Before The 2020 Pandemic: An Observational Study Exploring Public Knowledge, Attitudes, Plans, and Preferences Towards Death and End of Life Care in Wales

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Abstract

Background: Understanding public attitudes towards death and dying is important to inform public policies around End of Life Care (EoLC). We studied the public attitudes towards death and dying in Wales.

Methods: An online survey was conducted in 2018. Social media and the HealthWiseWales platform were used to recruit participants. Data were analysed using descriptive statistics and thematic analysis.

Results: 2,210 people participated. Loss of independence (84%), manner of death, and leaving their beloved behind were the biggest fears around death and dying. In terms of EoLC, participants sought timely access to care (84%) and being surrounded by loved ones (62%). Being at home was less of a priority (24%). Only 50% were familiar with Advance Care Planning (ACP). A lack of standard procedures as well as of support for the execution of plans and the ability to revisit those plans hindered uptake. The taboo around death conversations, the lack of opportunities and skills to initiate discussion, and personal fear and discomfort inhibited talking about death and dying. 72% felt that we do not talk enough about death and dying and advocated normalising talking by demystifying death with a positive approach.

Background

Contemporary Western societal approaches to death have been criticised for marginalising death and characterising it as a “forbidden subject”(1). This has hindered open conversations about death and End of Life Care (EoLC), despite UK research and policy(2–8) mandating the importance of effective conversations about the end of life(8–10). Cox and colleagues’(11) systematic review focusing on public attitudes towards death and dying in the UK has highlighted key concerns of fear of the unknown, experiencing distress, and becoming a burden to the family. ‘Understanding Dying’ was also identified as a theme in the James Lind Alliance national survey process of identifying the top ten research priorities for palliative and End of Life care(12). A supplementary analysis of participants’ narratives(13) emphasised a general lack of understanding of death and dying processes and the frustrations of participants when faced with a situation for which they felt unprepared. The need to reduce uncertainty of prognostication, a better understanding of what to expect, and how to recognise when death is near to
allow time for appropriate preparations were key concerns as well as the importance of supporting the wellbeing of staff, patients, and their families.

Death and EoLC is therefore clearly a public health issue\(^{14}\) with its high burden, social impact, and the need to prevent suffering. In the UK and worldwide\(^{15–17}\), there have been campaigns and initiatives to raise public awareness and facilitate discussions around death.

The COVID-19 pandemic has suddenly brought the reality of death to the forefront of public consciousness and highlighted new circumstances around death including the forced separation of patients and families at end of life, the challenges of future care decision making, and conversations on treatment escalation and place of care when dying\(^{18–21}\). The pandemic has highlighted death and dying as topics of immediate concern in our daily lives, forcing the way we think about death to re-assimilate at pace in what is likely to provoke a longstanding societal shift.

Public preference is vital in shaping effective palliative and EoLC services\(^{2, 22}\). Pre COVID-19 public data can be useful to determine the baseline in terms of how the public feels about death and dying and determine what, under the current circumstances, might be prioritised for public engagement. This paper describes the findings from a public survey carried out in 2018 with the aim of understanding people's attitudes towards death and dying in Wales. The specific objectives were to learn about people's:

- Fears about death and dying
- Preferences and priorities around EoLC
- Knowledge around the terminologies commonly used in EoLC and understanding about Advance Care Planning (ACP)
- EoLC plans
- Communication around death and dying

Most previous research has used either numerical or narrative data. In this study, we aimed to build on previous findings and use narrative from open-ended questions to integrate, interpret, and validate the findings emerging from the closed questions.

**Methods**

**Study design and setting**

This is a cross-sectional observational study using a mixed-method approach in data analysis.

Patient and Public Involvement (PPI) was fully incorporated into the development of the survey. The survey was conducted in Wales between 31/01/2018 to 21/05/2018. Self-selected participants responded using the Jisc Online Survey tool.

**Participants**
Adults residents in Wales and with capacity to consent were eligible for the study. An invitation to take part in the study and the link to the online questionnaire was advertised via social media and institutional websites. In addition, email invitations were sent to public and private organisations (Supplementary File 1: STROBE(23) checklist). We estimated that at least 123 respondents would be a good representation of the public in Wales considering the number of survey respondents in England(2).

