Choice and place of death

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INTRODUCTION: CHOICE IN A CONSUMER SOCIETY

Never before have people been faced with such levels of choice as now enjoyed by those living in affluent societies. Increases in technological possibilities and relatively high levels of disposable income have enabled large numbers of citizens to exercise choice in a wide variety of areas, from contraception to managing their infertility, from deciding where to educate children to which career to follow, which TV/DVD/computer to buy, where to holiday, etc.

However, when it comes to deciding about death, choice is a more contentious notion. As the comedian Woody Allen once said ‘I am not afraid of death, I just don’t want to be there when it happens.’ While individuals can influence their health and illness through lifestyle choices, most have relatively little choice with respect to the time, manner and place of their death. With recent moves to legalize euthanasia and physician-assisted suicide in some countries, choice regarding place and time of death has been extended. However, even where such choice is available, it tends to be utilized by relatively few people. Less than four percent of deaths in the Netherlands are reported to be by euthanasia.

Although most people say that they would prefer to die at home, there has been a persistent fall in the proportion of people doing so. This trend has continued, with only 22% of cancer deaths in the UK in 2003 occurring there. The reasons for this are undoubtedly complex, involving many aspects of health-care practice and provision as well as social changes. The trend in falling home death rate has continued despite the rise in community palliative care services in the UK over the last 20 years.

In line with the all-pervading choice agenda, health care professionals are increasingly expected to ascertain a patient’s choice about preferred place of death, record it and aim to deliver it. Little is known about how health-care professionals elicited patients’ preferences and the extent to which eliciting such views enables them to facilitate their realization. In this paper we present an overview of the UK policy context and the published literature on preferred place of death, and examine key issues which arise for health-care professionals in exploring preferred place of death with patients. The purpose is to inform debate about the role health-care professionals play in helping patients make this most important of choices, and in enabling their preferences regarding preferred place of death to be respected.

CHOICE OF PLACE OF DEATH

With the recent White Paper Building on the best: choice, responsiveness and equity in the NHS, the UK government’s agenda of choice across the spectrum of health-care, from birth to death, was laid out. With regard to choice and dying it proposed, ‘To offer all adult patients nearing the end of life, regardless of their diagnosis, the same access to high quality palliative care so that they can choose if they wish to die at home.’ In addition, through funding the End of Life Care Programme, the government specifically sought to implement this policy objective. Since April 2006, within the Quality and Outcomes Framework, additional payments are payable to GPs for keeping a register of palliative care patients and meeting regularly to plan care at which ‘preferred place of care’ for individual patients should be noted.

For choice to be meaningful, it must involve at least two—preferably a range of—available, high quality options; which in terms of place of death should arguably include home, hospice, hospital and nursing home. In addition, enabling patients to die in their place of choosing depends on health-care providers understanding the patient’s prognosis, likely mode of death, and the availability and accessibility of services. They will then be in a position to assist individual patients and their carers articulate their preferences so that plans can be made and reviewed over time, to enable choices to be turned into realities when the time comes.

PREFERRED PLACE OF DEATH

Between 50 and 90% of patients with cancer, when they express a wish concerning place of death, would choose
to die at home. Several studies from a variety of countries have linked patients’ expression of their preferred place with actual place of death, showing more patients achieve a home death if they have expressed a wish to do so. In addition, patients and carers agreeing on home as the preferred place of death has been shown to be a strong predictive factor in achieving a home death.

Seventeen factors strongly associated with place of death have recently been reported by Gomes and Higginson. Based on a review of 58 studies, they suggested a conceptual model for ordering such factors in terms of illness, personal and environmental issues, also highlighting the dynamic nature of the model since all factors interact with each other. Preferred place of death is recognized as changing throughout the disease trajectory, with some studies showing that a preference for home death decreases over time, whilst home remained the preferred place given more favourable conditions.

The pattern of change over time was studied qualitatively by Thomas et al. in an interview study of cancer patients and their informal carers. This highlighted the complexities surrounding the issue of preferred place of death. Contextual factors that influenced patients’ choice included extent of social network, perception of carers’ attitudes, symptom management, fear of loss of dignity and the views of the patient/carer of the care provided by available hospices, nursing homes, hospitals and community nursing and social services. Patients’ choices were characterized by uncertainty. Preferences were rarely categorical, but were qualified by speculation about the unpredictability of events.

