Living and dying with dementia in Wales: Barriers to care

Report
February 2015
## Contents

<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Foreword</td>
<td>3</td>
</tr>
<tr>
<td>Introduction</td>
<td>4</td>
</tr>
<tr>
<td>Theme one: Identification and planning</td>
<td>8</td>
</tr>
<tr>
<td>Theme two: Inequality of access</td>
<td>11</td>
</tr>
<tr>
<td>Theme three: Quality of care</td>
<td>17</td>
</tr>
<tr>
<td>Conclusion</td>
<td>22</td>
</tr>
<tr>
<td>Looking forward</td>
<td>23</td>
</tr>
<tr>
<td>References</td>
<td>24</td>
</tr>
</tbody>
</table>
Foreword

Dementia is a growing issue and concern for societies across the world. Our ageing population means that dementia will affect an increasing number of people in the coming decades. It represents one of the major health and social care challenges and therefore presents a major challenge facing Wales and the UK.

While much of the focus of research and discussions is rightly on the challenge of finding cures for the different forms of dementia, we must be realistic and acknowledge that the eradication of dementia is, at present, a medium to longer term ambition rather than a short term goal. Therefore we must place equal emphasis on the care which is provided to people with dementia and the support which is available to their families and carers.

We hope that this report makes a contribution to how we address the challenge of improving care and support available to people with dementia at the end of life as well as their families. By focusing on the barriers we can begin to improve that care and ensure all those who might benefit from it receive it.

As a charity working to improve end of life care and strengthen the focus on dementia, we are committed to working with all those organisations which provide support and care to people in the final stages of the disease. This report sets out a series of principles which we believe should underpin high quality end of life care, no matter where it is delivered. We do recognise the successes that have been achieved in Wales in supporting people with dementia, but we also highlight areas where improvements are needed. To that end, we have set out a number of recommendations which we believe would help contribute to better care for people at the end of life with dementia.

Our goal must be to ensure that everyone, regardless of their condition, where they live, or the services they use, gets the best possible care and support at the end of life.

Dr Jane Collins
Chief Executive, Marie Curie Cancer Care
February 2015
Introduction

The G8’s dementia summit\(^1\) held in London in December 2013 shone a global spotlight on dementia and in particular the challenge faced in the UK and globally by the projected soaring increase in dementia cases over the coming decades\(^2\).

Much of the focus of the summit and the resulting media coverage centred on the search for a ‘cure’ for dementia. However, given that dementia is a collection of different diseases, with different causes, the reality is that there can be no single cure for dementia. This means that we are looking for cures rather than a cure and that is a much longer term goal than may initially be imagined. So, it is important that we place at least as much emphasis on caring for people with dementia – now and in future – as we do on finding cures.

The Welsh Government has supported a number of initiatives aimed at meeting the challenges associated with caring for rising numbers of people with dementia. The National Dementia Vision for Wales\(^3\) sets out the specifically Welsh aspects of this challenge, such as dementia’s effect on rural communities and the communication needs of people whose first language is Welsh. In keeping with the vision laid out here, about 20 communities in Wales are now working towards making society more dementia friendly and enabling people with dementia to live well for as long as they can. The Dementia Friends programme is also active, with approximately 6,500 in Wales to date\(^4\).

There has, however, been much less focus on the inevitable conclusion of what is a progressive and terminal condition. The National Dementia Vision includes no reference to the end of life care needs of people with dementia. Together for Mental Health: A Strategy for Mental Health and Wellbeing in Wales, through which much of the Welsh Government’s approach to dementia is focused, makes only a general commitment to ‘Supporting the optimum state of wellbeing right through to the end of life for older people and those with dementia’\(^5\). Beyond exceptions such as Alzheimer’s Society’s My life until the end: Dying well with dementia\(^6\) and the work of the National Council for Palliative Care (NCPC), the final stages of the disease remain the forgotten aspect of what has been called a ‘silent epidemic’\(^7\).

This report explores how we can ensure that there is a stronger focus on dementia as a terminal condition and in particular the barriers which prevent many people with dementia from accessing and receiving appropriate high quality care at the end of their lives. There are three key barriers:

1) identification and planning
2) inequality of access
3) the quality of care experienced by people with dementia

This report considers how we can improve the care and support for people who are living and dying with dementia, as well as their families and carers, and also how we can reach those health and social care professionals who care for and support them. It focuses on the situation and improvements needed in Wales, but draws on developments and research from across the UK.
What is dementia?

Dementia does not have one single cause. It is caused by a range of underlying conditions including Alzheimer’s, vascular dementia, dementia with Lewy bodies and fronto-temporal dementia. Throughout the remainder of this paper, dementia is used as shorthand for this broad range of conditions.

