Results 410 ultrasounds were performed, 3 (<1%) of these were in the patient's own home. All scans were performed by a Palliative Medicine Consultant or Clinical Fellow trained in use of FASP.

Clinical indication was broadly split into malignant and non-malignant, 64% vs. 36% respectively.

Identified indications include: ascites 74% (302); urinary retention 19% (78); constipation or overflow 5% (9); deep vein thrombosis 1% (5); pleural effusion <1% (3); assessment of mass <1% (2) and spontaneous bacterial peritonitis <1% (1). 41% of the 302 ultrasounds for ascites, resulted in paracentesis. Major and minor complications rates, 0% vs. 8% respectively, were reviewed over 4 weeks. Failure of procedure was the highest minor complication rate at 2%. Persistent leakage from drain site, infection, local skin reaction and an asymptomatic drop in blood pressure had complication rates of <1% each. Data trends show the number of ultrasounds performed each year is decreasing, however, paracentesis rates remain relatively stable, suggesting more targeted use.

Conclusions Community ultrasounds reduce hospital transfers, admissions and need to access hospital-based radiology services. With appropriately selected patients, the low complication rates, support continued use of community paracentesis. Patient views and preferences on the location of performing ultrasounds and procedures would be valuable. This combined with projected cost savings, has the potential to demonstrate and endorse the importance of these community-based services.

Conclusions

This is the first study to determine epidemiological local data on the foregoing. There is clear variance in practice. We suggest that all samples should have biochemical analysis (transudates are to be given prophylactic antibiotics) and all patients should be referred for consideration of IPC.

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126 A SCOPING REVIEW OF THE IMPLEMENTATION OF NON-MEDICAL DEVICES FOR THE SELF-MANAGEMENT OF CHRONIC BREATHLESSNESS

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Background Non-medical devices such as the handheld fan (fan), mobility aids and inspiratory muscle training (IMT) devices offer important benefits for patient self-management of chronic breathlessness. We examined the published evidence regarding barriers and facilitators to implementation of non-medical devices for the self-management of chronic breathlessness from the perspectives of patients, carers and clinicians.

Methods MEDLINE, EMBASE, SCOPUS, EBSCO and the Cochrane Database of Systematic Reviews were searched. Papers were imported into EndNote and Rayyan for review against a priori eligibility criteria. These were: i) any study design including randomised controlled trials, observational, interviews, primary and secondary analyses of data, ii) patients; adults with chronic breathlessness, carers', and clinicians, iii) exposure to the fan, mobility aids and IMT devices, iv) and outcome data regarding implementation of these devices for self-management of chronic breathlessness.

Data relevant to use were extracted and categorised as potential implementation barriers and facilitators, and a narrative synthesis exploring reasons for similarities and differences conducted.

Results Seven studies were included; five fan and two mobility aid (wheeled walkers with seats) studies. All of the studies presented patient use of non-medical devices only. The fan was implemented for self-management of chronic breathlessness at home whereas mobility aids were used mainly for outdoor activities. Key barriers to implementation of non-medical devices were: appearance; credibility; self-stigma; technical specifications. Common facilitators were ease of use and feeling safe and secure with the device.

Conclusion Limited evidence exists for the implementation of non-medical devices by patients for the self-management of chronic breathlessness. We highlight the lack of studies on carers, clinicians, IMT devices and the use of applied implementation science theory. Further implementation research is...
needed to explore the interactions between patient, carer and clinician perceptions of non-medical devices and, ultimately whether this influences the benefits for patient self-management of chronic breathlessness.

### 127 IMPROVING THE IDENTIFICATION AND MANAGEMENT OF CONSTIPATION ON THE PALLIATIVE CARE UNIT, WANSBECK GENERAL HOSPITAL

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**Background** In 2020 the palliative care team at the inpatient palliative care unit at Wansbeck General Hospital identified the notable burden of constipation on patients, and the barrier to symptom improvement it caused. Our team designed a quality improvement project to identify and tackle this issue.

**Aims** Our team aimed to assess the documentation, identification and management of constipation on the unit. This was undertaken via instigating health care education, and (2) developing a patient information leaflet to empower patient control of constipation.

