to hide and avoid sensitive discussions about death and dying, pertaining to their own care and the care of other residents living in the care home.
been found in professionals who are regularly exposed to emotional
dying with residents as a defense mechanism. Similar behaviour has
suggested that some care home staff avoided the subject of death and
protecting their own emotional wellbeing. Specifically, analysis sug-
avoidance of conversations about death and dying was also about
was experienced by many care home staff.

Although, the findings from this present study support Hall and col-
& Patricia, 2019). However, only a few studies report on how care
home staff's personal backgrounds and cultures can influence how
proceeded in discussions about death and dying when delivering EoLC. Despite this, most
literature advocates for supporting care home staff to manage the
cultural needs and backgrounds of residents receiving EoLC (Givler
and Patricia, 2019). However, only a few studies report on how care
home staff’s personal backgrounds and cultures can influence how
they approach death and dying in the care home (Hall et al., 2011;
Xiao et al., 2017). For example, Hall et al. (2011) highlighted a lack of
confidence in staff from different cultures towards death and dying.
Although, the findings from this present study support Hall and col-
leagues it was found that reluctance to talk about death and dying
was experienced by many care home staff.

There was evidence to suggest that some care home staff’s
avoidance of conversations about death and dying was also about
protecting their own emotional wellbeing. Specifically, analysis sug-
gested that some care home staff avoided the subject of death and
dying with residents as a defense mechanism. Similar behaviour has
been found in professionals who are regularly exposed to emotional
and traumatic experiences (Drury et al., 2014). It has been found that
being continually exposed to traumatic experiences without support
can lead to compassion fatigue, which is described as a state of men-
tal exhaustion triggered by a decreased ability to cope with everyday
settings (Baranowsky & Gentry, 1999). Findings from this study sug-
gest that care home staff's way of managing compassion fatigue was
to avoid sensitive subjects with residents and relatives about death
and dying. However, this behaviour (avoidance) appeared detrimental
to care home staff's ability to engage in meaningful advance care
planning conversations with residents and relatives.

Care home staff’s reluctance to talk about death and dying was
particularly apparent with residents with dementia. Some care home
staff went straight to the next of kin to develop the advance care
care homes (Alzheimer's Research UK, 2020), but because research
has found that residents who are cognitively impaired such as those
living with dementia have a greater chance of being unnecessarily
admitted to hospital at the EoL (Perrels et al., 2014). These findings
suggest that need to provide care home staff with the knowledge to
be able to sensitively engage in discussions about death and dying
with a diverse range of residents, including those with dementia.

Despite the need for more education and support, findings sug-
gest that a lack of resources inhibited change. For example, despite
the apparent lack of issues in the care home workforce, there
was lack of educational focus on emotionally supporting staff to
build resilience around advance care planning. This is important
as care homes across the UK and Europe are struggling with lim-
ited resources, increasing demands and expectations (European
Commission, 2018; The King’s Fund, 2018). However, while costs
need to be taken into account and minimized, the evidence pre-
vented in this study highlights important quality issues in the provi-
sion of advance care planning which justify the need for care homes
to incorporate greater support for their workforce into their cost
base.

4 | DISCUSSION

The findings of this study showed that although the importance of
advance care planning was well recognized, the emotional labour
of frequently engaging in discussions about death and dying was a
problem for some care home staff. In some cases, care home staff’s
unmet emotional needs led them to rushing and avoiding discussions
about death and dying with residents and relatives. Despite these
difficulties, our findings indicated a sparsity of mechanisms to sup-
port care home staff’s emotional needs. In addition, a lack of train-
ing and knowledge, particularly amongst non-registered care home
staff and those with non-formal caring roles, appeared to inhibit
their ability to engage in meaningful care planning conversations
with those living with dementia due to preconceived assumptions
and communication barriers.

Whilst the numbers of advance care plans in place are import-
ant (Finucane et al., 2013; Kinley et al., 2014; O’Sullivan et al., 2016;
Temkin-Greener et al., 2017), findings presented in this study sug-
gest that more needs to be done to support staff to sensitively en-
gage in discussions about death and dying to improve the quality of
advance care planning discussions.

Although current education mechanisms such as shadowing
helped staff learn the practicalities of engaging in discussions
about death and dying (such as how to approach the subject and
what questions to ask), it was evident more needs to be done to sup-
port the emotional needs of care home staff. Specifically, findings
identified that some care home staff were reluctant to talk about
dead and dying. Our findings indicated that care home staff’s back-
grounds, cultures and personal experiences influenced how they ap-
proached death and dying when delivering EoLC. Despite this, most
literature advocates for supporting care home staff to manage the
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to care home staff’s ability to engage in meaningful advance care
planning conversations with residents and relatives.

Care home staff’s reluctance to talk about death and dying was
particularly apparent with residents with dementia. Some care home
staff went straight to the next of kin to develop the advance care
plan as they believed that residents with dementia would not un-
derstand and would become too upset if they discussed death and
dying with them. This poses implications for practice not only be-
cause of the rising numbers of residents living with dementia in UK
care homes (Alzheimer’s Research UK, 2020), but because research
has found that residents who are cognitively impaired such as those
living with dementia have a greater chance of being unnecessarily
admitted to hospital at the EoL (Perrels et al., 2014). These findings
suggest that need to provide care home staff with the knowledge to
be able to sensitively engage in discussions about death and dying
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sion of advance care planning which justify the need for care homes
to incorporate greater support for their workforce into their cost
base.

4.1 | Limitations

The study did not include residents receiving EoLC. This is recog-
nized as a limitation given that service users’ perspectives and ex-
periences are often overlooked (Spacey et al., 2020). Nonetheless,
bereaved relatives were included in the study which provides val-
uable serve user insight whilst also acting as a proxy for residents
themselves. The number of care homes involved in the study
was also a limitation, three care home were selected, limiting the
transferability and generalizability of the findings. However, these
homes were carefully selected to represent the main care home
types in England. This study did not explore the implementation of
any specific intervention to support advance care planning, there-
fore inferences about causality cannot be made. Nonetheless, our
findings have highlighted some of the processes which may influence the effectiveness of care planning in advance of EoLC in care homes.

5 | CONCLUSION

Findings provide care home staff, policy makers, regulators and researchers with a deeper insight into what needs to be done to support advance planning in care homes. For example, findings from our study suggest that more research and education initiatives need to go into supporting care home staff emotionally to improve their ability to engage in discussions about death and dying as part of advance care planning.

In the longer term, the findings generated in this study can be used to inform the development of future interventions designed to support care home staff engage in advance care planning. However, further research is recommended to replicate the study in different contexts such as rural care home settings to assess the generalizability of the findings.

CONFLICT OF INTEREST

No conflict of interest has been declared by the author(s).

AUTHOR CONTRIBUTIONS

All authors have agreed on the final version and meet at least one of the following criteria (recommended by the ICMJE*): (1) substantial contributions to conception and design, acquisition of data, or analysis and interpretation of data; (2) drafting the article or revising it critically for important intellectual content.

PEER REVIEW

The peer review history for this article is available at https://publons.com/publon/10.1111/jan.14822.

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How to cite this article: Spacey A, Scammell J, Board M, Porter S. A critical realist evaluation of advance care planning in care homes. *J Adv Nurs*. 2021;77:2774–2784. https://doi.org/10.1111/jan.14822
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