Dementia is a progressive neurodegenerative disease. This means that the symptoms become more severe over time. Symptoms usually include memory loss, confusion, speech difficulties, mood and behavioural changes and problems with self-care and other aspects of daily living. There are some treatments to alleviate the symptoms, but dementia is not curable and is a terminal illness.

The final stage of dementia is comparable with other terminal illnesses as the prognosis is often similar. While we argue that it is important to focus on the latter stages of the condition, we recognise that planning and intervention in the earlier stages are also essential to ensure a good death for a person with dementia. Better planning and support in the early stages of dementia can improve informed decision making in the latter stages. This applies also to those with dementia who may die from another condition.

It is also important that we acknowledge that the reality for many people with dementia is that they will be living with different conditions, including cancer and those associated with frailty and older age.

Dementia: The numbers

Around 850,000 people live with dementia in the UK, 45,000 of whom are Welsh residents. The vast majority (almost 95%) of those living with dementia are over the age of 65. There are, however, approximately 2,500 people under 65 in Wales who have dementia. Dementia also has an interesting gender profile: two thirds of people with dementia are women. The latest data suggests that dementia or Alzheimer’s is now the biggest single cause of death amongst women in England and Wales, having surpassed different forms of cancer for the first time. Alzheimer’s Society provides further key data regarding current and future trends in their report Dementia UK: Second Edition.

Providing definitive figures for the number of people with dementia is problematic because diagnosis is difficult, particularly in the early stages. Similarly it is difficult to identify the number of people dying as a result of dementia. Many deaths are not directly attributed to dementia, but rather to the immediate cause of death, such as an infection or another coexisting condition. In Wales in 2013, 1,555 women and 724 men had their death attributed to dementia or Alzheimer’s disease.

Between 2003 and 2013 the percentage of deaths from dementia and Alzheimer’s disease in England and Wales rose from 4.7% to 12.2% for women and from 2% to 6.2% for men. A recent cohort study in Northern Ireland concluded that:

‘The cause of death documented for [Alzheimer’s disease] patients may be affected by the physician’s knowledge of the patient or reflects the approach to management of patients with end-stage dementia.'
The likelihood is that, across the UK, the number of deaths from dementia is currently being significantly under-reported and these official figures are not a reflection of the real picture. For example, a significant number will be recorded as being caused by ‘senility’.

Dementia is therefore an important issue at the end of life for a growing number of people in the UK. According to the 2011 census, there were approximately 8.5 million people living in the UK over the age of 65. These individuals can be considered to be at a higher risk of developing dementia.

This figure is set to rise in the future because the UK has a rapidly ageing population. Between 2001 and 2011, the proportion of the UK population aged 65 and over increased from 15.9% to 16.3%. By 2030, the proportion of the UK aged over 85 is expected to double. Xie, Brayne et al estimate that one in three people over the age of 65 years who die will have some form of dementia.

As the proportion of the population aged 65 and over increases, so the number of people at risk of developing dementia increases. By 2021, the number of people with dementia across Wales is projected to increase by 31% and by as much as 44% in some rural areas. It is essential that we address the issue of end of life care for people with dementia now.

**Dementia: A long and unpredictable condition**

It is important to underline the unpredictable nature of dementia. People with dementia are also likely to live longer than those with some other life-limiting conditions. The period from diagnosis to death varies, depending on what condition has caused the dementia symptoms, but all have a relatively long disease trajectory.

People with Alzheimer’s have been known to live for up to 10 years after the initial diagnosis, although this depends on the age when the diagnosis is received. People diagnosed with fronto-temporal dementia can expect to live between six and eight years, while people with dementia with Lewy bodies can live between six and 12 years. Those diagnosed with vascular dementia can expect to live a shorter length of time after diagnosis – approximately five years.

The length of time a person can expect to live with dementia also depends on the stage that the disease is at when they are diagnosed. Dementia’s relatively long and unpredictable disease trajectory (see Figure 1) makes it difficult for doctors to provide an end of life prognosis – and perhaps makes doctors unwilling to do so. A study of nursing home deaths found that most people with dementia were given a prognosis of more than six months, but 71% of these people died within the six month period.
The period for which a person with dementia might require care is long in comparison with a number of other terminal illnesses. People with dementia also often struggle with daily living and taking care of themselves at an earlier stage than those with some other conditions due to the nature of the symptoms. Many people with dementia will find themselves living in care homes. We must therefore consider how well social care and palliative care services are coordinated and aligned in different settings across the country so that an individual can routinely access palliative care, regardless of where they live.

The progressive and degenerative nature of dementia, coupled with current and projected levels of dementia diagnosis, mean that it is essential that we explore how we can make all relevant health and social care services more dementia friendly and appropriate, particularly for those approaching the end of life.

