People with dementia and family carers are welcoming of a model of dementia palliative care, but sceptical of its implementation


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Abstract

Introduction: A palliative care approach can improve quality-of-life for people with dementia. It is the preference of many people with dementia to remain living at home until death, with the appropriate care. To develop a successful model for dementia palliative care in the community, it is essential to assimilate the perspectives and experiences of those affected. The guiding research question for this study was: What are people with dementia and family carers’ views on a model for dementia palliative care?

Methods: Focus groups (n=3) were conducted with bereaved or current family carers (n=11), and people with dementia (n=2). Discussions centred around a proposed model of dementia palliative care. These were transcribed and analysed using thematic analysis.

Results: Three main themes were identified: living and dying well with dementia; reducing carer burden to fulfil the wish for home care; and lack of faith in the healthcare system. An overarching theme which unified the analysis was: “Dementia palliative care is a dream, but not a reality.” This reflected participants’ repeated “wish” for this “ideal” model of care, but simultaneous scepticism regarding its implementation, based on their prior experiences of healthcare services.

Conclusion: All participants were welcoming of the proposed model for dementia palliative care and were generally positive about palliative care as a concept relating to dementia. There was consensus that the model would allow people to live and die well with dementia, and reducing the carer burden would fulfil the wish for home care. However systemic changes in the healthcare system will be needed to facilitate a truly person-centred, holistic, individualised and flexible model of care.

Keywords: Dementia, Palliative Care, Advance Care Planning, End-of-life, Home Care, Community Care, Model.
**Introduction**

Dementia is a syndrome with an ever-growing population. It is estimated that 55 million people globally have dementia, with nearly 10 million new cases every year, and it’s the seventh leading cause of death in the world (World Health Organization, 2023). Dementia is caused by many different diseases and affects each person differently. Cognitive and non-cognitive symptoms include memory loss, reasoning and communication challenges, personality changes and a reduction in the ability to complete daily activities (National Institute for Health and Care Excellence, 2018).

An approach originally associated with cancer, palliative care can improve quality-of-life for all people with life-limiting illnesses, and their families, by addressing physical, psychosocial and spiritual needs (World Health Organization, 2020). While it is now recognised that people with dementia can benefit from palliative care (Fox et al., 2020), they don’t have equal access to effective palliative and end-of-life care (Dempsey et al., 2015; Harrison Dening, 2016). The European Association of Palliative Care (EAPC) white paper (van der Steen et al., 2014) on optimal palliative care for older people with dementia highlighted key domains: applicability of palliative care; person-centred care, communication and shared decision-making; optimal treatment of symptoms and providing comfort; setting care goals and advance planning; continuity of care; psychosocial and spiritual support; family care and involvement; education of the health care team; societal/ethical issues; prognostication and timely recognition of dying; avoiding overly aggressive, burdensome or futile treatment (van der Steen et al., 2014). Core components of a model for palliative care for people with dementia living at home have also been put forward by service providers (Fox et al., 2020).

A study of bereaved carers (Mogan et al., 2022) highlighted the disparity between the consensus on the core aspects of end-of-life care (Poole et al., 2018) and peoples' lived experience. Carers who had looked after a person with dementia at home during the last six months of life experienced: poor continuity of care, lack of expertise, limited advance care planning, and a loss of autonomy (Mogan et al., 2022). The importance of autonomy and
treating the person with dementia with dignity and respect has emerged in several studies involving people with dementia and families about their perspectives on end-of-life care (Davies et al., 2017; Dening et al., 2013; Mogan et al., 2022; Russell et al., 2008).

People with dementia and their family carers concur that certain components are essential to end-of-life care: being cared for in place, being comfortable at the end-of-life, and having a skilled care team (Poole et al., 2018), with these views mapping to the EAPC domains. However, some differences of opinion exist (Poole et al., 2018). In a qualitative study of people with early-stage dementia and current or bereaved carers, carers put more value on future planning, prioritised decision-making on daily care, and felt poorly equipped to manage end-of-life care. On the other hand, people with dementia attached little value and importance to future planning, prioritised decision-making on medical care, and assumed their family would be capable of coordinating their end-of-life care (Poole et al., 2018).