**Questionnaire development**

A literature scoping process identified previous studies to inform the conceptual design. A questionnaire was then developed and tested for content and face validity by members of the research team. Relevant questions were adapted from existing survey questionnaires where possible including the Dying Matter Coalition(2) and the VOICE survey(24). In addition, the findings from the PeolcPSP study(12) helped inform some of the questions and refine the questionnaire. A panel of 10 topic experts and three colleagues with no expertise on this subject reviewed the draft of the questionnaire. The revised questionnaire was further validated by eight academics and seven research partners.

The final draft of the questionnaire covered six sections and included open and closed questions (with Likert scales) aiming to gauge participants’ i) views about death, ii) feelings about death and dying, iii) knowledge on commonly used terms in EoLC and relevant facilities, iv) preferences in EoLC, v) plans around death and dying, and vi) socio-demographic information. A consent statement followed a summary of the eligibility criteria, anonymisation issues, and data management (Supplementary File 2: Survey Questionnaire).

**Consent**

Participants were self-selected and proceeded to the survey if they agreed to the consent statement. After consenting, participants could avoid responding to any of the questions if they wished). A Welsh version of the questionnaire was available on request.

**Data analysis**

Participants responses would be the outcome measure. Descriptive statistics and regression analysis were used to analyse the numerical data. We studied the representativeness of the study sample by comparing the characteristics of the study sample to the Welsh population statistics. IBM SPSS statistical software package (V. 25) was used to run the analysis.

The narratives from the open-ended questions were analysed using Thematic Analysis(25). Following the six steps, the researchers first familiarised themselves with the data. The narratives were then systematically coded following a pattern of similar ideas whilst ensuring data relevant to these codes were collated across all survey data. Themes were derived in a deductive manner using the questions as a coding framework. Subsequent themes and subthemes were derived along the analyses using an inductive approach. A second independent coder validated the coding. Themes and subthemes (including any discrepancy in the coding) were further refined and reviewed following discussions with the study team. Verbatim quotes were used throughout the report as illustrative examples. NVivo (V. 12) was used
to manage and catalogue the data. Given the mixed methods approach, mixed-methods(26) and survey(27) study reporting guidelines are used throughout the paper.

Results

From 31/01/2018 to 21/05/2018, a total of 2,210 people responded to the survey. 12 requested the Welsh version. The sample was representative of all geographical areas across Wales. Males, people with chronic illness and disability, and people with a lower level of education were underrepresented (Supplementary File 3: Table S1). The participants’ circumstances at the time of the study are listed in Table 1.

| Circumstances of study participants (n = 2204)                                      | n   | %   |
|-----------------------------------------------------------------------------------|-----|-----|
| I am a bereaved carer/ family member/ partner/ spouse/ friend who lost a loved one in the last 5 years | 342 | 15.5 |
| I am a carer/ family member/ partner/ spouse/ friend of someone who is thought to be in the last few years of their life | 190 | 8.6  |
| I am a health / social care professional (non-clinical)                             | 122 | 5.5  |
| I am a health care professional (clinical)                                          | 204 | 9.3  |
| I am a health researcher/ an academic who has an interest in the subject            | 59  | 2.7  |
| I am a member of the public who has an interest in the subject                      | 949 | 43.1 |
| I am a professional working with people who are thought to be in the last few years of life | 99  | 4.5  |
| I am a volunteer working with people who are thought to be in the last few years of life | 16  | 0.7  |
| I consider myself to be in the last few years of my life                           | 107 | 4.9  |
| Other                                                                             | 116 | 5.3  |

Overall, missing data was negligible (below 2%). Around 10% of the participants did not answer two questions: having a chronic mental illness and if moved from another country.

The participants’ narratives merged under four main themes: enablers and barriers to talking about death and dying, fear of death, and EoLC preferences in terms of services and personal priorities (Fig. 1). A sample of quotations from each theme is reported in Supplementary File 3: Table S2.