Although the same standards of care should be offered to all patients with end-stage illness, as is common with issues relating to end of life care the majority of studies into preferred place of death have focused on patients with cancer. Available evidence suggests that the preferences of patients with other illnesses, and the factors influencing them, are not dissimilar from those with cancer. In a qualitative study of elderly patients’ views regarding home as a place of care when dying, concerns were expressed regarding the quality of care which could be provided at home, the appropriateness of their children providing intimate care, and ambivalence towards having professional carers in the home. One telephone survey of elderly patients who had recently been admitted to hospital with cardiac failure, chronic obstructive airways disease or pneumonia found that approximately equal numbers expressed preferences for home compared with hospital care. In a follow-up interview study with a selection of these patients, familiar themes regarding not wanting to be a burden on family carers emerged.

DETERMINING PREFERRED PLACE OF DEATH

It has been suggested that enabling people to die in their preferred place requires greater empowerment of patients and families, early and continuous risk assessment, and better training in palliative care for health-care professionals, including those that work in primary care. However, there is very little evidence to inform how this should best be done. Issues regarding preferred place of death will often be discussed along with other concerns of the patient, including questions regarding likely prognosis and mode of death. The difficulty in predicting prognosis and mode of death may confound such discussion. The very concept of a terminal phase with non-cancer patients is problematic with, for example, patients with end stage heart failure or chronic obstructive airways disease tending to follow a trajectory of steady decline with episodes of acute deterioration and recovery, rather than the relatively rapid decline as tends to occur for cancer patients with advanced metastatic disease (Box 1).

Whilst determining prognosis for cancer patients is problematic, with a tendency for overestimation of survival, clinical prediction criteria for prognosis in patients with end stage non-cancer diagnosis have been shown to be ineffective; in addition, one study showed that primary care physicians tended to over-estimate the likelihood of death at one year in patients with end stage heart failure.

Each individual patient’s expectations of the dying process and the way that this shapes their wishes is unique. Helping patients to identify and articulate their preferences, and how these evolve in response to the patient’s changing condition, requires well-developed communication skills, including considerable sensitivity to differences in values. Such discussions can only take place effectively within the context of an ongoing and trusting patient-clinician relationship, and may require a considerable investment of time.

The extent to which patients wish to discuss their preferred place of death varies from individual to individual. While some will have openly acknowledged that they are dying, may be clear of their wish and will articulate clearly their preferred place of death without being prompted, others will be in denial regarding their terminal illness, using this as a protective mechanism; the subject of where they wish to die may be off limits. Still other patients—possibly the majority—will be ambivalent. They may not voice their concerns or wishes openly but may be ready to discuss the issue if sensitively broached. They may be clear of what they would want in optimal circumstances, but may recognize that circumstances as they approach the time of dying may not favour this and that an alternative place of death might occur.
to avoid inappropriate management, particularly hospital admission, at the end of life (Box 2). Informing secondary care providers of a patient’s wish to die at home can enable forward planning so that discharge can be arranged at short notice if necessary. However, the imminence of death is often difficult to recognize, and problems with estimating prognosis may have the consequence that hospital admission (e.g. for symptom control), although intended to be short-term, will result in the patient not being discharged. This may be especially problematic in non-cancer diagnoses, where prognosis is even more difficult to predict. While further research may enable new models which improve accuracy of prediction for both cancer and non-cancer diseases to emerge,""},29,30 given the non-linear complexity of terminal illness, the precision of such models will always have its limitations.31

**EFFECTS ON THE HEALTH-CARE PROFESSIONAL**

Little is known about the effect on health-care professionals of discussing preferred place of death. The process may stimulate negative emotions, with concerns that the patient will react emotionally to the subject being broached. They may have personal fears regarding discussing such a difficult and emotive subject openly, including uncertainty about how best to handle the patient’s and carer’s feelings, or worries about being asked to make assurances which they may not be able to fulfil. They may have concerns about mortality for themselves or their family and friends, or they may have unresolved grief from past bereavements which makes the subject of death particularly difficult to discuss. Personal knowledge of the patient over time may help to prepare the professional and the patient for discussion of the issue, but conversely it may also make the issue more difficult to discuss openly if an emotional tie has developed between them. Professionals with a close relationship with patients may tend to overestimate prognosis and fail to recognize the impending onset of the terminal phase.32

The health-care professional may feel it particularly difficult to discuss preferred place of death with a patient when providing adequate community services or securing admission to a hospice may not be possible. Preferred place of death may be a hollow concept and the promise of choice a cruel sham if services are not available because of lack of funding or other resources. Whilst simplistic rhetoric promising patients a planned death is seductive, the reality may be that the complex uncontrollable nature of the dying process exposes real choice as being fool’s gold.