People with dementia may also have limited insight into the existing burden on carers, viewing burden as something that might happen in the future (Dening et al., 2013). Family carers need support around end-of-life as they manage the challenging dual role of being a grieving carer and a decision-maker (Hennings et al., 2010; Mogan et al., 2022).

Generally, the preference of people with dementia and family carers is to be supported to live and die at home, with family carers concerned about the person with dementia being left alone or isolated in other settings, and keen for the person with dementia to have opportunities for social engagement and physical contact (Kupeli et al., 2016; Seow et al., 2023). While family carers generally agree with healthcare professionals on what's important for end-of-life care for people with advanced dementia, an additional theme identified by family carers was for end-of-life care to be provided “at home” or in a “home-like environment” (Kupeli et al., 2016). From the family carer’s perspective, a suitable physical and social environment is key to achieving an adequate quality-of-life for the person with dementia (Russell et al., 2008).
To fully address the needs of palliative care for people with dementia, a unified, interdisciplinary and evidence-based approach is required that links from the local to national and global levels, informed by priorities and guided by frameworks and models of care (Fox et al., 2017). The Model for Dementia Palliative Care project sets out to develop an acceptable, evidence- and practice-based model for palliative care for people with dementia living in the community in Ireland (Developing a Palliative Care Model for People with Dementia in the Community, n.d.). A draft model was developed using a programme theory approach and was presented as a logic model linking programme inputs to components and intermediate and long-term outcomes, while taking contextual and external factors into account and ensuring all elements translated into a model relevant to the community setting (paper in preparation). An important aspect of model development involved sharing drafts with stakeholders, including groups of people with dementia and family carers, to obtain critical feedback and iterate the model design. The guiding research question for engagement with these groups was: What are your views on our model for dementia palliative care?

Methods

Design. A qualitative design was employed, using in-depth focus groups comprising people living with dementia and family carers. The discussion was framed in the context of the draft model for dementia palliative care in Ireland. The draft model was shared with participants in advance by email. During the focus groups, following initial rapport building questions, the six components of the model were presented: comprehensive care, person-centred practice and care, integrated care, accessible care, care for carer/supporter, and end-of-life care (more fully explained in Table 1). Each component was discussed sequentially with the key principles narrated by the facilitator (SF), along with examples of what the component might look like in practice (see detailed excerpt from one of the six components in Table 2). After
each component was presented by the facilitator, an open discussion ensued, allowing participants to share their views on each component. The conduct and reporting of this study adhered to the COREQ guidelines (Tong et al., 2017).

### Sampling and recruitment.
Inclusion criteria were broad: a person living with dementia, or a (current or previous) family member of a person living with dementia. As recruitment required potential participants to read a plain language study information sheet, and opt-in to the research, it was unlikely that people with advanced dementia would take part, although they were not precluded from doing so. Using convenience sampling, participants were recruited through the Alzheimer Society of Ireland (ASI). A study poster and information sheet were circulated via email to members who expressed an interest in hearing about research studies. A list of those expressing interest was gathered by the ASI and shared with the research team who then made direct contact, answered any further questions, and arranged times for participation.

### Data collection.
All focus groups were conducted by a female senior postdoctoral researcher experienced in dementia and qualitative research (SF). The focus group discussions took place online during March 2022 via videoconferencing software and were digitally recorded with permission. The audio data was transcribed verbatim with potential identifier information removed to uphold anonymity. Files were saved to an encrypted, password protected drive to ensure confidentiality.
A total of 13 participants took part in three focus groups (Table 3). Participants were given numerous dates, and selected to take part in the focus group at the time most convenient for them. The focus groups opened with general introductions and discussion to build rapport. Focus groups lasted between 91-106 minutes (average 96 minutes). Two of the focus groups included a person with dementia (both with a diagnosis of young onset dementia, and both at early stage), while the other 11 participants were family carers. The family carer relationship to the person with dementia was either daughter (n=8) or wife (n=3). In some cases, the family carer’s loved one had passed away, and in others they were alive and either living with the family carer or in long-term care. Of the sample, 92% were female, with only one male participant who was a person with dementia. All participants were resident in Ireland, living across various urban and rural settings. Participants were not required to have prior knowledge or experience of palliative care to be eligible for inclusion but were asked about this at the beginning of the focus groups. Participant’s familiarity with palliative care varied from no prior experience to one participant who previously worked as a paediatric palliative care nurse.

