palliative care services. Final results are due to complete by June 2013.
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A Hospice Motor Neurone Disease Clinic: Challenges and Benefits of a New Service

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Background/context The Hospice in North Hampshire had offered support and care to MND patients but in an unstructured way. Close working between the Palliative Medicine and Neurology Consultants led to a recognition that both early and advanced care could be improved if services were to work more closely together.

Aim The aim was to set up and run a monthly multi-disciplinary clinic for people with MND in the Hospice, allowing access to all members of the multi-disciplinary team (neurologist, consultant in palliative medicine, care centre co-ordinator, hospice at home team, social worker, physiotherapist, OT, speech and language therapist, Association Visitors) via one clinic. The clinic link with the MNDA would improve access to information about services. The outpatient hospice setting allows informal support for pts and carers from Assoc Visitors.

Outcomes To improve the patient and carer experience.
- To ensure access to hospice services early in diagnosis.
- To improve both peer and carer support through joint working with the MNDA.
- To develop a specialist skill base with opportunities for support and training of other professionals.
- To develop patient professional relationships which allow easy transition in the end stages of illness.

Application to hospice practice Many centres across the country run a multidisciplinary MND clinic but traditionally in the hospital setting. Evidence is good for improvement in patient care from a hospital model of service delivery model. Hospices have many advantages with their experience in coordinating care and community focus. The early links with the Hospice allow continuity of care throughout the patient journey.

Changing a Lymphoedema Service through Service Redesign

Debbie Murphy. St Catherine’s Hospice, Preston, UK

Background Very well established Lymphoedema service facing number of escalating issues:
- Increasing referrals
- Over budget
- Frustrated staff
- Lack of care pathways and long term management
- Lack of innovation

Aim
- Reduce financial strain
- Provide clear care management strategy
- Introduce self-management strategies
- Ensure full access for individuals with Lymphoedema.

Approach Used
- Poor team work - looked at strengths and the weaknesses of each team member as a team.
- Cost reduction – compression garment formulary assessed on cost effectiveness, size of range, availability on FP10, education supplied by the company.
- Care pathways – patient journey plotted dependent on severity of condition.
- Discharge policy developed, packs sent to the GP and to patient.
- Long term self-management - six week education programme combined with fun exercise class (Le Bed), gives patients tools to manage own condition.
- Following discharge from Le Bed, Lymphoedema Information Fun Event (LIFE) established after feedback from patients - provides ongoing information about service development. Group used as patient reference and involvement group e.g. feedback on leaflets etc.
- Investment in new technologies (lymph assist machines, lasers, oscillator) - enhances treatments offered, produces quicker results.
- Kinesio tape introduced
- Children’s Lymphoedema Service initiated
- Bra fitting service developed.

Outcomes
- Greater sense of team - clear lines of reporting, goal setting and monitoring.
- Expansion of team with a clear mix of skills
  - support worker undertaking more routine work, specialist nurses concentrate on more complex cases
  - part time specialist appointed
  - Lymphoedema physiotherapist (still to be appointed)
  - training and development - manual lymphatic drainage, Le Bed instructor training, bra fitting, wound management, child protection training etc.
- Number of new patient referrals has increased year on year (see table)
- Negotiated and secured contracts with three additional CCG’s
  - 2008/2009 = £23k overspend, 2011/2012 break even (including reinvestment in new technologies and education)
  - Number of intensive treatments increased from 27 in 2010/2011 to 119 in 2012/2013
- Patients self-managing - the discharge pathway and Le Bed class – leading to greater self esteem
- Lymphoedema manager recognised as Lymphoedema nurse of the Year 2013, British Journal of Nursing

Think Again! Treatment Changes for Patients with Head, Neck and Facial Swelling

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Background Few patients referred for head, neck and facial lymphoedema as widely viewed that very little could be done, apart from prescribing tight fitting compression garments which are very constrictive and have a low patient compliance.

Aim
- Improve access for patients experiencing lymphoedema to the head, neck and face.
- Identify alternative treatment methods.
Meaning of pain in advanced cancer is inextricably linked with loss and bereavement. Meaning of pain is a dynamic, temporal process underpinned by the following themes: (i) nature of cancer-related pain, (ii) multidimensional impact of pain on patients’ function and behaviour, (iii) ‘Web of Loss’: a complex network of losses that perpetuates cycles of further loss and (iv) hope and appreciation of life. The data suggests meaning of pain has an indirect influence on behaviour mediated by the coping strategies patients adopt. Meaning of pain has potential to promote both constructive and maladaptive responses in functional behaviour.

Conclusions Findings further our understanding of the experience of cancer-related pain and inform bio-behavioural approaches to pain management in palliative care. Recommendations include (i) tailoring pain management to dedicate greater attention to the behavioural dimension of cancer-related pain (ii) timely identification and management of preparatory grief in patients with advanced cancer and (iii) promotion of constructive coping strategies to support patients to make sense of their pain and maintain functionality within the limitations of advancing disease.

P162 LOST IN TRANSLATION: PHYSIOTHERAPISTS’ ATTITUDES AND BELIEFS TOWARDS PALLIATIVE CANCER PATIENTS IN THE ACUTE HOSPITAL SETTING

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Abstract Aim The study aimed to elucidate any attitudes acute hospital physiotherapists may have towards palliative cancer patients.

Background With advances in medical technology, increasing numbers of patients are ‘living with’ a terminal cancer diagnosis for longer periods; this coupled with changes in government policy, is resulting in patients being treated closer to home rather than in specialist centres, bringing physiotherapists into increasing contact with this patient group.

Methods Semi-structured interviews (analysed utilising Colaizzi’s method of analysis), of the first twelve physiotherapists from an acute hospital in the UK who volunteered for the study and who met the inclusion criteria.

Results Two clusters of themes were identified, namely Attitudes and Issues. There were four sub-divisions of Attitudes; terminology, physiotherapists’ perceptions, other professionals’ perceived attitudes towards physiotherapists, and patients and their families perceived attitudes towards physiotherapists’ role with terminally ill cancer patients.

“I’d say there are a lot of physio’s that think it’s probably a little bit pointless”

“try and maintain their quality of life ... by their thinking... and hopefully try and meet it.”

There were six sub-divisions of issues: communication, education, emotions, boundaries, referring onward and time.

“I think they’re [physiotherapists] scared that that’s going to be asked of you...will I walk again?”

Conclusions Some physiotherapists understand and apply their skills to improve the quality of life of palliative cancer patients. However it is suggested that the majority of physiotherapists regard these patients as ‘dying from’ rather than ‘living with’ a...