**What prevents people with dementia accessing high quality end of life care?**

There is a growing evidence base which suggests that people with dementia are not being appropriately identified for end of life care, and that they have less access to, and receive poorer quality, care than people with other terminal illnesses. These three barriers extend through a person’s dementia journey, from before receiving a formal diagnosis through to accessing appropriate care in the last days of life.

It is only by addressing the barriers people with dementia face in these three areas that we can ensure that there is a stronger focus on the needs and experiences of people living and dying with dementia, and their families and carers.
Theme one: Identification and planning

Barriers: Lack of timely and appropriate diagnosis

Dementia is often difficult to diagnose in its early stages and people who have signs of the condition may be slow to seek help. In the earliest stages, symptoms such as confusion and memory loss may be dismissed by people with dementia, families and professionals as normal signs of ageing.

A formal diagnosis from a doctor or other health professional is generally required for people to access many services, and this restricts the support people receive if there is any delay in them receiving a diagnosis. There is no general diagnostic test for any type of dementia. Diagnosis currently involves looking at an individual’s clinical history, cognitive testing and possibly neuro-imaging.

The two key issues here are:

- Reluctance or fear on the part of people with dementia to identify or acknowledge symptoms;
- The ability of health and social care professionals to accurately identify and diagnose dementia, particularly in its earliest stages.

Rates of diagnosis vary significantly across the UK and within the individual nations. In Wales it is estimated that the average rate of dementia diagnosis is 42.9%, with an estimated 25,000 people living with dementia without a diagnosis. This compares to a diagnosis rate of 64.8% in Northern Ireland.

Timely diagnosis has a major potential impact on subsequent access to support, treatment and appropriate care.

Barriers: Lack of recognition of dementia as a terminal condition

There appears to be a failure to identify dementia as a terminal illness in some quarters; for example, by health and social care professionals. This has implications for the type of care which is provided for people with dementia. Indeed, most people with dementia have, historically, not had access to specialist palliative care.

In 2004, the World Health Organisation (WHO) found that less than 1% of people in hospices in Europe had a primary diagnosis of dementia. If dementia is more routinely identified as a terminal diagnosis, health and social care professionals will be able to provide more appropriate support and care to people with dementia.

Barriers: Failure to identify dementia as a cause of death

This issue of poor identification is also reflected in the rate at which dementia is recorded as the cause of death on death certificates. The ultimate cause of death is often attributed to an infection or other common illness and is stated as such when the death is formally registered.
This creates a culture in which dementia is not recognised as a cause of death in its own right, potentially reinforcing the earlier lack of recognition of dementia as a terminal condition. This means that dementia is often overlooked despite the fact that it may have played a key role in a person’s decline and ultimate death.

For example, where a person with dementia loses the ability to swallow and therefore to eat and drink, their body, without the use of artificial feeding, will slowly begin to close down and they will ultimately die at the point at which their organs fail. In some instances, this death might be simply recorded as being as a result of heart or multiple organ failure and not the progress of dementia which precipitated the ultimate decline.

The number of deaths with a mention of dementia recorded on the death certificate almost doubled in England between 2001 and 2010, rising from 6.6% to 12.0%. In Wales there was a smaller increase in the proportion of deaths recorded as having a cause of vascular or unspecified dementia or Alzheimer’s disease over the same time period, from 3.3% to 4.7%. This rose to 7.4% in 2011 following changes to practices for coding causes of death. As Sleeman et al argue ‘Improved death certification is essential to fully understand place of death in dementia’. If there is to be an accurate understanding of dementia for people at the end of life in Wales, it is important that these improvements in recording practices continue.

**Barriers: Ineffective advance care planning**

Once a person is identified as having a terminal illness, advance care planning is crucial. In theory it should be particularly important for people with dementia and other degenerative cognitive conditions, whose diagnosis means that they will increasingly lose capacity as their condition progresses. However, the reality is that for many people with dementia, advance care planning can be extremely problematic. A study from the US indicates that people with dementia were less likely to have advance care directives than people with other conditions. This is supported by evidence from the UK.

It is difficult to judge when advance care planning discussions should be initiated. Delayed diagnosis may mean that even in the earliest stages, people with dementia may not have retained the mental capacity required to make advance decisions. Even in the earliest stages of dementia, people can struggle to imagine their future self and, understandably, frame their views according to their current situation. Further, it is suggested that preferences change as the condition progresses.

Finally, many people with dementia in the early stages may be unwilling to accept that their condition will result in death and this can disrupt conversations over planning for this outcome. This means that it is often left to family members to plan. The families of people with dementia are supported in their decision making by their ‘knowledge of the person with dementia’s previous views, clear diagnostic information, and family support’.

