**A RESPONSE TO GOSPORT: REGIONAL PROJECT ON ANTICIPATORY PRESCRIBING AND BENCHMARKING PRACTICE CROSS-HOSPICE IN YORKSHIRE**

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**Background and introduction** The findings of the enquiry into events at the Gosport Hospital, published in June 2018, demonstrated shocking misuse of opioid and benzodiazepine medications and syringe drivers leading to the shortening of many lives. Hospices frequently utilise opioid and benzodiazepine medications, including via syringe driver, and need to be able to demonstrate that their practice is safe and appropriate.

**Methods** Individual hospice units developed their own responses to the Gosport report. Both Prince of Wales and St Catherine’s Hospice undertook retrospective audits of hospice inpatient unit care. St Catherine’s Hospice presented their work for discussion at the regional ‘Learning group’ meeting.

**Results** The two sites utilised different collection proforma and reviewed care given to 93 hospice inpatient unit patients, demonstrating safe and appropriate practice at both hospices. Anticipatory prescribing was personalised to the patient, considering factors such as renal function, and was judged ‘appropriate’ in 82–100% of cases. 80 patients died and, as anticipated, there was significant use of opioids, benzodiazepines and syringe drivers. The use of opioids and syringe drivers was found to be proportionate and appropriate. Areas for improvement included recording of indication for both prescription and administration of medications. Through discussion of these results, it was acknowledged that a regional approach to auditing would be advantageous in standard setting and comparison. Both pilot proformas had strengths and weaknesses and are being utilised to develop a regional audit proforma with plans to establish regular audit and re-audit cycles.

**Conclusions** As specialists with a high use of opioid and benzodiazepine medications, there is a need to demonstrate that our use of these medications is safe, consistent and of exemplary standard. The hope is that this approach will allow identification of any areas of difference in practice and offer quality assurance across region and Hospice.

**PLACING DEATH AND DYING: ON THE WORK OF MAKING PLACE AT THE END OF LIFE**

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**Background** Palliative care policy and professionals are concerned about the location of care, epitomised in phrases like ‘preferred place of care’ and ‘preferred place of death’. There has been an emphasis on home being the ideal place. Conversely, it is recognised that high quality palliative and end-of-life care can be delivered in a wide variety of settings.

**Methods** This presentation is part of the ongoing Forms of Care project about palliative care. Ethnographic methods include observation of over 60 in- and outpatient palliative care team meetings, 1 year of shadowing palliative care team members in their daily work, and 15 interviews with palliative care staff, including doctors, nurses, social workers, and physiotherapists. Three patients and their families were also interviewed multiple times. Data was thematically analysed, drawing on social theory.

**Results** Palliative care professionals consider place as part of the care they deliver. Hospital was typified as a place of control, known infrastructure, skilled staff and support, and constant monitoring. The home was a place of being a ‘guest’ and unknown quantity, creating a shift in power dynamics between professionals, and patients and their families. The drive for patients to be cared for at home came with added costs in terms of resources and the emotional labour. However, many cases extended beyond this binary. We highlight the work both professionals and patients do to keep a place stable. We suggest such efforts of ‘placing’ show how place is not merely a geographical location, but made through relations, interactions and activities.

**Conclusion** In the push to enable patients to die at home, what tends to be overlooked is the work that goes into ‘placing’: the work of ‘making place’. The research indicates the importance of understanding how placing gets done as a form of care.

**ADVANCED CARE PLANNING IN PATIENTS ADMITTED FROM CARE HOMES TO THE EMERGENCY DEPARTMENT**

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**Background** Early identification of patients in their last year of life is a national focus, enabling the expression of preferences through Advance Care Planning (ACP). This is particularly relevant to patients in nursing homes, who die on average within 18 months of admission. This service evaluation aimed to establish if ACPs are followed appropriately by care homes and hospital staff.

**Methods** Retrospective data collection was undertaken for patients aged >75 admitted to the Emergency Department over a 6-week period (January-February 2019). A total of 120 patients from care homes were included. Review of the Coordinate My Care (CMC) database identified which patients had existing ACPs.

**Results** 17% of patients had a CMC record on admission. 26% had one created subsequently, and 57% had no CMC record. The most common recorded diagnoses were Dementia (45%) and Frailty (25%). 30% of patients were ‘For treatment of reversible conditions in the home/hospice setting’. Hospital admissions for these patients should theoretically be avoided, but in 83% of cases there was no acknowledgement of their CMC record. For patients who had a CMC record created during or post admission, 61% were identified as not for readmission. This group had the highest proportion of subsequent deaths (26%) and lowest number of subsequent admissions (0.4 per patient).

**Conclusion** Less than 20% of patients from care homes had a pre-existing CMC record, despite being a group to target for ACP. 30% of admissions were inappropriate and 16% of patients with newly created CMC records had subsequent inappropriate admissions (including two inpatient deaths). Further work is required to identify the challenges to managing patients within the community and factors contributing to hospital admission. Finally, when patients had a pre-existing CMC record there was poor recognition of this. Improved
awareness is necessary to ensure that ACPs are reviewed appropriately.