To ease the presentation of the results, the answers to the closed questions are integrated, contextualised, and validated with the themes emerging from the participants’ narratives.

Fears about death and dying
The fear of being helpless and dependent was overwhelming across participants’ responses (Supplementary File 3: Fig S1). In their narratives, participants expressed their reasons for fearing death, these included: the manner of death, experiencing suffering and pain, losing dignity, facing the unknown, leaving wishes and deeds unfulfilled, and leaving loved ones.

“I don’t fear death but the manner of dying” (PID 32)

“If it happened now - where would my children live?” (PID 404)

**Preferences and priorities around death and EoLC**

The majority of the participants would wish to gain quality of life over survival (Supplementary File 3: Fig S2). In terms of EoLC services and personal priorities, participants showed a strong focus on being able to access care provision for themselves and their families and being surrounded by their loved ones (Fig. 2).

Similar to the numerical data, the participants’ narratives strongly converged towards aspects of service provision (i.e. timely access, peaceful environment) and presence (and wellbeing) of family and friends rather than a physical place of care.

“Having someone to discuss how end of life meds and treatment is administered” (PID 576)

“I'd like to be kept alive until my family can get to me to say goodbye! One lot lives in Scotland, the other in Australia” (PID 1262)

“The calm space to say goodbye well, and support to ensure everything that needs to be asked has been” (PID 1001)

“Knowing that there is someone there to give support to my loved ones during my illness/following my death” (PID 758)

Some wanted to avoid the burden and decisional conflict for families, particularly where capacity is lost.

“My family’s wishes would take priority over mine - I don’t want them to be more upset than necessary. This is particularly so if I have advanced dementia and am not aware of what is happening” (PID 214)

**Knowledge around EoLC and understanding of Advance Care Planning (ACP)**

The majority of the respondents were very familiar with many terms related to EoLC, however, only 50% and 45% were familiar with ACP and Advance Directives, respectively (Fig. 3).

Participants expressed doubts about their wishes being respected and frustration about the absence of a formal procedure or standard practice to deal with advanced care plans in the healthcare settings.

“Then it is important that they are listened to and not ignored. My mother’s medical power of attorney was ignored as the care home staff didn’t agree with it” (PID 366)
“Yes – BUT [there] needs to be a method of communicating this to the professionals for complete security. Too many Living Wills are ignored or are unknown” (PID 287)

Participants were confused about the appropriate timing for preparing the plan and opportunity of reassessment without which the plan might lose its relevance and efficacy.

“I think it’s ok to state your preferences in advance but there must be flexibility in the system because circumstances change, and these changes could affect the decisions one has made.” (PID 400)

Overall participants acknowledged that ACP would help increase their autonomy of choice whilst avoiding any unwanted treatment.

“Having a sense of control over how I will be treated and peace of mind that my wishes will be considered and respected.” (PID 56)

In practice, only 16% of the sample had formalised any EoLC plans such as Living Will (Fig. 4), by contrast, 61% indicated their intention to do so. Notably, 80% of the participants had formalised their decision around organ donation, which might be related to the opt-out system introduced in Wales in 2015.

**Communication around death and dying**

The majority of participants felt ready to talk about death and dying and valued strongly a sense of control and choice over EoL treatment and care, personal arrangements (e.g. financial) (Supplementary File 3: Fig S3) and life-sustaining technologies (Supplementary File 3: Fig S4).

However, 72% thought that, as a community, we do not talk enough about death and dying. Three main factors (Fig. 5), acting at three different levels, prevent conversations about death and dying: i) social perception and practice, ii) lack of opportunities and support, and iii) personal emotions and values.

**Social perception and practice**

Participants believed that people would not be interested in talking about death as this was not socially appropriate. Talking about death was considered a social taboo and perceived as impolite or morbid.

“Upsetting subject for others. Uncomfortable to think about - our society teaches us that death is a taboo subject” (PID 872)

**Lack of opportunities and support**

Participants found it hard to find the right time to talk about death or did not feel they had family members and friends with whom they would talk.

