**Abstracts**

**P-114**  
**MORTALITY AUDIT OF CARE HOME PATIENTS KNOWN TO THE COMMUNITY PALLIATIVE CARE TEAM**  
Maxine Concannon, Michelle Tombs, Claire Capewell. St Catherine’s Hospice, Preston, UK  
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**Background**  
The percentage of patients dying in care homes is steadily increasing, with 70% of permanent residents dying in their care home (Public Health England, 2017). Therefore, it is important to understand what is happening to the local population of care home residents at the end of life and their preferences for care.

**Aims**  
To assess the impact of involvement of Specialist Palliative Care, for care home patients, on deaths in usual place of residence, hospital admissions in last 90 days of life as well as length of stay in hospital in the six months prior to death. To assess the impact of advance care planning.

**Methods**  
A retrospective review of patients who were resident in care homes was undertaken. These patients were known to the Community Palliative Care Team and died between 16 September and 16 December 2018. The electronic patient records were used to gather the data, alongside requesting information from Business Intelligence at the local hospital Trust to obtain information regarding hospital admissions.

**Results**  
74% of patients achieved their Preferred Place of Death (PPD) in this audit. 2/3 of those not achieving PPD had requested hospice transfer. 89% died in the care home and only 4% died in hospital. However, 41% of patients had three or more admissions to hospital within the last six months of life. 93% had documented evidence of having a DNACPR (Do Not Attempt Cardio-pulmonary Resuscitation) form in place and there was evidence of advance care planning for 85% of patients.

**Conclusions**  
Further exploration is required around the reasons these patients are being admitted to hospital. Our action plan includes signposting care homes to available education, sharing learning with generalist colleagues and encouraging After Death Analysis at GSF meetings.

**Patient Care**

**P-115**  
**INTRODUCTION OF NEWS2 IN A HOSPICE SETTING**  
Helen Ireland, Kate Davies, Jane Dochane, Bethany Wright, Gaye Senior-Smith. St Peter’s Hospice, Bristol, UK  
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**Background**  
Following a clinical incident there was a review of the current system for recording and reporting patient observations. This involved an initial audit of current practice and scoping of regional and national practice.

**Aims**  
The project aim was to provide all clinical staff with a tool to identify critically unwell patients and respond safely and appropriately.

**Methods**  
The project team looked at the local hospitals NEWS (National Early Warning Score) charts and embarked on designing a bespoke chart for the hospice setting. Following several rounds of consultation and different iterations it happened that the NEWS 2 (Royal College of Physicians, National Early Warning Score (NEWS) 2, 2017) tool was released nationally. This led to further review and highlighted our error in trying to adapt a nationally recognised tool which was enabling common language, transferability and standardisation across different settings.

**Results**  
Following cascade training to all clinical staff, NEWS 2 is now embedded in use in the Inpatient Unit. Staff have said it ‘highlights when we should respond to triggers’ and it ‘helps decision making overnight and when the medical on call needs to be contacted’. Weekly snapshot audits have also shown examples of clearer documentation regarding planned frequency of observation and clinical response and allowed assessment and teaching at the point of care. A more formal audit is being undertaken in June 2019 and the results of this will also be reported.

**Conclusion**  
Despite some initial resistance to change we have shown that it has been possible and acceptable to introduce the use of NEWS 2 in the hospice inpatient setting. Following this it is now being rolled out to other departments including Day Services and the Clinical Nurse Specialist Team.

**P-116**  
**WHAT ARE HOSPICE STAFF PERCEPTIONS OF CARING FOR PATIENTS WITH APHASIA?**  
1Louise Shelley, 2Janet Holt. 1Leeds Teaching Hospitals NHS Trust, Leeds, UK; 2University of Leeds, Leeds, UK  
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**Background**  
Aphasia is an acquired disorder of language and can be caused by conditions including stroke, brain tumour, head injury and dementia (Royal College of Speech & Language Therapists, 2019). Due to the nature of these conditions, it is likely that people with aphasia will access palliative care (Hawksley, Ludlow, Buttmer &Bloch, 2017).

Speech and language therapists (SLTs) are well placed as ‘communication experts’ to support patients with aphasia to communicate. Although the role of SLTs in palliative care is well documented (Chalda, Mathisen & Carey, 2017) the literature suggests that SLTs are not usually integrated within a hospice MDT (Pollens, 2004).

**Aim**  
To identify views of hospice staff on their interactions with patients with aphasia.

**Research questions**
- Do staff have an understanding of SLT and do they feel SLTs have a role in palliative care?
- Do staff feel confident in interacting with patients with aphasia and/or do they feel that service improvements need to be made to improve these interactions?