Data analysis. Qualitative data was coded and analysed in accordance with Braun and Clark’s (Braun & Clarke, 2013) seven stages of thematic analysis: 1. Transcription, 2. Reading and familiarisation, 3. Coding, 4. Searching for themes, 5. Reviewing themes, 6. Defining and naming themes, 7. Writing – final analysis. The process was managed with MicroSoft Word and Excel software packages. The transcriptions were first read through to gain familiarisation with the content, with brief written memos used to record items of potential interest. Coding of the full dataset was undertaken independently by two researchers (SF and MF) before comparing the proposed codes. Codes relevant to the research question were applied, at a semantic level. The coding was then used to develop
provisional themes and sub-themes, with particular attention given to identifying both positive and negative examples for each theme. The researchers consulted regularly, and after an iterative review of codes, themes and sub-themes, the final themes were defined before proceeding to completion of the analysis and final write-up.

**Ethical considerations.** Appropriate guidelines for conducting psychosocial research with people with dementia were followed, including the Alzheimer Europe position paper (Gove et al., 2018). Written informed consent was obtained from all participants. Ethical approval was granted by the local Research Ethics Committee (ref: 2019-077).

**Results**

Overall, three main themes and nine sub-themes were identified in the dataset, relating to our guiding research question: What are people with dementia and family carers’ views on a model for dementia palliative care? The overarching theme unifying the analysis was: “Dementia palliative care is a dream, but not a reality.” This underscores participants' repeated language around the “wish” for this “ideal” model of care, but their dubiousness that it could be implemented based on their poor prior experiences of services. A thematic map is presented in Figure 1.

<<<Figure 1>>>  

**Theme 1: Living and Dying Well with Dementia**

Palliative care as applicable in dementia
All participants were welcoming of the proposed model, agreeing that it would allow people with dementia and their families to live and die well, and were generally positive about palliative care as a concept relating to dementia. One bereaved carer (Family Carer-6), whose husband had dementia, experienced specialist palliative care in a psychiatric hospital for the final six months of his life and described the care as “wonderful”. Others with experience of specialist palliative care through family members, or those with professional experience, recognised the importance of early palliative care for an improved end-of-life experience.

Participants said that the model represented what they wanted and needed and would “do anything” to make it happen. Some described it as a “wish list” for ideal care. One participant said it was upsetting to see the model written down as it is exactly what they wanted, but so removed from their actual experience.

While participants felt that palliative care is applicable to dementia, they perceived a wider stigma in society. They felt a wider public understanding of palliative care and openness to talking about death and dying would benefit everyone, including those living with dementia.

The wish for the core tenets of dementia palliative care to be actualised

Participants felt the person-centred care focus of the palliative care approach for dementia would greatly improve overall care. Participants agreed that the ultimate goal of dementia care, and in particular dementia palliative care, was to have the best quality-of-life possible, with their personhood respected:

“It’s a lot of behavioural issues Mom has, she doesn’t need a nurse, a doctor, she needs someone who cares and is going to put music on and rub her head and calm her down when she’s stressed and love her and that’s what she needs.” Family Carer-3

Relatedly, the holistic principles of the model were welcomed, in particular consideration of non-physical needs. A common discussion was the need for counselling for the person with
dementia and/or their family members. Some family members found that while counselling was accessible to them, it was less accessible for the person with dementia. Additional to formal counselling, participants also felt that health and social care practitioners (HSCPs) should take a more holistic approach to care; simply asking them about how they were feeling and coping, would greatly improve their experience and outcomes.

“I think just have somebody to listen to [the person with dementia] so they could get information on how they felt and give them the time to do that, and sometimes people just rush in and out of appointments, they don’t really talk to them and nobody really listens to them” Family Carer-7

Regarding other core tenets, the groups agreed that an effective model of dementia palliative care needs to be individualised and flexible.

