MOVING UP THE ‘TRIANGLE’ TO CREATE A RESEARCH ACTIVE HOSPICE

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Background There is a welcome and appropriate movement, encouraged by Hospice UK, towards increasing the role of research in palliative and end of life care.

Aim To enable Hospice Isle of Man to reposition as a Research Active Hospice.

Methods A survey was undertaken to ascertain the current level of research interest and awareness in Hospice Isle of Man.

Results to date The survey was completed by 25 staff and 45 volunteers. The majority (88%) of clinical staff had some research training but most had no recent opportunity to engage in research; over 70% agreed that research added value to their daily work but only 40% were confident in their critical appraisal skills. As a result further training e.g. in critical appraisal skills within journal clubs is being initiated. Amongst the volunteers 22% were identified to have research skills but very few were actively engaged in research. These volunteers will be invited to participate in future projects. Funding from the Scholl Foundation and the Manx Lottery Trust has enabled research staff to be recruited and has facilitated protected time for some clinical staff to participate in research with peer support and skill sharing in, for example, developing outcome measurements, appropriate analyses and interpretation of data and considering how the value of routine data can be improved. Initial uncertainty about the role of research and researchers is gradually being replaced with appreciation of their support in enhancing research awareness and activity among staff.

Conclusion Hospice Isle of Man had a low level of research activity however a step-change has been achieved by utilising existing skills and interests of the clinical staff. Key drivers were obtaining funding for protected research time for key clinical staff and employing experienced researchers to support and complement the clinical staff to develop an active research culture.

ARE WE ACHIEVING PREFERRED PLACE OF DEATH (PPD) IN PATIENTS REFERRED TO OUR PALLIATIVE CARE SERVICE?

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Aim To review whether a cohort of patients referred to the Countess Mountbatten Palliative Care service achieved their preferred place of death (PPD).

Method A clinical notes review of the first 100 patients referred to the service from 1st June 2017 specifically looking at preferred place of death, actual place of death and, if relevant, reasons PPD wasn’t achieved. Length of time known to service and likelihood of achieving PPD were also studied.

Results Of the 100 patients studied 35 had PPD of Home, 23 had PPD of Hospice, three had PPD of acute hospital, 27 PPD remained unknown and 10 had no preference.

PPD of Home was achieved in 17 of the 35 patients (49%). PPD of Hospice was achieved in 22 of the 25 patients (88%) and PPD of acute Hospital was achieved in all three patients (100%). Patients known to the service between 2 to 12 weeks were the most likely to achieve their PPD (64%).

Discussion Lack of care and equipment at home, carer distress and difficult symptom control were the main reasons for not achieving PPD of Home. Reasons for not achieving PPD of Hospice included rapid deterioration making patient transfer inappropriate. Where PPD was unknown, most patients were only known to the service for a short period before death and the majority of patients were acutely unwell and PPD discussion was felt to be inappropriate.

Conclusions Rapid access to care and equipment in the home could result in more patients achieving their PPD. Increased community support and a hospice at home service could potentially provide this.

HOSPICE CONTRIBUTION TO THE UK NIHR NATIONAL CLINICAL RESEARCH PORTFOLIO: 2018 UPDATE

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The National Institute for Health Research (NIHR) is responsible for funding, delivery and monitoring of clinical research studies in the NHS. The main subset of NHS studies relevant to hospices is categorised under ‘Supportive and Palliative Care’ and ‘Psychosocial Oncology and Survivorship’ (SPC/POS). In the NIHR, the Clinical Research Network (CRN): Cancer Cluster is responsible for overseeing the delivery of this subset within the NIHR portfolio, not only in cancer but also extending to advanced non-malignant disease.

Since January 2018, non-NHS sites such as hospices are also officially recognised as sites for NHS-based studies. Of course, hospices have been involved and indeed leading in clinical research for many years – the Cancer Cluster office is now regularly tracking this important contribution. In 2016–2017, there were a total of 49 studies open in the SPC/POS group, of which 43 were actively recruiting. There is a balanced mixture of observational and interventional research. Participating sites included 22 hospices and 125 hospital trusts. (Some hospices may be listed under their local trust name.) There is wide variation across England and the devolved nations with respect to participation in these studies.

Funding for studies relevant to hospices came mainly from NIHR and other major charities, including Marie Curie, Macmillan, research councils. A few studies have commercial sponsorship.