“Too busy living right now, not a priority” (PID 599)
Others expected support from health professionals but felt that time pressure or lack of communication skills did not allow for sensitive discussions. It was also feared that discussions would be documented. Lack of information around available services hampered these discussions as well.

“From previous experience Healthcare professionals often find it difficult to discuss” (PID 309)

**Personal emotions and values**

Many participants acknowledged their fear and discomfort around talking about death and dying, especially when still grieving for the death of someone close. The fear of hurting other people's feelings, especially those of family members, prevented initiating the discussion.

“I'm scared of dying and talking about death makes me anxious” (PID 779)

Participants identified opportunities to improve talking about death with normalisation emerging as an overarching theme, subthemes are summarised in Fig. 6. Participants believed that a cultural shift around the negative perception of death with a focus on demystifying death and a more positive approach to encouraging societal conversations are needed, and this shift would enhance the acceptance of death as a part of life. In an attempt to break the stigma around death, participants advocated a public health approach by including death and dying as part of school curricula, using national print, TV and social media, and other public platforms to promote conversations and more open signposting to existing EoLC and ACP services. GPs and other health and social care professionals could prompt discussions to avoid very late conversations with unprepared patients and families. This would be made easier by raising awareness within communities around issues such as palliative and End of Life Care, funeral arrangements, bereavement services, organ donations, and living will and help them to acquire the necessary competency to empower their decision making around planning for care at the end of life. Notably, some suggested adopting more coercive approaches such as making funeral plans compulsory and ACP universal in care-homes settings. A limited knowledge around where to retrieve the relevant information was also evident (Supplementary File 3: Fig S5).

**Discussion**

We used an online survey to study people's knowledge, attitudes, plans, and preferences towards death and EoLC in Wales. A total of 2,210 participants took part in the study. The participants contributed 2,610 pieces of narratives across eight open-ended questions.

Participants especially feared specific aspects linked to the process of dying such as the loss of independence (84%), the manner of death (narratives), and having to leave their beloved behind (54%).

In terms of service provision and personal EoLC preferences, participants favoured aspects of service provisions such as timely access to care (84%) and being surrounded by family and friends (62%). Being at home was less of a priority (24%).
Only 50% of the sample were aware of ACP. ACP was perceived as a means to increase the autonomy of choice around EoLC and to avoid unwanted treatment (narratives). However, this depended on the presence of standard procedures, support for the execution of plans, and the ability to revisit plans (narratives). These factors might explain why a large proportion of respondents intended to arrange EoLC plans but, except for organ donation, only a minority had formalized any EoLC plans.

Communication around death and dying were hindered by the taboo around death conversations, the lack of opportunities and skills to initiate discussion, and personal fear and discomfort. Respondents advocated normalising talking about death and dying - a cultural shift encouraged by demystifying death with a positive approach. Health professionals can initiate and support this conversation, but communication skills and manageable workload pressure were crucial for facilitating this.

Participants sought a public health approach to breaking the stigma around death and dying and endorsed the use of four channels to achieve this: a) using social media and other public platforms, b) providing formal education, c) developing formal and legal actions, and d) improving signposting and access to information.

The study findings provide robust estimates to corroborate previous research in terms of people's fears around the process of dying(11–13) and reinforces the evidence(28) that the preferred place of death is associated with specific environmental factors. Again, similarly to the Dying Matter Coalition survey(2), nearly 90% of the respondents felt comfortable discussing issues around death, but the percentage of people who had started any discussions with health care professionals remains small (a change from 2–3%). However, there was an increase from 7.5–16% in the number of people formally expressing future health care wishes. More recently, a mixed-method study(29) explored attitude to discussion death and dying amongst adults with advanced or terminal condition. For this study population, death was far from a taboo; instead, they found talking about death and dying “liberating”. Participants sought honest and open conversations about their choices and, similarly to our study, they expressed a need for more support to achieve what they define as a good death; including the choice of a safeguarded assisted dying law alongside good palliative care. This could in part explain why only 12% of the respondents had completed an Advance Decision to Refuse Treatment. The uptake of ACP remains low outside the UK as well. Yadav and colleagues(30) conducted a systematic review of the prevalence of Advance Directives (AD) among US adults and found that only one in three adults had carried out some form of AD. A similar percentage applied to residents of long-term care facilities across European countries(31).