**Methods**  
A qualitative descriptive exploratory design (Sandeflowski, 2000). Four doctors and four nurses working in hospices in Yorkshire were interviewed using a semi-structured interview guide. Hospices which identified that they commonly care for patients with aphasia were included. A purposive sampling strategy was adopted. Data were analysed using thematic analysis (Tesch, 1990).

**Results**  
The following themes emerged:
- Knowledge of and attitudes towards speech and language therapy;
- Knowledge and awareness of supporting people with communication impairments;
- Palliative care in the hospice;
- Using family to support;
- Directions for change.
Differences were apparent between the two hospices in the frequency of referrals to SLT. Staff had skills in caring for patients with communication impairments but some spoke of a desire for further training and closer links to SLT.

Conclusions Findings from this study and the literature suggest limited specialist communication support for patients accessing palliative care. Clinicians are encouraged to reflect upon the approach in their own setting to consider if changes could be implemented.

P-117 IMPLEMENTING A HOSPICE TRAINING PROGRAMME FOR THE MANAGEMENT OF PATIENTS WITH AN ALTERED AIRWAY

Joanne Reynolds, Liz Reed, Princess Alice Hospice, Esher, UK

Background Only 40% of hospitals demonstrated good practice in the care of patients with tracheostomies. Six recommendations have been made to improve the care they receive. These include having a protocol and mandatory training to ensure staff are competent in recognising and managing common airway complications (National Confidential Enquiry into Patient Outcome and Death, 2014). There is little research or evidence about the quality of care patients with an altered airway receive in adult hospices.

These patients usually have complex care needs and present a significant challenge for hospices in that staff are not caring for patients with an altered airway regularly enough to maintain competence and confidence in this area.

Aims In-patient (IPU) clinical staff will be competent and confident caring for patients with altered airways.

Intervention Mandatory, continuous education programme titled ‘Essentials in tracheostomy and laryngectomy care in the hospice setting’.

Methods

- Pre- and post-education intervention on-line survey to all clinical staff undertaking the educational intervention;
- Practical simulation of the care of patients with an altered airway;
- Assessment of confidence and competence in caring for patients with an altered airway.

Findings This educational intervention is on-going. Analysis of the accumulative data will be undertaken in September 2019 and reported in the proposed paper.

Conclusion This appears to be the first evaluation of its kind; addressing retention of knowledge and assessment of competence/confidence in an adult hospice caring for patients with an altered airway.

P-118 INVESTIGATING ANAEMIA AND TARGETING TREATMENT IN A HOSPICE POPULATION: A QUALITY IMPROVEMENT PROJECT

Tom Steele, Helen Bonwick, Laura Chapman. Marie Curie Hospice Liverpool, Liverpool, UK

Background Anaemia is common in palliative care. Results of the national audit of blood transfusion practice in hospices, published (online first) last year, recommended more rigorous investigation of anaemia and increased use of alternative therapies (Neoh, Gray, Grant-Casey, Estcourt et al., 2019). This project aimed to improve the proportion of patients with anaemia at Marie Curie Hospice Liverpool having haematologic investigations and receiving appropriate, targeted therapies.

Methods Several changes have been implemented through this project. To raise awareness, education sessions were delivered and practical written guidance on performing and interpreting haematologic investigations disseminated. A protocol was implemented to ensure outpatients attending for blood transfusions had investigations completed. Finally, an electronic template was developed to prompt investigating and recording results. Between August 2018 and February 2019 continual retrospective analysis was performed using laboratory and patient records. All patients who had haemoglobin checked at the hospice had clinical details, haematologic tests (ferritin, iron studies, B12 and folate; if performed within 6 weeks before or 2 weeks after) and resulting treatments recorded.

Results 125 patients with anaemia were assessed over the project (excluding repeat sampling). Amongst those having blood transfusion, assessment of both iron status and B12/folate improved from 2/12 (17%) in the first two months to 7/7 (100%) in the last two months of the project. Including all patients with Hb <100 g/dL, iron status assessment improved from 7/22 (32%) to 12/18 (67%) and B12/folate from 5/22 (23%) to 9/18 (50%). Results led to alternative therapies in 13 cases; oral iron (n=5), intravenous iron (n=6) and folinic acid (n=2).

Conclusions This quality improvement project has led to more patients being investigated for underlying causes of anaemia and increased usage of targeted therapies. Research is required to establish if this approach impacts symptoms or transfusion utilisation and the role of specific treatments, particularly intravenous iron therapy, in this population.