“If you’ve a row of twenty houses, and you have twenty people with Alzheimer’s in every house, every single one of those people is going to need different care, different hours, different times, and they can have the same dementia, be the same age, but their stages will be different, their needs will be different, their carers needs will be different and it has to be individual care tailored to everyone” Person with Dementia-2

The need for care and settings that are appropriate to the individual’s personality and preferences were discussed:

“[my mother] isn’t suddenly going to start doing day care because it wouldn’t have been her personality whereas my dad did enjoy that now” Family Carer-10

Participants recounted the negative impact of receiving care that was inflexible and misaligned to their needs, such as home carers arriving early in the morning when the person with dementia was still asleep and who was then negatively affected for the day after being woken. Carers who were working full-time were exasperated with the inflexibility and poor communication channels of the healthcare system:
“when my mum was in hospital...I’d be asking how she was and what could I do for her and you’d be speaking to the nurses and [they’d say] ‘oh the doctors will be doing their rounds at 11:00 o’clock if you want to pop in’, like you can’t just walk out of work and pop into the hospital on the off chance that somebody might be there to talk to you” Family Carer-7

Participants identified a significant gap in current services as their unavailability during evenings and weekends, which can preclude continued care at home, particularly in advanced dementia. This left people feeling unsupported and precipitated potentially avoidable crisis events. Avoiding the emergency department is particularly important to participants with experience of advanced illness.

“Personally, I would like to really ensure that these weekend elements of it is gone, that it is 24/7 because you know not being able to contact the GP on a Friday evening…it just bugs me that people think you know nothing happens on Saturday and Sunday” Family Carer-2

Importance of preparing for a comfortable and dignified death

There was a consensus that while advance care planning (ACP) can be difficult and uncomfortable, it is hugely beneficial:

“We took it on board early in the time the care planning and it has taken a load off of my mind, it has taken a load off my wife’s mind, it’s taken a load off my kid’s mind. I think it’s of vital importance, the earlier the better because…it’s the people that matter made the decisions with me, and they know what I want, and I can’t stress how important that is”

Person with Dementia-2

Many participants had never been engaged in a discussion about ACP by a HSCP, only learning about different care choices around end-of-life at a late stage.

“I think people aren’t aware of the choice of not going to hospital, for somebody with dementia at the end because for us (husband’s name) went to hospital in February and it
was then that the doctors raised it with us, you know, would you think about not coming back and I thought, oh my God, of course I would like I didn’t know, I didn’t know that I could think about not coming back” Family Carer-6

Bereaved carers added that education around what to expect at end-of-life earlier in the course of the illness would have greatly helped them cope at end-of-life. Carers valued honesty around end-of-life, and HSCPs being forthright about what was happening. Knowing how much time the person had left ensures that the family have the chance to say goodbye.

“on the end-of-life care and the dignified death I think it’s very important that we know that it’s the end-of-life and that you know, people are honest with us” Family Carer-7

Fear of pain was a common thread in the conversations. Many participants expressed worry about pain at end-of-life for themselves or their loved one, especially as they feared a person with dementia can’t express their pain at advanced stages. One bereaved carer shared her experience:

“it still took a couple of weeks before the people in the nursing home recognised that she needed pain treatment i.e. morphine and I kind of wish that she could have had that morphine maybe two months beforehand and things would have been a lot easier for her and for the family as well” Family Carer-7

More positive experiences tended to be those where there was better support including palliative care involvement:

“we were so blessed that the GP associated with the nursing home was fantastic and … was very clear about the main issue … that she wouldn’t be in pain, I mean my mum was nonverbal for the, you know, the last two years of her life, so I mean literally you’re trying to judge from somebody’s eyes if they’re in pain and it’s really, really difficult and uncomfortable but I have to say … the nursing home was used to palliative care and … they dealt with it amazingly well” Family Carer-10
Key aspects of good end-of-life experience included being in a familiar, comfortable, and private environment; surrounded by loved ones; with their personhood and dignity respected:

“at the end of the day, all we all want just to be hopefully in our own homes but if that's not possible that we're comfortable, that we have a palliative care team there with us and to help our families and ourselves have a peaceful exit because there's nothing more than we want. We just want it to be as less dramatic as possible, we want to have conversations, we want to have laughter, we want to have sometimes music, sometimes storytelling, sometimes just kindness” Person with Dementia-1

Although assisted dying was not included in the model, it was raised as a key issue in two of the three focus groups. Participants in this group, including a person with dementia, were open to conversations around assisted dying and felt that it should be included in a model of dementia palliative care. Participants felt that it should be a choice for them and having the option would give them a sense of control over their illness, and alleviate worries about pain.