The wishes of a terminally ill person can also be expressed in the form of advance directives or Living Wills. In England and Wales, an advance directive is the only legally binding form of expressing views on care and is situation specific.

In the absence of an advance directive, families may express the wishes of the person with dementia. This is known as a proxy directive or a general values directive; however, these two forms of directive are not legally binding.
A person with dementia may also nominate a ‘welfare attorney’ or assign lasting power of attorney. This gives the attorney the power to make decisions based on the current situation. Attorneys should be specified well in advance of loss of competence. The alternative to advance care planning is the identification of someone trusted by the patient who can represent their wishes.

In the absence of advance instructions, decision making ultimately sits with healthcare professionals, who must ensure that “the treatment has to be necessary either to save life or to prevent a serious deterioration in health and must be in the patient’s best interests”. In such instances, families can express their wishes regarding treatment, but ultimately any decision is the responsibility of the healthcare professionals involved. This can create conflicts between health professionals and families; although in reality the wishes of family and care givers will usually be discussed in a calm atmosphere with the relevant health professionals and a consensus reached on the best way to proceed.
Theme two: Inequality of access

Barriers: Access to palliative care

As has been discussed, for people with dementia to receive timely palliative care and holistic support, identification and recognition that they have dementia is a crucial first step. For some people, once dementia has been diagnosed, the next step might ideally be a referral to a palliative care service.

However, a recent hospital-based study in an acute medical ward in the UK suggested that people with dementia were less likely to be referred to palliative care and less likely to receive palliative medication than people who did not have dementia – just ‘17% of patients with dementia that subsequently died were referred to the palliative care team during their admission’.\textsuperscript{51}

This echoes the findings of a recent study in Scotland which found that just 20% of those with frailty or dementia had been formally identified as requiring palliative care, compared to 75% of those with cancer\textsuperscript{52}.

Data from the NHS Wales Informatics Service shows just 3.8% of people who died in Wales in 2012 with dementia as their underlying cause of death were recorded as having a palliative care diagnosis or having received hospital-based specialist palliative care, which is appropriate for people with more complex needs\textsuperscript{53}.

This data chimes with the widely held perception that palliative care services have historically failed to reach a significant number of people with dementia. The reasons for this may be explained by an acknowledgement that the roots of the modern hospice and palliative care movements developed in response to the needs of people with cancer. Additionally, dementia is a condition which is primarily addressed through social care, and many people with dementia live in care homes.

The National Institute for Health and Clinical Excellence (NICE) has produced guidelines on dementia care including palliative and end of life care for people with dementia\textsuperscript{54}. The guidance is clear:

‘Dementia care should incorporate a palliative care approach from the time of diagnosis until death. The aim should be to support the quality of life of people with dementia and to enable them to die with dignity and in the place of their choosing, while also supporting carers during their bereavement, which may both anticipate and follow death.

Health and social care professionals working with people with dementia and their carers should adopt a palliative care approach. They should consider physical, psychological, social and spiritual needs to maximise the quality of life of the person with dementia and their family.

Palliative care professionals, other health and social care professionals, and commissioners should ensure that people with dementia who are dying
have the same access to palliative care services as those without dementia’.  

Importantly, palliative care is identified as a valid and important treatment approach for individuals with dementia from the point of diagnosis. The authors of the Behavioural and Pain BEPAID study advocate the early introduction of palliative care for people with dementia.

The European Association for Palliative Care’s White paper offers a model (see Figure 2) which plots the changing care goals and priorities throughout the course of the dementia.

**Figure 2. Importance of palliative care in dementia and dementia severity**

It is therefore vital that we enhance the role of palliative care for people with dementia, not just in the last few days of life, but much earlier in the development of the condition because of its emphasis on symptom management.

We need to ensure that examples of excellent links between local palliative care services and care homes and hospital services are replicated across the country and extended over a much longer time period. We also need to ensure that people with dementia in the last days of life, whether they are in their own home, their care home or in hospital, can access palliative care specialists who then support colleagues to assess and manage pain.

It is important to consider what prevents many people with dementia accessing palliative care. The links between oncology departments and palliative care teams are well established. However it is not clear that the links between dementia specialists and palliative care teams are similarly well developed across all parts of the country. There may be additional barriers in the divide between health and social care. As previously noted, decision making about the move from active treatment to palliative care is complex. For people with dementia there is usually no single point equivalent to the conversation which someone with cancer may have with an oncologist about the transition from active treatment to palliative care. The degenerative nature of dementia should make the transition to palliative care easier; however, the data suggests that this is not the case. We have to
find a way to better identify people with dementia at the point at which they need support from and will benefit most from palliative care teams.

Importantly it has been suggested that access to palliative care should be based on the patient’s need, not their life expectancy, as it can offer huge benefits even for those who may have a less imminent expectation of death\textsuperscript{58}.