**CONCLUSION**

In this paper, we have summarized issues that influence choice and preferred place of death. Ascertaining a patient’s
Box 2  Advance care planning around preferred place of death

Case 1: Mr A, a 76-year-old ex-lorry driver with NYHA stage IV heart failure, diabetes, painful neuropathy of his lower legs and gout in his feet.

Mr A is housebound and cared for by his 69-year-old wife, who is in reasonable health. He has had several admissions to hospital with acute left ventricular failure (LVF) over the last two years, which he always finds distressing. He discusses future management with his cardiac failure nurse on one of her regular visits, when he says he has ‘had enough.’ They explore alternatives to hospital admission for any further episode of acute LVF, and Mr A agrees to a multidisciplinary discussion of plans for managing him at home if at all possible. This discussion takes place at the next primary care team meeting between the GP and district nurse which the cardiac failure and Macmillan nurse attend.

A plan is drawn up for Mr A to have subcutaneous diamorphine via a syringe driver and furosemide as required at home if and when further LVF occurs. The cardiac failure nurse checks the appropriate dose of furosemide with his cardiologist and the Macmillan nurse and GP decide on the appropriate dose of diamorphine. A prescription for the agreed drugs is written so that anticipatory medication can be kept in his home. Instructions are written in the district nursing notes kept in his home and faxed to the out-of-hours primary care provider.

Six weeks later Mr A suffers breathlessness and acute chest pain on a Sunday afternoon. He is visited by the on-call GP who gives him intravenous furosemide and arranges for the district nurse to commence the diamorphine driver.

All has gone to plan but the outcome is still not certain. The following two scenarios are amongst several possibilities:

Scenario A: Mr A’s pain is controlled and his breathlessness improves. Mr A becomes unconscious in the evening and dies the early hours of the next morning, surrounded by his family.

Scenario B: With controlled pain and improved breathlessness, Mr A remains bed-bound for several days, but with review by the cardiac failure nurse who receives telephone advice from the cardiologist. His ACE inhibitor dose is adjusted and he recovers to the level of his previous functioning, though he remains frustrated at his poor quality of life.

Case 2: Mr B, a 63 year old retired headmaster with lung cancer

Mr B has discussed his wish to die at home with his GP since early after his diagnosis with advanced disease six months ago. He has had two admissions with shortness of breath, for which he has had a right pleural tap and a transfusion on one occasion. He has been increasingly short of breath for a week and the district nurse takes blood for a full blood count. His haemoglobin is 7.6. His GP visits and discovers that he also has reduced air entry in his right lung base. They discuss the situation and Mr B agrees that he should be admitted onto the oncology ward for a transfusion and possible pleural tap. On admission an ultrasound scan of his chest reveals mostly solid tumour with a small amount of loculated fluid in his right chest. Transfusion is arranged for three units of blood.

Despite transfusion the breathlessness increases and Mr B becomes quite distressed. He is using continuous oxygen, and a syringe driver with diamorphine and midazolam sedation is commenced. He settles to some extent but is quite drowsy and confused. His wife and family feel that they could not cope even with additional support at home and request that he remains in hospital. Mr B is too confused and drowsy to enter into discussions. He dies in hospital after a further 48 hours.

The Macmillan nurse spends time with Mrs B exploring her feelings with regard to her husband’s wishes for a home death not being met. She is upset and feels that she has failed him. Several meetings are needed to support Mrs B.

Finally she accepts the inevitability of the hospital death, but still feels sad that her husband did not die at home as he had hoped.

wishes with regard to place of death is an important function in end of life care, and should be approached sensitively and regularly reviewed as the patient approaches death, within a realistic evaluation of the feasibility of different options. Discussions will need to involve the patient’s family and/or other informal carers. Preferences when clarified can aid in advance care planning to enable patients to achieve their goals; however, the complexities surrounding clinical course and limited resources are likely to confound patients’ plans even with health-care services of the highest standard.

Since the issue is difficult to discuss, a professional with an established relationship with the patient—such as the GP or district nurse—is arguably the most appropriate person to perform the task, using a sensitive approach, well developed communication skills and providing ongoing relational continuity. Whilst clearly recording the patient’s preference in the health-care record is important, it must not become a mechanical ‘tick box’ exercise. The recent change to the Quality and Outcomes Framework in the UK has introduced a financial incentive for practitioners to record patients’ preferred place of death. While this may encourage more patient-centred end of life-care, perversely it could lead to a more perfunctory approach. Investigating the experiences of health-care professionals in this difficult and sensitive task is an important area for further research.
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