**Theme 2. Reducing carer burden to fulfil the wish for home care**

**Carers need their own support to facilitate dementia palliative care at home**

Carers spoke about the immense toll of caring for their loved ones with dementia. One person with dementia stated that the carers “have it a lot tougher than the person with dementia” Person with Dementia-2). Carers made huge sacrifices such as moving home with young families from abroad, moving in with their loved one, or quitting work to care full-time. Many of the carers had little or no support; one spousal carer had only one “night off” in the previous three years. One carer living with her husband with a rare and progressive type of young onset dementia starkly outlined her “sacrifice” as a carer:

“you would be made to feel guilty if you weren't prepared to sacrifice your life and look after your spouse...I’m no longer living with the person I married, it’s a completely different person, I don't know him, you know, and yet I have to do everything for him, he can’t eat, he
can't go to the loo, he can't take his tablets, he can't, you know, get up from a chair, everything, without my help and yet there's no recognition of that sacrifice" Family Carer-9

Carers felt aggrieved by a healthcare system that didn’t consider their needs or provide the support they needed to care for their loved ones. They pointed out that carers are as individual as each person with dementia. For example, to facilitate them caring at home they need some respite, but at a time and duration that allows them to truly engage in something meaningful to them, to do “anything other than dementia” or “switch off”.

Carers wanted to be the ones primarily caring for their loved one, but they felt to do this, they too needed care and support. Having overnight support was also important for carers wanting to support the person with dementia at home for end-of-life.

Some participants expressed a need to balance what the person with dementia wants and what the carer needs, particularly in the context of home care.

“it’s a battle between mom’s needs and what she deserves for her and what we're able to do long term” Family Carer-3

One participant expressed her frustration at having no home support or GP support, and being left with no option but to bring her mother to hospital during a crisis, where the following exchange occurred:

“I brought my mother into A&E because I honestly didn’t know what else to do, and I was in a state, she was in a state...and then a doctor came up to me and said 'd'you know the best place to care for someone with dementia is in the home?' [laughs ironically] I just, and she had been living with me and I wasn’t- I didn’t bring her there because I didn’t want to care for her at home, I brought her there because I didn’t know what else to do!” Family Carer-7

There was a deep frustration that their care and sacrifice was not seen and not valued:
“it’s only a pittance anyway, you know and if you look at cost of me saying, ok, I’m too ill to look after him which I probably soon will be if it carries on, you know, and dump him at a hospital door that’s going to cost an awful lot more” Family Carer-9

Home care is preferred, but there needs to be other options

Overwhelmingly participants were supportive of a model which facilitates home care, and most family carers wanted to care for their loved ones at home, if possible:

“my mom was an amazing mom of 11 and she was a nurse herself and I just can't imagine putting her anywhere else because she deserves to be at home.” Family Carer-3

However, it appeared that an overemphasis on home care can lead to feelings of grief and failure if the person cannot be cared for at home for any reason:

“If you’ve an overemphasis on doing everything at home it can lead into a sense of, I suppose, failure you know when the person can’t be at home, and I think there’s various reasons why somebody with or without dementia…why a person can't be at home” Family Carer-6

Overall nursing homes were seen as sub-optimal care settings, and this perceived lack of suitable alternatives put pressure on carers to continue caring at home even when this was becoming unsustainable. Feelings of fear, regret and failure were common at the experience, or idea of, “putting” one’s loved one in a nursing home.