The NIHR CRN is committed to helping the hospice sector to increase its access to clinical trials, through the NIHR Charities Consortium. It will do this by disseminating information and supporting the research workforce, working with Hospice UK. The Consortium held its first national conference on hospice and community-based research in April 2018. This will become an annual event and there will be prizes to reward hospices and other SPC/POS research teams for making significant contributions to the national NIHR portfolio.
Patient care

P-75 ‘COLD BEDROOMS, COOLING BLANKETS, AND CUDDLE COTS’ PROVIDED BY CHILDREN’S HOSPICES: MAPPING PRACTICE

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Background The ways parents are supported at the end of their child’s life and after death can profoundly affect the grieving process. Within children’s hospices cold bedrooms, cooling blankets and cuddle cots are provided to enable families to be close and have unrestricted access to their child after death. Such facilities are thought to support parents in the initial stages of grief. However, there is wide variation in practice and almost no evidence to inform parents’ decision-making during advance care planning.

Aim We aim to map current practice and explore parents’ experiences of using cold bedrooms and/or cooling blankets and/or cuddle cots – either in the hospice or at home – and the perceived impact of this on immediate and long-term bereavement.

Methods This study comprises two stages. First, a cross-sectional survey to gather data on previous and current practice, and reasons for inclusion of these facilities within the portfolio of care and support, collected via structured telephone interviews with heads of care in all UK children’s hospices (n=52). Second, qualitative interviews with parents (n=30) exploring stories of the early days of bereavement and the support received from the hospice, their needs and desires in those early days and the perceived role of these facilities on early and subsequent experiences of bereavement. Survey data will be analysed using descriptive statistics and content analysis. Interviews will be analysed using narrative and thematic approaches.

Results We will not have data to report but hope presenting this work will alert others to this research and generate connections and meaningful conversations with others with an interest in this area.

Conclusions Findings will have implications for practice within children’s hospices and also for statutory services which are beginning to offer this provision.

P-76 ASKING SENSITIVE QUESTIONS ABOUT SEXUALITY ON AN UNCERTAIN LIFE COURSE

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Background The term ‘sensitive research’ first emerged in the 1960s. Since then there has been a growing acceptance that many topics are ‘sensitive’ in nature, including those that separately address sexuality and disability at the end of life but maybe further compounded when such topics are explored together.

Aims This presentation will focus on four intersecting ‘sensitive’ issues: sexuality, disability and young adults with life-limiting or life-threatening conditions on an uncertain life-course, who may or may not be approaching the end of life. The sensitivities of conducting research about sexuality with young disabled adults and the dilemmas encountered by both researchers and practitioners will be addressed.

Methods and results This presentation draws on the in-depth, qualitative research interviews of the first author’s doctoral thesis about intimacy, relationships, sexuality and the reproductive choices of 13 young adults with life-limiting conditions, two partners and the views of 10 parents and 10 practitioners (Blackburn, Earle & Komaromy, 2014). The findings highlight multiple taboos and ‘sensitivities’ when researching sex with young adults who may or may not be approaching death. These include addressing posthumous consent in advance of death, access to research participants and the implications for General Data Protection Regulations. Affective aspects for addressing the research findings and reflections on why research with young adults about sexuality, disability, dying and end of life is often regarded as taboo and sensitive will be explored.

Conclusions There is a delicate balance as to whether research about sex encroaches too sensitively on participants’ lives, whilst recognising that there are important ethical considerations for the research relationship between participants and researchers.

P-77 SEXUALITY AND RELATIONSHIP CHOICES IN PEOPLE WITH LIFE-LIMITING OR LIFE-THREATENING CONDITIONS

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Until recently, surviving into adulthood with a life-limiting or life-threatening condition (LLTC) was unusual (Beresford & Sturtard, 2014). Young adults with LLTCs are a growing population. It is therefore timely to explore their personal choices and considerations about sexuality, intimacy, and reproductive choices. This research aimed to understand and make the sexual knowledge and experiences of people with LLTCs more accessible and meaningful.

This presentation focuses on the first author’s PhD thesis; centring upon 13 young adults, age 16+ and featuring some, but not all, life-limiting or life-threatening conditions, but including LLTCs, such as, Cystic Fibrosis and Duchenne Muscular Dystrophy. This research also sought the views and contributions of family supporters: two partners, ten parents and ten care practitioners, in order to gain their understanding about the sexual knowledge and experiences of people with LLTCs. This research used a qualitative approach, underpinned by a life course theoretical perspective. What has emerged is a picture of the transitions which occur from childhood to adulthood, from comparative wellness to progressive illness, uncertainty about the life course and most importantly, the sexuality of people with LLTCs.

The findings aim to influence policy and practice. They suggest that the voices of young adults are insufficiently heard, and that there is a gap in the provision of information particularly in relation to their sexuality and reproductive choices. As an increasing number of young adults strive for independence and sexual citizenship, efforts need to be congruent with and grounded in the views of those most personally involved; the young adults themselves.