Results Two patients with PDOC were admitted to the hospice over a 12 month period for withdrawal of CANH. A 28-year-old man who had been in a vegetative state for 15 months secondary to a cerebral abscess and a 73-year-old woman who had been in a vegetative state for 6 years following a hypoxic brain injury. They both died at the hospice 9 days after discontinuing CANH. Key themes identified were educational needs of hospice staff especially an understanding of how reflex reactions may be misunderstood as purposeful actions in PDOC, the value of providing an opportunity for staff to debrief following episodes of care and the importance of the hospice team being involved in the development of a detailed end of life plan prior to hospice admission.

Conclusion There are numerous medical, ethical and legal challenges encountered in making the decision to withdraw CANH and then in providing end of life care for these patients. Learning from our work may help other professionals caring for similar patients as we predict an increase in referrals to hospices to be involved in managing the care of similar patients.

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INEQUALITIES IN HOSPICE CARE PROVISION: A SYSTEMATIC LITERATURE REVIEW AND NARRATIVE SYNTHESIS

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Background Hospice care is often perceived as being associated with stereotypical patients: white, middle-class, young and with cancer.

Aim To review the literature concerning who is/is not referred for hospice care, with a focus on diagnosis, age, gender, marital status, ethnicity, geography and socio-economic status.

Design of study Systematic literature review and narrative synthesis.

Methods Searches of Medline, PsycINFO, CINAHL, Web of Science, Assia and Embase databases from January 1987 to December 2017 were conducted, with further citation, reference and hand searches. Studies of adult patients living in the UK/Australia/New Zealand and Canada and receiving a range of hospice care (inpatient, outpatient, day hospice, hospice at home, community specialist (Macmillan) nursing) were included. Case studies, opinion pieces and systematic reviews were excluded. Of 28 433 articles reviewed, 152 met the inclusion criteria. Extracted data were analysed using NVivo with a narrative synthesis of emergent themes.

Results Since 1987, there has been extensive research demonstrating inequalities in hospice care provision. There is some evidence that some inequalities are reducing, particularly for those with non-malignant diseases. Nonetheless, there are still important groups with less access to hospice care, including the oldest old, ethnic minorities and those living in areas of high socioeconomic deprivation. Gender and marital status have little effect on hospice care provision. The literature concerning hospice care provision for the LGBTQ community and those living with HIV/AIDS is very limited.

Conclusion Prognostic uncertainty and fragmentation of care remain barriers to receipt of hospice care for those with non-malignant disease. The oldest-old, the socio-economically deprived and those from ethnic minorities continue to receive less hospice care. There is a persistent challenge to the hospice movement to provide equitable access to all.

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THE DEVELOPMENT OF A FRAMEWORK TO PERSONALISE HYDRATION MANAGEMENT IN CANCER CARE: THE USE OF NON-INVASIVE TECHNOLOGY TO EVALUATE FLUID STATUS AND DEHYDRATION-RELATED SYMPTOMS

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Background The role of hydration in causing or alleviating suffering in advanced cancer is poorly understood. The evidence for the efficacy of clinically assisted hydration in advanced cancer is inconclusive. Bioelectrical impedance vector analysis (BIVA) is an accurate validated method of assessing hydration status. Previous work with BIVA demonstrates...