“nobody wants their loved one to be in a nursing home, particularly if it’s not something that they wanted themselves but sometimes it's just not possible for the person with dementia to still be safely at home” Family Carer-1

This carer also spoke about the very positive experience her father had in a specialist dementia long-term care facility.
“I left in tears when I dropped him in because I thought I'd just betrayed him, I thought it was the worst in the world that I have done for him but actually he had a great time and I've seen people who are on respite there, they've had a lovely time, people who haven't enjoyed it at all … let’s maybe not throw out some of the existing models because it can work for people” Family Carer-1

Another reason for endorsing home care was the accessibility it offered. Participants spoke about the difficulty in physically accessing services, needing to arrange time off work, transport, getting bloods done prior to an appointment. They would greatly prefer if care could come to them at home (e.g. nurse come to the house to take blood samples).

Complex grief before and after death of the person with dementia

Participants spoke of the often traumatic, slow decline in dementia, as one bereaved carer put it:

“I felt like I was losing [my mother] in the last few years of her life and it was kind of death by a thousand cuts, it was really like heart-breaking" Family Carer-10.

It can be very difficult for carers who’ve devoted a significant time of their life to caring for someone to accept they are at end-of-life. One participant had found the following analogy helpful, likening the person’s decline towards end-of-life to “letting go, not giving up”.

It was evident in discussions that the carers who felt supported in their caring role were better able to cope following the death of their loved one. Carers who are entirely consumed with caring with no formal support can face a huge adjustment after the death of the person with dementia:

“I struggled as well because I kept my job my full-time job while I was looking after my mum getting home care and…a lot of people are tempted to give up their work and I would say
don't, do it if you can, if you could possibly manage without it because it's important because after you know after the person's gone” Family Carer-7

Good palliative care at end-of-life is important not only for the person with dementia, but also for the family as a bad experience can leave a carer with lasting trauma:

“If the person with dementia doesn't have a good journey to the end the person that's left has to deal with that…you don't you don't want to be thinking back and thinking, oh I wish I could have done this or I wish we'd known this” Family Carer-7

Those who had good palliative care support around end-of-life seemed less likely to suffer from complicated grief:

"when I knew that my mother didn't have long I was at peace in some ways because I just knew all I had to do was be with her and make sure she got the pain relief she needed” Family Carer-7

**Theme 3: Lack of faith in the healthcare system**

**Disillusionment with the healthcare system**

While participants generally welcomed a model for dementia palliative care, they were disillusioned by their experience of the healthcare system thus far and were doubtful that the system could change to accommodate the model:

[regarding the dementia model of care]: “on paper looks fantastic but oh my god how you’re going to get to implement it I don't know. Instead of adding they're taking from it” Person with Dementia-2

A small number of participants had been involved with earlier initiatives, such as dementia training programmes, or dementia guidelines, and expressed frustration that these initiatives, while they were a source of great promotion and excitement, sometimes were not rolled out, or else failed to have much impact.
“The talking has got to stop and we’ve got implement these things” Person with Dementia-1

A few participants compared their experience of palliative care for loved ones with cancer with their experience of dementia care and were pessimistic about whether palliative care would be applied to them or their loved ones with dementia in the same way.

“if I say I have cancer I, I know from my husband, would get every piece of care under the rising sun. I mean, I could not have asked for better healthcare than he got. But I say that about my people the people that I know that are at advanced stage of Alzheimer's and you know, no, they don't get the care, the families don't get the care” Person with Dementia-1

**The fight for dementia services in a fragmented healthcare system**

A prominent theme underlying much of the focus group discussions was the “fight”, “struggle” or “battle” for dementia services. Geographic disparities were apparent.

“[Dementia services are] really on a demand basis, and by demand, I mean the people who shout the loudest and who are most persistent tend to be rewarded if it's available in their area” Family Carer-1

Another carer spoke of only getting services as she “played the system” (Family Carer-5).

