what services are offered to meet which needs; communicating regularly with referrers, and providing education or training to support referrers to improve their ability to meet patient needs directly.

### REVIEW OF POTENTIAL GEOGRAPHICAL DISPARITIES IN ACCESS TO ANTI-FIBROTIC MEDICATION AND CLINIC-BASED SPECIALIST PALLIATIVE CARE FOR PATIENTS WITH IDIOPATHIC PULMONARY FIBROSIS

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**Background**

Idiopathic pulmonary fibrosis (IPF) is a progressive interstitial lung disease of unknown origin (NICE, 2013). Disease-modifying anti-fibrotics are restricted to patients who meet specific criteria and are prescribed by regional IPF centres (NICE, 2016; NICE, 2018). Patients should be offered best supportive care from diagnosis (NICE, 2013). In January 2016 specialist palliative care (SPC) support was embedded in a regional IPF clinic. Initial service review (Kavanagh, 2017) suggested potential geographical disparities in access to both anti-fibrotics and clinic-based SPC.

**Aim**

To repeat the 2016 review to assess whether geographical disparities have changed over time.

**Methods**

The sex, age and postcode of all patients with known IPF prescribed anti-fibrotics or seen by clinic-based SPC between January 2016 and April 2019 were collected retrospectively and geographically plotted. Chi squared goodness of fit was used to calculate statistical significance.

**Results**

398 patients were started on anti-fibrotics (male:female 330:68, mean age 74.7). Geographical plotting shows significant variation by region (P<0.01) ranging from 6.1 to 16.2 per 100,000 population, with comparatively low numbers of patients from Cumbria and North Durham (See figure 1). 80 patients were seen by SPC (male:female 65:15, mean age 74.9). Again, geographical plotting shows significant variation by region (P<0.02) ranging from 0.9 to 4.5 per 100,000 population, with a higher proportion of patients from Newcastle-Gateshead (See figure 2).

**Discussion**

Geographical distribution of patients seen by clinic-based SPC appears to have changed little since 2016, with clear centralisation to Newcastle-Gateshead. Speculatively, reasons for this could include travel distances and local palliative care provision. Geographical disparity in anti-fibrotic prescription is more difficult to account for and requires further investigation.

**REFERENCES**


### FAMILY-CENTRED CARE IN HOSPICES: WHAT ABOUT THE CHILDREN?

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Palliative care professes to take a family-centred approach to patient care. However, children under 18 may not feel included when a parent is being cared for in a hospice. A systematic literature review of PsychINFO, Medline, Embase, Scopus and Web of Science was conducted, with no restriction on publication date. Nineteen studies met the inclusion criteria, all reporting primary data from children under 18, whose parent has a life-limiting illness. Findings from the review reveal that:

- Children and young people often feel excluded when a parent is dying.
- Communication about prognosis does not routinely happen with children.
- For children, the terminal period is generally more stressful than after the death.
• There are discrepancies between the support children would like from professionals and
• The support they typically receive.

Recommendations highlighted by the review:
• Regular training for all hospice staff in relation to working with children – including recognition of how parental illness may affect children, the importance of communication, and children’s understanding of death at different ages.
• Consideration of how children can be more routinely involved, for example inclusion in family meetings.
• Encouraging staff to explore patients’ wishes and feelings in relation to their children.
• Consideration of whether the hospice environment is ‘child-friendly’.
• Flexibility around working hours – children are often ‘invisible’ and visit parents in the evening and at weekends.
• Consideration of creative methods of including children, perhaps by using technology.

Hospices are encouraged to consider these recommendations and promote a model of family-centred care which includes children under 18.

Background

There has been rapid growth in older caregivers (OCs) age 65+ in the UK. Playing a fundamental role in the life of their care-recipient, helping to sustain preferred place of care, with many delivering over 50 hours of weekly care. OCs may themselves have age and caring role associated health issues. Spending cuts mean fewer now qualify for support. Understanding the breadth and extent of OCs unmet needs can inform support to help OCs maintain their wellbeing and caring role.

Aim

To identify and describe the unmet health and social care needs of caregivers aged 65+ in the UK.

Methods

A systematic search was undertaken across a wide range of databases and grey literature from inception to February 2018. Two reviewers undertook study selection, data extraction and quality assessment. Qualitative Metasummary was used to synthesise findings.

Results

In total 10 of 4407 identified records satisfied the inclusion criteria. Eight unmet need domains were identified, many were interrelated and displayed interplay: psychosocial, service, informational, constancy of care, future care planning, physical health issues, reluctance to utilise services, and financial needs.

Conclusions

The review has identified eight domains of unmet need that may impact on OCs wellbeing and ability to care. OCs are a heterogeneous group with wide-ranging support needs, hence personalised carer’s needs assessments are vital. Services need to be accessible, flexible and sufficiently available. Service information should be actively promoted. An extensive research gap on the unmet needs of OCs has been exposed.

Systematic Review Registration: CRD42018086759.

REFERENCES


52 EXPERIENCES OF SUPPORTING NUTRITION AND HYDRATION FOR SOMEONE LIVING WITH DEMENTIA AT THE END-OF-LIFE: A SYSTEMATIC REVIEW OF QUALITATIVE EVIDENCE

Background

People living with dementia often face eating and drinking difficulties as life is ending. Family and friends find this emotionally challenging and professionals also struggle to provide information and support. The aim of this systematic review was to 1) explore the views of people with dementia and those involved in supporting eating and drinking for someone with dementia (paid and unpaid carers) at the end of life; and 2) explore views and experiences of artificial nutrition and hydration (ANH) for people with dementia.

Methods

A systematic search was conducted in Medline, CINAHL, Embase and PsycINFO, for English-language qualitative studies published between 2000 and 2019, and forward and backward citation tracking. Studies were synthesised using narrative synthesis and thematic analysis methods.

Results

18 papers were included with six focussing on carers’ and 13 on professionals’ views. Professionals and carers reported on challenges associated with supporting nutrition and hydration and several strategies used to facilitate oral intake (i.e., modifying the environment and food provision). When ANH was considered, six main factors were identified as influencing decisions: 1) personal perceptions; 2) national context; 3) understanding of the natural dying process; 4) involvement of different parties (i.e., professionals in multidisciplinary teams, carers and the person living with dementia); 5) lack of information; and 6) prognosis.

Discussion

Decisions around eating and drinking and the introduction of ANH require clear and respectful communication among all involved (including those living with dementia), as their personal/professional experiences, perceptions and knowledge impact on their behaviours and emotions.