relationships over time; that physical care tended to take priority over time spent exploring issues especially when time was limited. Reasons for not exploring and documenting were common for all staff groups.

Conclusions and recommendations The results were presented to the wider hospice team who made suggestions for change. These included: offering additional staff education and training in communicating with cognitively impaired people; proactively consulting informal carers to gain a deeper understanding of patients’ needs; the use of a person-centred document for informal carers to complete with patients if able, that encouraged identification and exploration of psycho-social concerns; using routine handovers to highlight when these areas had not been addressed. A working group will be formed to implement and test changes to practice as part of the quality improvement in this area.

Abstracts

90 EMERGENCY ADMISSIONS FROM CARE HOME TO HOSPITAL AT THE END OF LIFE: AN ANALYSIS OF NATIONAL DATA 2006–2015 FOR ENGLAND

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Objective Hospital admissions of care home residents close to the end of life are a source of concern and frequently described as ‘inappropriate’ and avoidable. However, the magnitude of this ‘problem’ on a national scale has not been investigated to date. This recently completed study investigated trends and characteristics of emergency admissions from care home to hospital where death occurred within seven days.

Methods Longitudinal and cross-sectional analysis of linked Hospital Episode Statistics with ONS mortality data for England 2006 to 2015: 1,858,830 care home residents aged 25 + who experienced a last week of life emergency admission to hospital, with comparator group 

91 HOW THE TRACEY JUDGEMENT HAS CHANGED DNACPR DISCUSSIONS IN PALLIATIVE CARE

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Background The decision published from the Court of Appeal of England and Wales in 2014 in the case of Tracey vs Cambridge University Hospital (‘the Tracey judgement’) changed the requirements for discussing Do Not Attempt Cardiopulmonary Resuscitation (DNACPR) decisions with patients.

Methods A retrospective case note review aiming to identify changes in practice when discussing DNACPR decisions following the Tracey judgement. 150 case notes from 2013 (before the Tracey judgement) were compared with 150 case notes from 2015 (following the Tracey judgement) at five hospices in the West Midlands. The notes were analysed to identify whether the judgement had resulted in a change in how frequently DNACPR decisions were discussed with patients or their families, as well as any changes in the rationale for not discussing such decisions.

Results Discussions with patients increased from 31% to 60% and with relatives from 29% to 39%. The most common reason that DNACPR decisions were not discussed with patients or relatives changed from a belief that these discussions would cause distress in 2013 (23%) to the patient lacking capacity to engage in such discussions in 2015 (40%). There was a lack of consistency and clarity in defining the concept of ‘physical or psychological harm’ following the Tracey judgement.

Conclusion Healthcare professionals specialising in palliative care in the region and time frame studied are discussing DNACPR decisions more frequently with patients and families following the Tracey judgement but clarity on what constitutes ‘physical or psychological harm’ caused by these discussions is still required.

92 AN AUDIT OF CURRENT GUIDELINES AND RECOMMENDATIONS FOR MOUTH CARE IN PALLIATIVE AND END OF LIFE CARE

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Background Oral complications in palliative care patients are very common. Guidelines for oral care of these patients are available from several different sources within the UK. However, these guidelines are often inconsistent and it is unclear whether the recommendations are implemented in practice. This project aims to determine if the guidelines for oral care translate into the care provided to patients, and identify any additional techniques that are not published but provide benefits to patients in end of life care.

Methods A literature review of the guidelines and published evidence was conducted to collate a list of recommendations. A pilot survey and qualitative research was conducted with Special Care Dentistry Consultants, Palliative Care consultants, geriatricians, and those providing day-to-day care for palliative care patients. This focused on what treatment was being provided to maintain oral care and comfort for palliative care patients.
Abstracts

93 PATTERNS OF GENERAL PRACTICE PRESCRIBING IN THE LAST YEAR OF LIFE

Amelia Hartfield, Rupert Payne, Stephen Barclay, University of Cambridge; University of Bristol

Background Little is known about patterns of prescribing in the last year of life in primary care.

Methods Recently completed analysis of UK Clinical Practice Research Datalink (CPRD), a large database extracted from GP records that is representative of the UK population. Prescribing data for the last 12 months of life were extracted from 1 18 571 patients’ GP records who died between September 2010 and August 2015. Two clinicians used British National Formulary codes to identify eight palliative care drug categories: antiemetics, anti-secretory, benzodiazepines, steroids, strong injectable opioids, strong patch opioids, strong enteral opioids, and weak oral opioids. The percentage of patients prescribed each medication was calculated at weekly intervals during the year before death, further stratified by cause of death and age at death. Random effects logistic regression models were run adjusted for age-group, sex, and causes of death.

Results Levels of prescribing significantly increased for all drugs at time of death compared with one year before death (with the exception of weak oral opioids and steroids where results were inconsistent across ages, genders, and causes of death). The largest increase was for 90+year old male patients prescribed strong injectable opioids (OR: 21032, 95% CI: 3499 to 12641) and the greatest decrease for females prescribed steroids who died of external causes (OR: 0.52, 95% CI: 0.34 to 0.70). Patients who died of cancer generally showed the most marked increase in prescribing towards the end of life compared with other causes of death.

Conclusions Primary care patterns of prescribing in the last year of life has not been previously investigated. The findings of this recently completed study of a large nationally-representative dataset reveal increasing GP prescribing in response to the rising symptom burden of all patient groups as death approached. The implications for clinical practice and future research will be discussed.

94 GUIDELINE DEVELOPMENT FOR THE MANAGEMENT OF AGITATION IN THE LAST WEEKS OF LIFE

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Background Agitation is a distressing and common symptom at the end of life; despite this the evidence for the management of this symptom is poor.

Aim
- Audit clinical practice around management of terminal agitation with patients and healthcare professionals (HCPs) against regional standards.
- Use audit outcomes to update regional standards and guidelines

Method
- Systematic literature review examining the evidence base.
- Survey of HCPs working in specialist palliative care.
- Retrospective case note review of patients with a diagnosis of terminal agitation receiving specialist palliative care in hospital, hospice or community settings.

Results Below is a summary from 185 responses from 11 different healthcare settings in the Merseyside area from retrospective case note review:
- Hospice 54.5%, Hospital 29.7%, Home 11.8%, other including Nursing homes 3%.
- Main reversible causes documented included: pain; breathlessness; constipation and urinary retention.
- 94% of patients had medications reviewed.
- Non-pharmacological interventions formed part of the overall management plan.
- 82% of patients had psychological and spiritual support offered to the patient and or family.
- First line pharmacological intervention is Midazolam, Levomepromazine is second line.
- In 92% of cases a Syringe driver (CSCI) was used.
- Majority of deaths occurred in 3 days or less
- Communication with patient and family formed a key theme

At the time of death:
- Midazolam prescribed in 146 cases (mean 26 mg – range 0–60 mg).
- Levomepromazine prescribed in 61 cases (Mean 56.8 mg – range 5–250 mg).
- Haloperidol 25 cases (Mean 3.5 mg – range 1–8 mg).
- Phenobarbital 9 cases (Mean 922 mg – range 200–1200 mg).

Conclusion This Audit revealed the challenges and wide variation in practice with regards to management of terminal agitation. These findings have informed an update of the regional palliative care guidelines.

95 THE ‘SUPPORT OF CARE CYCLE’: INTEGRATING ETHICS INTO HEALTHCARE PROFESSIONAL SUPPORT AND DEVELOPMENT IN HOSPICES

Craig Gannon, Princess Alice Hospice

Background High-quality hospice care is dependent on an expert caring and resilient workforce who can cope with the