Others felt that a situation must become very dire before help is offered:

“the prompt for [services] to get involved in the case of my mother, for example, was when she became a problem to them or to the system or to society” Family Carer-7

Participants were jaded and fatigued by their “fight”:

[regarding looking for services to keep the person at home] “I [am] fighting all the time, sometimes I can't fight. I'm tired and all of us get tired of fighting.” Family Carer-3

The word “lucky” was used a lot when participants talked about services they did have, with some participants appearing embarrassed or apologetic mentioning that they had any
services, knowing that others had no support at all. Another participant expressed that their
father was “lucky” with the timing of his illness because there was support available as he
neared end-of-life, whereas another family struggled to get support for their mother:

“It’s all about timing and also I know my mother would go and do whatever we felt as family
needed. I know she would just she loves us all and she’d do, she’d say, oh…I'll be grand, but
it's not fair. It's not fair that dad got to die at home and he got he got all the support he
needed. It's just not fair. She's amazing.” Family Carer-3

Some felt that cost was a personal barrier to them accessing services, but more felt that
even if they were willing to pay privately for services, suitable care services are not available.
Participants were also sceptical of how elements of the model could work, when their
experience was that many HSCPs in existing services are under-skilled in dementia care.
Some recounted unsatisfactory experiences with GPs and hospital staff. Commonly, the
group’s experience of home care staff was that they were not sufficiently trained in dementia
care.

“we are paying about two and a half grand a week for carers and the carers are not as good
as family. The reality is they don't seem to have experience with Alzheimer's or with
dementia” Family Carer-3

Some worried that unskilled care workers might have such a negative impact they may make
the person with dementia “worse”. Even where there were good care workers who the
person with dementia liked, they might lose them after a hospital stay when they are
assigned to someone else.

**Fight for information**

Finding information was also framed as a “fight”. Participants often had to find information on
their own, and sometimes came upon very useful information about a support or service only
by chance. Carers, even those with backgrounds in healthcare themselves, found it very
difficult to navigate the system in relation to dementia. They said that HSCPs rarely or never
came to them with information about supports or services, that they had to figure this out on
their own, adding to the stress they were experiencing.

"from a dementia point of view it’s still quite fragmented … no one person has all the
information" Family Carer-5

Families would appreciate having a "one stop shop" or a named HSCP who knows them to
coordinate their care, signpost them to relevant services, and provide information as needed,
as is proposed in the model:

“The single point of contact is very important to have one person to coordinate everything”
Family Carer-5

Others elaborated that this should be proactive management, where the contact person
follows up with them. Families felt that improved information sharing between healthcare
services also needed to be communicated with them:

“like it's very easy to say all these things, but getting those group of people together, a GP
should be communicating with the MDT [multidisciplinary team], and the MDT should be
communicating with palliative care, palliative care should so who is who is the person that's
going to communicate with all these people and are the family going to get that
communication back to them or is it always just going to them communicating often, some
office, some are saying which is what I often heard in the hospital, oh we’re having an MDT
meeting about that [laughs] we’re having an MDT meeting about that. Great, could you tell
us about what happened in that MDT meeting at some stage?” Family Carer-3

Family carers’ role as information providers was also framed as a fight; carers felt that they
are a valuable source of information, but that they are not being listened to by HSCPs. Thus,
while they welcomed the carer as being placed centrally in the model, they expressed some
doubt as to how this would work in practice:
“[the model] mentions here as well that the carer and supporters’ insights is valuable but how does that fit in with, you know, the medical profession and how they interact with us and confidentiality” Family Carer-7

There was also frustration expressed by many that the healthcare system is very fragmented with no facility to transfer information. Some worried that advance care plans or advance healthcare directives may not be followed in different settings.

“then you go to different hospitals than you’re originally in and if they weren't your notes, they don’t have the notes, they're starting from scratch” Family Carer-3

Discussion

This study provides a detailed account of people with dementia and their families’ views on a proposed model of palliative care for people with dementia living at home in the community. The overarching theme was dementia palliative care is a dream, but not a reality. People with dementia and their family carers agreed that the core tenets of palliative care and dementia care, including care that is person-centred and holistic, delivered in an individualised and flexible way, represents their ideal care model. However, their unsatisfactory prior experiences of healthcare services led to scepticism that this model could be implemented.