**Methods** A 8 month QI project was undertaken on the unit, assessing the written and NerveCentre documentation of bowels, prior to and following the instigation of staff education and patient leaflets, promoting patient lead documentation of their constipation. This project followed several PDSA cycles and identified an initial absence of patient perception of autonomy over constipation, as well as prescriber’s limited knowledge of the regional guidance. The data from the measures gathered continuously using an electronic pro forma, with alternations following each cycle with further staff and patient education, and alternations to the language of the leaflets. A flow chart was generated following the measures indicating response to intervention.

**Conclusion** The prevalence of overall constipation halved from 46 to 22%, with an increase on the NerveCentre documentation of 19%. In those for whom constipation was appropriately identified 100% used the regional guidance to manage this. It was noted however that following the initial spike of identification of constipation this deteriorated after the change was instigated, demonstrating the need for ongoing education to both patients and staff.

### 128 WHAT ENABLES AND CONSTRAINS SHARED DECISION-MAKING ABOUT OPIATES FOR PEOPLE LIVING WITH CHRONIC BREATHLESSNESS? A SYSTEMATIC NARRATIVE REVIEW AND FRAMEWORK ANALYSIS OF PUBLISHED DATA USING A PROGRAM THEORY OF SHARED DECISION-MAKING

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10.1136/spcare-2021-PCC.146

**Background** Despite the evidence supporting the use of opioids to manage chronic breathlessness, these are rarely prescribed, indicating the need for enhanced SDM in this field. The Waldron 2020 SDM program theory explores the relationship between various factors in patients’ and healthcare professionals’ (HCPs) engagement in SDM. This systematic narrative literature review assesses enablers and constraints of SDM in the prescription of opioids for chronic breathlessness and evaluates whether the Waldron’s SDM program theory can explain the operation of these factors.

**Methods** A literature search was conducted on Medline, Embase, PsychInfo, Cochrane Database of Systematic Reviews, CINAHL, Scopus, and Web of Science (1946 “ June 2020). Studies were selected if they took place in Europe, Canada, or Oceania, and addressed an element present in decision making for the management of chronic breathlessness with opioids. Thematic analysis identified patient and HCP factors involved in the prescribing of opioids for chronic breathlessness. Their effect on SDM was assessed using Waldron’s SDM programme theory.

**Results** Six themes were identified: concerns with side effects, knowledge of opioid use for breathlessness, acceptability, prior experiences, awareness of symptom severity, and system support. HCPs were reported to have more negative attitudes towards opioid use for breathlessness, particularly in the context of COPD, than patients. HCP’s support by the MDT improved engagement in SDM. Open, clear communication addressing patients’ questions and concerns, alongside HCP’s revisiting of the discussion enabled SDM. A simple change in attitude from either the patient or HCP could be determining to increasing engagement with SDM.

**Conclusions** Research is scarce on SDM in opioid prescription for breathlessness. Limitations include the lack of quality appraisal of evidence. The Waldron SDM program theory appropriately supports SDM in this context. This review informs patient-centred care on management of chronic breathlessness with opioids through SDM, and highlights gaps in the literature.

### 129 BALANCING PATIENT BENEFIT AND RISK OF RESISTANCE: AN AUDIT OF HOSPICE ANTIMICROBIAL PRESCRIBING

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**Background** Antimicrobial stewardship aims to improve the safety and quality of patient care and reduce the risk of antimicrobial resistance. Symptoms of infection are often part of the natural deterioration towards end of life. Clinical evidence for the benefit of antimicrobials at the end-of-life is scarce and results inconclusive. Within our hospice, our aim of prescribing effectively whilst observing the principles of advance care planning led to an audit of practice as part of our antimicrobial stewardship.

**Methods** A retrospective audit was completed of all patients admitted over 3 months, where antimicrobials had been prescribed, considering whether antimicrobials were prescribed in accordance with local trust guidelines; presence of indication, review and stop dates on each prescription and whether advance discussions were documented in relation to the use of antimicrobials.

**Results** Of 56 admissions identified, 23 (41%) patients had 37 individual antimicrobials prescribed. The commonest sources of infection were chest (18), urinary tract (4), skin and soft tissue (4).