Barriers to a higher uptake of ACP include scepticism about wishes being respected(32), social grade(33), communication skills and competency(32, 34–36). Abel and colleagues (2020) sustain that a shift of focus from medical treatments to health and wellbeing would facilitate an early engagement and ease this difficult conversation across all care settings(37).

Participants seek a public health approach to normalise talking about death(3–5). An effective public health programme should include an educational component, a community component, and government/service role(38). Initiatives such as Dying Matters in Wales are actively working towards this.
In Wales, a Clinical Lead and ACP facilitators at the Health Boards and charities such as MacMillan, Marie Curie, and Byw Nawr work towards raising public awareness of ACP in the community and educating facilitators and health care professionals. In 2015, the UK statistics indicated that a child loses a parent on average every 22 minutes(39). Bereavement can lead to social isolation and further consequences such as alcoholism, depression, and antisocial behaviour. This loss of human capital is, at least in part, due to unacknowledged loss. In some countries, death education is part of the school curricula(40). In the UK, this is currently limited to programs of support for bereaved young people(39).

As it is the case for other online survey-based studies, this study has strengths and limitations. The large sample allowed for robust estimates and rich narratives. An extensive literature review, questionnaire piloting, and PPI support secured the questionnaire content and face validity. The qualitative data provided additional dimensions and validations of the numerical findings. The survey was online and, in 2018, around 10% of the adult population of Wales reported not using the internet. Male, singles, and people with a lower educational background were under-represented, future research should consider oversampling from these groups. However, given the high percentages of participants agreeing to the attitudinal answers, it is unlikely that this seriously biased the estimates.

**The impact of COVID-19**

COVID-19 has severely altered the health environment and transformed the delivery of care. Health professionals might be unable to rely on customary norms to create rapport and continuity of care: Consultations occur through screens and protective equipment, communication with families and friends is done remotely, and difficult conversations might have to take place in emergency contexts, shortly after meeting the patient, because disease progression is such that the patient needs to be diverted to the EoLC pathway. However, today’s technology can support this consultation mode(41). In Wales, the NHS Wales Video Consulting (VC) Service is a video consultation service(42) rolled out by Welsh Government to offer video consultations where possible, the system has currently covered 90% of GP practices in Wales.

The daily updates on mortality counts and rates are crucial in highlighting the public health message and have instilled a deep awareness around death and dying. However, six months into the COVID-19 pandemic, mass-media headlines continue to focus on these statistics only. There is now a need to re-direct the conversation and pay attention to the process of dying and how we make sense of mortality.

In the UK and across the world, the pandemic has inevitably propelled Palliative Care (PC) centre stage(21) and turned it into a driver of best practice(43) to ensure access to EoLC is in respect of patients’ priorities and preferences. The PC philosophy is becoming the vehicle to safeguarding patients’ autonomy and avoiding decisions being driven by the fear of the pandemic(44). However, this is only possible if PC becomes more visible, starting with national (and international) guidelines and policy documents clearly embedding the contribution of PC when treatments no longer aim to cure and shift to comfort.
COVID-19 has also created an unprecedented number of fast-tracked research, which highlights the need to test and share examples of best practice as soon as they become available. A follow-up survey is underway to remedy some of the sample bias (45, 46) and to gauge real-time data about how COVID-19 has affected people's attitudes towards and engagement with EoLC planning.

Conclusions

The findings resonate with and reinforce knowledge from before the pandemic: People are ready to talk about death and dying, and COVID-19 has acutely increased awareness of this topic. Participants advocate the adoption of a public health framework and acknowledge the need for a combination of top-down and bottom-up initiatives across levels and settings to increase awareness, knowledge, and service utilisation. This would encourage earlier discussions and preparation of advance care plans and support health professionals and people towards shared decisions which closely align with people's end of life wishes and preferences.