Both family carers and people with dementia want good quality care at end-of-life including being cared for in place, being comfortable at end-of-life, and having a skilled healthcare team, which is consistent with views expressed in a previous study (Dening et al., 2013; Poole et al., 2018) and in the EAPC framework (van der Steen et al., 2014). Family carers want their loved one’s individual needs met, with personalised care that fosters respect and dignity (Davies et al., 2017). Themes of dignity and respect were common in previous literature (Davies et al., 2017; Dening et al., 2013; Mogan et al., 2022; Russell et al., 2008). In contrast to an earlier publication (Dening et al., 2013), people with dementia who
participated here were acutely aware of the impact of their illness on their family carers, perhaps as these people had young onset dementia and were also involved in advocacy whereas the previous study included people with late onset dementia.

Engaging in ACP appeared to bring peace of mind to people with dementia and their families; previous research has associated ACP with better end-of-life outcomes (Dixon et al., 2018). However, as identified elsewhere (Mogan et al., 2022), many people with dementia and families had rarely been engaged in ACP by a HSCP or had limited knowledge of ACP (Seow et al., 2023) and there are barriers to undertaking ACP which need to be addressed, with the help of HSCPs (Dickinson et al., 2013). Fear of pain at end-of-life was common, and compounded by worries about the inability to communicate pain to others. This is consistent with other research which shows how important it is for families that their loved ones are pain-free, with carers feeling the only way to maintain control and independence was through assisted dying and euthanasia, but if end-of-life care was better, they would not need to contemplate euthanasia (Russell et al., 2008). Notably, assisted dying was discussed openly and frankly in our focus groups. This may have been influenced by current debate in Ireland around this issue, following the establishment of the Joint Committee on Assisted Dying in 2023. A model which would facilitate ACP, pain and symptom management and support grieving carers was greatly welcomed.

Literature supports the position that the preference of most people with dementia and their carers is for them to stay living at home in the community until end-of-life (Kupeli et al., 2016; Russell et al., 2008; Seow et al., 2023). As consistently outlined in the literature (Mogan et al., 2022; Russell et al., 2008), carers can experience immense burden while caring for loved ones, especially at end-of-life. This stress can be exacerbated when the views of people with dementia and family carers diverge (Poole et al., 2018), which reinforces the importance of supporting the family carer with care decisions. Carers felt they needed formal support to provide care at home. There was a feeling of pressure to provide care, with emotions of guilt, sadness and grief, when carers were not supported to do so, or they made a choice to move.
the person with dementia to another setting such as a nursing home, which was typically seen as a less preferable option.

While largely positive and welcoming of a model for dementia palliative care, participants were doubtful that it could be implemented. They were disillusioned by a healthcare system which made promises that were not kept and plans that weren’t implemented. This theme is consistent with a UK study, which found people with dementia and family carers had a lack of trust in medical decision-making (Dening et al., 2013). Participants had to fight for services, a theme also found in other research which highlights family carers’ struggle to navigate “unfamiliar territory” (10, p.122), struggling with their lack of knowledge and experience, and poor communication with professionals; family carers had to fight for information and did not feel listened to, which was also a theme in other studies (Mogan et al., 2022; Smith et al., 2021). In Ireland, private home care is available, however even carers who could afford this found it unsatisfactory. Carers felt that HSCPs, particularly home care workers, were under-skilled in dementia care, as reported elsewhere (Mogan et al., 2022), manifesting in a lack of trust.

**Conclusion**

In this study, focus groups were conducted to answer the research question, what are people with dementia and family carers’ views on a model for dementia palliative care? Following analysis of a rich discussion, we can summarise that this cohort concur that dementia palliative care is very important and valuable, but they are disillusioned by their experience of the healthcare system and somewhat doubtful that a new model of care which is truly person-centred, individualised, and flexible, could become a reality. While consistent components of a model for dementia palliative care for people living at home have been agreed, cultural and systemic changes in the healthcare system and among the public more generally are needed to facilitate the implementation of a model of dementia palliative care that is acceptable to those who are directly affected by dementia.
Abbreviations

ACP  Advance Care Planning
ASI  Alzheimer Society of Ireland
DPC  Dementia Palliative Care
EAPC  European Association of Palliative Care
GP  General Practitioner
HSCP  Health and Social Care Practitioner
MDT  Multi-Disciplinary Team

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Ethical Approval

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Availability of Research Materials

Raw data (i.e. interview transcripts) are not publicly available due to their potentially identifiable nature.
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