**Barriers: Access to hospice care**

The picture regarding access to palliative care for people with dementia that we have presented so far is highlighted further by the fact that in the UK it is still unusual for people with dementia to die in a hospice\textsuperscript{59}. ONS figures for deaths in Wales in 2012 show just one woman and no men whose deaths were categorised as caused by ‘mental and behavioural disorders’ (the category in which deaths from vascular and unspecified dementia should be recorded) died in a hospice\textsuperscript{60}. This reflects the wider picture of hospice use which is still predominantly used by people with cancer.

It is important that we explore why hospice use is so low for people with a dementia diagnosis. Is it because hospices are unsuitable places for people with dementia, or are there other reasons which explain the low take up of both hospice and palliative care? Do people with cancer represent the majority of those using hospices simply because oncologists are in the habit of referring those for whom there is no further active treatment to hospices and palliative care teams? Do we need to establish the same sort of relationships between other condition-specific specialists and hospices and palliative care teams? In some cases, care homes may be performing the same role as hospices.

Given the different nature of dementia and divergent needs of those with the condition, it is perhaps not surprising that there are relatively few dementia specialists working in palliative care settings\textsuperscript{61}. As we have already noted, the modern hospice movement and the palliative care specialism grew out of caring for people with cancer. People with dementia have very different needs to those with cancer and caring for them requires very different skills.

In addition to issues relating to referrals, there are also questions about the spaces and layouts of hospices and whether these need to be different to meet the needs of people with dementia. It is important to note that for some people with dementia, the move to a hospice (or any other unfamiliar care setting) can cause unnecessary confusion and distress at a stage in the condition when the person may be less able to cope with change. They may therefore not be considered an appropriate environment for care. We also need to explore what more can be done to build connections between hospices and social care.

To address the current under-use of both hospice and specialist palliative care, one key message that we need to send out is that hospices are open and available for people with dementia and that staff and volunteers have the training and skills needed to deliver services to people with dementia. This will require commissioners to prioritise end of life care focused on dementia and resource it fully.

We must also learn from those hospices which are already providing high quality care for people with dementia.
Barriers: Access to funding

Access to funding for those approaching the end of life is complex.

As their condition deteriorates, people with dementia will have increasing health and social care requirements; they will need greater help with personal care as well as specific healthcare and hospital-based services related to their dementia.

While it is clear that people with later stage dementia will have considerable health needs, this is not reflected in their access to Continuing NHS Health Care (CHC). CHC is the system whereby the NHS picks up the costs of caring for someone whose primary needs are for healthcare in a non-hospital setting. It is suggested that disproportionately low numbers of people with dementia are receiving CHC.²²

A report by the Wales Audit Office in 2013 noted concerns expressed by Alzheimer’s Society and Age Concern that:

‘Because of differences in the DST [Decision Support Tool] used in Wales and England, it may be more difficult for some people in Wales, most notably those with dementia, to meet CHC eligibility criteria.’²³

Following a Welsh Government consultation on the national framework for CHC in Wales, the same screening tools and DST as used in England have been adopted. As a result, the needs level in the DST now extends to ‘severe’ level needs for the cognitive domain, giving this domain parity with others.

However, it is likely that many of the same issues raised about the effectiveness of CHC in England are also true for Wales. The All Party Parliamentary Group (APPG) on Parkinson’s Failing to Care report found that better data collection is needed to determine patterns and inequities of access to CHC for people with conditions like dementia.²⁴

The APPG on Parkinson’s analysis also highlights fundamental problems with assessment of applications for CHC funding. First, while there is a nationally recognised decision support tool and a national framework which sets out an assessment process in England, there is no compulsion to use it. Consequently, there are different local assessment approaches. This ensures that the system effectively has an in-built inequality.

Second, the guidance and the tools are often interpreted in a way that does not reflect the reality of different conditions, including cognitive degenerative conditions such as dementia, Parkinson’s or motor neurone disease. The focus of the assessment is the wording of the tool, rather than the needs of the individual:

‘It is not used as a support but being interpreted legalistically. The focus of the assessor is on the semantics of each word rather than the essence of the patient’s condition. It is often used literally and not interpreted intelligently.’²⁵

This can lead to perverse decisions:
'Parkinson’s UK had been given an example of where someone with Parkinson’s in receipt of NHS continuing care had undergone a reassessment. Since the previous assessment, the individual’s mobility had deteriorated to the extent that they were now permanently bed bound. The use of the DST then gave the individual a lower score in the mobility category because they were at a reduced risk of falling. This subsequently rendered the individual ineligible for NHS continuing care.'