**57** WHO WOULD KNOW? EMBEDDING FUTURE CARE PLANNING FOR ALL PATIENTS APPROACHING THE END OF THEIR LIVES

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**Background** Future Care Planning (FCP) empowers individuals to receive the care and treatment they choose towards the End of Life (EoL). Developing personal Advance Care Plans (ACPs) and Anticipatory Clinical Management Plans (ACMPs) improves patient and family satisfaction and reduces ‘burden-some’ treatments yet is not universally offered. An evaluation of current FCP across a locality was undertaken to establish current practice, explore barriers to implementation and guide future work.

**Methods**
1. Case note review of patients known to specialist palliative care services (SPCS) for evidence of FCP and information-sharing between organisations (n=52)
2. Review of ACMP content and information-sharing between organisations (n=50)
3. Health-care professional survey of knowledge, experience and expectations of FCP (n=39)

**Results**

- Preferred Place of Care was documented for 85% of patients known to SPCS, but there was limited documented evidence of other forms of FCP (uDNACPR: 38%, ACMP: 2%, ADRT/LPA: 0%). FCP undertaken by SPCS was rarely available in hospital records. The number of ACMPs is increasing (49 (2017/18) c.f. 198 (2018/19)), predominantly for care home residents, with 60% developed by newly appointed frailty practitioners. They consistently included key EoL topics but only 38% were available within hospital records. Staff had limited knowledge and experience of FCP, and the processes for recording and sharing patients wishes. They felt more training and more time would enable them to facilitate more ACP conversations.

**Conclusions**

- Not all patients are currently offered an ACP conversation. Although information packs are available further promotion is required to ensure they are given. There is a need to continue current network-wide ACP training; additional targeted support for those in key roles has been introduced. Collaborative working between SPCS, frailty, acute and primary care teams is ongoing to ensure a consistent approach, and shared documentation and processes to ensure patients wishes are respected as they move between care settings.

**58** ANTICIPATORY PRESCRIBING AND USE OF SYRINGE DRIVERS – CROSS BOUNDARY AUDIT OF PRACTICE

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10.1136/spcare-2020-PCC.79

**Background** Anticipatory prescribing and use of syringe drivers at end of life is now established practice. A national report outlined the need to ensure safe, appropriate and individualised prescribing for patients at end of life, considering available guidance to aid decision making. Within our locality we deemed it vital to demonstrate we are complying with guidance.

**Aims**

- To ensure safe and effective care of patients at end of life, with an individualised approach to decisions and compliance with guidelines, policies and procedures.

**Methods**

- Questions were developed through collaborative discussion and data collection tool produced. Retrospective audit cross settings was undertaken based on place of death to avoid duplication.

**Results**

- 71 cases included for full analysis. Largest proportion in Hospice. Spread of gender, age, diagnosis. Substantial data was collected on core drugs at end of life. Using agitation as an example, 93% had an individualised approach to prescribing: all had a dose and route; 1 no frequency although this was felt to be a documentation error; 31% had a maximum dose; 93% had an appropriate dose adjustment for renal impairment. In the last 24 hours of life the total dose of midazolam: Hospice median 1.25 mg and mean 8.2 mg; hospital 0 mg and 1.7 mg and community 0 mg and 5 mg respectively. Similar results were found for other drugs in terms of quality of prescriptions and individualised nature of decision making.

**Conclusion**

- Individualised approach to prescribing was evident overall, with a high standard of prescribing. Some incomplete prescription although felt due to data collection errors. Doses of medications in line with network guidelines. Need to improve discussion with patients and families regarding medications, syringe drivers and side effects. The three organisations involved developed individual action plans including improvements in documentation and where electronic prescribing had commenced, embedding this.

**59** IMPROVING IDENTIFICATION AND RECORDING OF THE GOLD STANDARDS FRAMEWORK (GSF) WITHIN RUSSELLS HALL HOSPITAL, DUDLEY

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**Introduction** Measuring performance in End of Life Care within an acute setting is a complex and difficult task. GSF implementation within a hospital setting can improve quality of care, patient satisfaction and an increase in advance care planning. This study aims to show how by developing an internal hospital application and protocol for GSF patients, improved data collection can be transformed from data into intelligence and impact on delivery of high quality, individualised end of life care.

**Methods**

- Most wards within Dudley had different ways of capturing GSF identified patients and recorded information manually. Informatics and the Specialist Palliative Care team developed an internal application to monitor all patient admissions against a GSF outcome and to provide additional information such as preferred place of care and ACP. This provided the ward with a visual management tool, easily identifying end of life patients and the ability to capture essential end of life metrics.

**Results**

- Since the launch of this application, analysis of the data has demonstrated an increase in the percentage of adult...