Abbreviations

ACP Advance Care Planning
AD Advance Directives
EoL End of Life
EoLC End of Life Care
PC Palliative Care
PID Participant's ID
PPI Patients and Public Involvement
SMREC School of Medicine Research Ethics Committee
VC Video consultation

Declarations

Ethics approval and consent to participate

Participants gave written informed consent to participate. The research protocol was approved by Cardiff University's School of Medicine Research Ethics Committee (SMREC Ref 18/49).

Consent for publication
Not applicable.

**Availability of data and materials**

The datasets used during the current study are available from the corresponding author on reasonable request.

**Competing interests**

The authors declare that they have no competing interests.

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**Authors’ contributions**

AB and AN conceptualized and, designed the study and secured funding, II sought Ethics approval and developed and conducted the survey. II and ML carried out the data analysis and drafted the manuscript. All authors approved the final manuscript.

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**Figures**

**Figure 1**

Themes and subthemes emerging from the analysis of the open-ended questions
Figure 2

Participants’ preferences for EoLC services and personal priorities

Terms participants were familiar with(%)
Figure 3
Participants’ familiarity with EoLC terminology

![Figure 3]

**Statements about EoLC plans (%)**

| Statement                                                                 | I have done this | I intend to do this | Don’t know | I do not intend to do this | I will not do this |
|---------------------------------------------------------------------------|------------------|---------------------|------------|---------------------------|-------------------|
| Formally expressing my future health care wishes and preferences (e.g., making a living will) (n=1,133) | 16               | 61                  | 18         | 4                         | 0                 |
| Talking to someone about my end of life care wishes (n=1,187)             | 37               | 62                  | 10         | 7                         | 0                 |
| Talking to my doctors/nurses about my end of life care wishes (n=1,173)   | 1                | 63                  | 27         | 7                         | 1                 |
| Talking to someone about whether I want my body to be buried or cremated or donated (n=1,196) | 66               |                     | 30         | 31                        | 0                 |
| Making financial arrangements for the funeral (n=1,192)                    | 27               | 53                  | 12         | 7                         | 1                 |
| Making a decision on organ donation (n=1,197)                              | 80               |                     | 13         | 5                         | 1                 |
| Asking any family member/friend whether they have made a living will (n=1,172) | 24               | 28                  | 32         | 15                        | 8                 |
| Asking any family member/friend what type of care support they would want at the end of their lives (n=1,181) | 28               | 42                  | 21         | 8                         | 1                 |
| Asking any family member/friend about their funeral wishes (n=1,186)      | 43               | 36                  | 15         | 6                         | 1                 |
| Asking any family member/friend where they would like to die (n=1,135)     | 22               | 43                  | 24         | 9                         | 2                 |
| Asking any family member/friend about their financial preparations for end of life care (n=1,141) | 23               | 52                  | 28         | 14                        | 3                 |

Figure 4
Participants’ statements about EoLC plans

Social perception and practice

Lack of opportunity and support

- Personal fear and discomfort
- Feeling of inappropriateness
- Respecting others’ feelings

Lack of opportunity

- Stigma and social practice

Lack of others’ interest
Figure 5

Factors that inhibit talking about death and dying

“Just talk openly to normalise dying as part of living not a feared thing” (PID 1262)

“People don’t know what dying looks like and how to recognise it. This is sometimes a problem because loved ones call an ambulance because they need to be sure that patient has died” (PID 095)

Removing taboo and changing attitudes

Talking more

Raising awareness on death and dying and service availability

i) using media and other public platforms,
ii) providing formal education,
iii) developing formal and legal actions,
iv) improving access to information

“It needs to be made less taboo, discussed in schools at secondary level.” (PID 142)

Figure 6

Normalising talk about death and dying – focus and mode

Supplementary Files

This is a list of supplementary files associated with this preprint. Click to download.

- SupplementaryFile1ReportingchecklistSTROBEFinal.docx
- SupplemetaryFile2PADDWSurveyQuestionnaireFinal.doc
- SupplementaryFile3TablesandFiguresFinal.docx