Third, there is inconsistency in the way that assessments are carried out. For example, assessments do not always take into account the level of medication a person is receiving and may not consider the person in an unmedicated state.

Fourth, there is the question of who carries out the assessment. Is the individual charged with undertaking the assessment the most appropriate person to do so? Are they known to the person with dementia and do they understand the different manifestations of their condition? There are clear issues relating to training and understanding of specific conditions and involvement of appropriate healthcare professionals.

It is important that the tools and guidance which are used to assess CHC eligibility reflect the realities of different conditions, including cognitive degenerative conditions such as dementia. The whole CHC system arguably needs a radical overhaul to make it dementia-friendly.

**Barriers: Discrimination**

Alzheimer’s Society has previously argued that individuals with dementia face discrimination in the care they are given because there is a lack of targeted services and care standards vary. Alzheimer’s Society argues that those with dementia may experience “poor rates of diagnosis and a lack of appropriate services” because it occurs at a later stage in life and is assumed to just be a symptom of getting old.

It has also been suggested that those with dementia have poorer access to mental health services compared with younger people with mental health conditions. Older people with dementia may, therefore, not be getting optimum care simply because of their age. Dementia affects people’s ability to defend themselves against discrimination, so issues of age discrimination may be more pronounced.

Younger people with dementia may also experience disadvantage in terms of access to services. Most people with dementia are aged over 65. This means services are targeted at older individuals and there could be a risk of younger people with dementia being excluded.

In Wales, there are approximately 2,500 people living with early-onset dementia aged between 45 and 65. Specialist services for younger people with dementia are rare, and dementia services targeted at older people may not accept them – which means it can be difficult for younger people to access support. Dementia services targeted at young people are needed so that these people get the support they require and are not excluded from services and discriminated against.

Connolly et al have suggested that people with dementia who are from ethnic minority groups might face a ‘double disadvantage’. Marie Curie commissioned research by Kings College London into the Palliative and end of life care for Black, Asian and Minority Ethnic (BAME) groups.
in the UK\textsuperscript{74} and has published an accompanying policy paper\textsuperscript{75} which explores these issues further. This work underlines the predicted growth in demand for end of life care from BAME communities over the next two decades.
Theme three: Quality of care

Barriers: Inconsistency in care standards in hospitals

ONS figures\(^7\) suggest that the numbers of people with dementia who die in hospital are similar for Wales as in England, where the figure is almost two in five (39.6\%)\(^7\). The National Care of the Dying Audit for Hospitals underlines the considerable variation in care on offer for people with different diagnoses in hospitals across England\(^8\). Similar information is not currently available for Wales; a comparable audit could reveal important information about care for the significant number of people who die with dementia in hospital each year.

In 2009, it was estimated that people with dementia over 65 years of age accounted for 25\% of hospital beds at any one time. When admitted to hospital because of another condition, they spend longer in hospital than those with the same condition who do not have dementia\(^9\). Data from the NHS Wales Informative Service (NWIS) shows that people who died in 2012 with dementia as their underlying cause of death spent more than 50 days of their final year of life in a hospital bed\(^1\). This is considerably higher than for other causes of death, such as cancer and chronic lower respiratory diseases.

Hospitals can be incredibly difficult places for people with dementia. In 2013, a Royal Society of Psychiatrists audit of hospitals in England and Wales highlighted low performance against essential standards for many hospitals in the care of people with dementia\(^1\). Headline findings for hospitals in Wales included that very few Executive Boards were regularly reviewing information on readmission rates, delayed discharges, and in-hospital falls relating to people with dementia, and important elements of mental health assessments were not being routinely carried out\(^1\). Furthermore, organisational checklists provided by Welsh hospitals showed that no mandatory provision of dementia awareness training to doctors, nurses and healthcare assistants was in place (although 88\% had been provided with training in the previous 12 months).

A key problem appears to be the inadequate identification of dementia when people with dementia are admitted to hospital. The Care Quality Commission (CQC) review of hospital episode data in England support this conclusion, finding that in 29\% of cases a person’s dementia was not recorded, despite having been recorded during previous admissions\(^1\). As the CQC note:

> ‘An NHS Confederation briefing on dementia showed that hospitals often code the primary reason for admission to hospital, not the dementia. Not identifying, and then not coding and recording, the dementia leads to hospitals underestimating the numbers of people with dementia occupying beds.’\(^1\)

Barriers: Inappropriate hospital admissions

Lack of training and awareness of end of life care among health and social care professionals, staff shortages and the non-availability of GPs (particularly out of hours) can all result in people with
dementia being admitted to hospital at the first ‘sign of decline’. Evidence from England shows that people with dementia in care homes are more likely to experience inappropriate admissions to hospital than people without the disease. Inappropriate admissions can cause unnecessary disruption for a person with dementia in the final days and weeks of life and they may end up not dying in their preferred place or in the most appropriate place for them.

Studies have shown that unplanned admission to acute hospitals also negatively impacts on survival times for people with dementia, which were half those without dementia.

This underlines the need for good care planning and excellent primary care and community support to help avoid unnecessary hospital admissions for people with dementia. In many instances this may include a role for external medical expertise, such as community nurses, going into the home to support care and nursing professionals at key stages.

It is important that we learn from those many care homes which are providing high quality care, working well with other professionals locally and able to deal with what are often the complex needs of people with dementia at the end of life.

In Wales in 2012, figures suggest that over half of deaths with mental and behavioural disorders as the underlying cause (of which the majority are attributed to vascular and unspecified dementia) occurred in a care home; however, nearly 4 out of 10 occurred in hospitals:

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As with many other conditions, most people with dementia would like to die at home or in their usual place of residence, for example a care home, and their families would also prefer this. However, a significant proportion of people with dementia are likely to die in hospital. Spending periods of time in hospital can be extremely distressing for people with dementia. Research has shown that those admitted to hospital with dementia are more likely to die during an admission and in the six months after their hospital admission.

**Barriers: Lack of continuity of care**

Continuity of care is extremely important as changes in health or social care staff can create unnecessary distress. It is clear that familiarity benefits both the person with the dementia (and their family) and those providing care, in helping people to stay in their own homes for longer. Similarly, unnecessary transfers from an individual’s usual place of residence can have a disconcerting impact, causing confusion and exacerbating uncertainty.
Evidence from England has highlighted issues relating to poor information exchanges between different providers as people with dementia moved between services. In 27% of care homes and 22% it found:

> ‘aspects of variable or poor care regarding the arrangements for how they shared information when people moved between services’.

Work is also needed to ensure that nursing and care home staff, social care providers and local GPs are confident about working with people with dementia and are able to take the steps required to prevent unnecessary admissions, which lead to breakdowns in continuity and place the individual at greater risk of decline.

**Barriers: Poor pain management**

Dementia may not in itself cause pain, but as most people with dementia are over 65 they may be suffering from conditions such as arthritis or cardiac disease which do. Researchers have shown that as dementia progresses, people become less able to identify pain:

> ‘Compared with the control group, people with dementia were less able to identify painful situations and used fewer categories to define their concept of pain’.

They conclude that clinicians should bear this in mind when assessing people with dementia for pain.

Health professionals tend to have to rely on close observation of changes in people’s behaviour to identify the level of pain that they are experiencing and manage pain correctly. As the disease progresses, reporting of pain by the person generally decreases and becomes less accurate. Studies have underlined the degree to which people with dementia in hospital are perceived to be in pain in the last few days of life:

> ‘50% of patients were noted by clinicians to be in pain in the last 48 h (hours) of life. Mean pain score throughout admission for these that died was 2.69 on the PAINADii, in comparison to 1.49 for those who were discharged. Of those that died, 45% of carers were a little dissatisfied/dissatisfied with the overall quality of care provided on the ward’.

Managing pain is more complicated for people with dementia and this means that people with dementia are more likely to experience inadequate pain management than people with some other terminal conditions.

It is also important that we consider whether the conceptions of pain which are used in palliative and end of life care are largely rooted in conditions such as cancer and whether we need to explore if it is more appropriate to talk of pain and discomfort for people with dementia. We need to build on existing expertise and resources such as the National Council for Palliative Care’s *How would I know? What can I do?* guidance. We need to do this as part of wider work to
increase awareness and understanding among health and social care professionals of pain and discomfort for people with dementia, particularly in the last few days of life, but also throughout the progression of their dementia.

**Barriers: Inappropriate interventions**

The use of aggressive treatment for people with dementia is common and can include tube feeding and antibiotic treatment for infections\(^\text{100}\). Evidence suggests, however, that this treatment is inappropriate and does not improve survival\(^\text{101}\). A study into family satisfaction with nursing home care suggested that aggressive treatments such as feeding tubes decreased satisfaction with care\(^\text{102}\).

It would appear that such interventions are not what most families want or what their loved ones need. Families desired a focus on improving people’s comfort and an increase in advance care planning, both of which could be provided under a palliative care approach\(^\text{103}\). Palliative care is likely to be a more appropriate intervention for people at this stage.

**Barriers: Failure to adapt practice to reflect the different nature of dementia**

Many of the common practices and approaches which underpin good end of life care have been developed in response to conditions such as cancer and chronic obstructive pulmonary disease (COPD). In most instances, individuals with such conditions retain an acute awareness of what is happening around them until a relatively late stage of their condition. They retain cognitive ability in respect of decision making, awareness of pain and of the responses of those around them. As we have seen, for those with advanced dementia this is rarely the case.

What is required is a fundamentally different approach to care for those people with dementia. For example, health and social care professionals are trained to routinely tell family and carers that the last of the senses to go prior to death is hearing and that the person can hear what they say to them. Such advice is often accompanied by a caution to avoid saying anything which might cause distress to the person.

While a person with dementia will similarly retain the ability to hear family members, this does not mean that they have recovered the ability to comprehend what is being said to them, a faculty which may have been lost some months or years before. It is therefore important that health and social care professionals adapt their approaches and relevant practice to reflect the very different reality of dementia.

**Barriers: Lack of support for carers**

The characteristics of dementia include behavioural and personality changes, which can cause difficulties for those caring for people with dementia\(^\text{104}\).

Those with moderate or advanced dementia can be physically aggressive, experience hallucinations, be agitated or anxious and have sleeping disorders\(^\text{105}\). These symptoms can cause emotional and physical stress for the terminally ill person and carers\(^\text{106}\). People with dementia and their carers need specific support that may not be required by those with other terminal conditions. We need to improve the support available for those caring for people with dementia.
Families of those with dementia may also require a different type of bereavement support to those dying of some other conditions. While most people grieve after a person has actually died from a terminal illness, the families of those with dementia may experience different stages of loss as the condition progresses, for example when the person with dementia no longer recognises them\textsuperscript{107}. Bereavement support needs to reflect this important difference.
Conclusion

An ageing population and the resulting projected growth in dementia will pose a particular set of challenges to the health and social care system in Wales over the coming decades. In order to meet these challenges we need to start removing the barriers which currently prevent people with dementia from accessing high quality end of life care.

The bottom line is that we will not be able to meet the needs of the growing number of people with dementia by continuing to provide care and support in the way that we do today. We need to recognise that dementia is a terminal condition and that people with dementia have very different needs and experiences to people with other terminal conditions.

Many of our responses to terminal illness and ultimately to death and dying have been shaped in response to cancer. The modern hospice movement and palliative care approach have been shaped in response to the needs and experiences of people with different forms of cancer.

Our challenge over the next decade is to build on the progress which has been made in many areas in transforming core services so that they are able to respond to the needs of people with conditions such as dementia. This will require action across many different aspects of health and social care and the involvement of a number of organisations.

To this end, Marie Curie and the Alzheimer’s Society are committed to working together to raise awareness and understanding of key aspects of dementia and end of life care and to address the barriers identified throughout this report. Our task over the course of 2015 will be to identify the practical steps which will enable us to remove each of the barriers identified by this report.

Over the coming decades, a growing number of people will experience the different forms of dementia. The challenge for society will be to ensure that these people are supported to live their lives as they want for as long as they can. And when they approach the last weeks and months of their lives, they are able to access the most appropriate care, advice and support.

To make this a reality we need to start redressing the balance now, and we must all play our part.
Looking forward

We have identified target areas for improvement and change in order to enable people with dementia in Wales to live and to die as well as possible.

Strategy

Currently, there is little strategy in Wales about how the end of life care needs of people with dementia will be met in the future. We believe this must change. We look forward to working with the Welsh Government, Health Boards and other partners to begin removing the barriers which prevent people with dementia having the best quality of life possible until the very end and to ensure end of life care planning for people with dementia is properly person-centred.

Advance care planning

People who have been diagnosed with dementia and their families should be better supported to plan ahead for their care through tools such as the Lasting Power of Attorney. Health and social care professionals must be trained to sensitively and effectively engage people with dementia in advance care planning when it is right for them, recognising that wants and needs may change over time.

Hospitals

People with dementia are more likely to have inappropriate hospital admissions than people who do not have the disease. Better advance planning, empowering communities to become more dementia friendly and supporting care homes to deliver the best care they can all play a part in reducing this.

Still, almost two out of every five deaths from dementia occur in hospital and people with dementia often spend a considerable amount of their final year of life as a hospital inpatient. We recommend that dementia awareness training should be mandatory for all hospital staff so that they can appropriately identify and respond to the needs of people with dementia as soon as they enter the unit in order to minimise the potential harm of unnecessary admissions. Introducing a biennial National Care of the Dying Audit for Hospitals in Wales would allow for effective monitoring of improvements.

Care at home

Most people with dementia want to die at home, which for many will mean the care home where they normally live. Figures for Wales suggest that over half of deaths with dementia as the underlying cause occur in care homes. The palliative care needs of people dying with dementia need to be better understood and effectively provided for in the community, including appropriate levels of support in nursing and residential care homes.
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At Marie Curie, our mission is to help people living with a terminal illness and their families make the most of the time they have together by delivering expert care, emotional support, research and guidance.

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