The Palliative Care needs of people with Motor Neurone Disease (MND) and their informal carers: a systematic review of qualitative research

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Project team

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• Prof Ian Watt – Professor of Primary and Community Care
“The hospice movement has largely concentrated on cancer; we need to look at other things, at what we can do for people with strokes and motor neurone disease. We need to go on learning so that in 10 years’ time we are doing things better than we are now”

Dame Cicely Saunders 2002

https://cicelysaundersinternational.org/dame-cicely-saunders/cicely-saunders-institute
MND

- Motor neurone disease (MND) or Amyotrophic Lateral Sclerosis (ALS) is a neurodegenerative disease
- It has a sudden onset, rapid progression and brings with it the potential for complex and disabling symptoms and care needs
- It is currently without cure, with the average time between diagnosis and death being two to three years
- The progression of symptoms and subsequent onset of disability are likely to occur over weeks and months rather than years.
- Delays in diagnosis can count for a significant proportion of total illness duration
The need for palliative care

- Up to 20% of people with MND opt for physician-hastened death in countries where this is a legal option (Maessen et al 2009)

- People diagnosed with MND and their caregivers often fear the dying process (Whitehead et al 2011)

- These fears are linked with choking to death, pain, loss of body control and breathlessness (Oliver et al 2007)
Why a systematic review?

**What** - A systematic review summarises the results of available good quality studies

**How** - Uses a systematic approach to searching for studies and the methods used to synthesise (combine) them together

**Why** - Aim to generate a greater understanding from existing research by considering connections, similarities and differences between each study.

**The result** - The combination of research can provide more information than the findings of each study in isolation.
Why qualitative research?

• Qualitative research is a methodology that explores the perceptions and views of individuals.

• Commonly used in health care to understand more about people’s experiences of their situation, illness or services.

• Synthesising qualitative research can help maximise the contribution of the primary qualitative studies to the evidence base for practice.
What we did

• Searches were run in: CINAHL, MEDLINE, PsycINFO, Social Science Citation Index
• From 403 results we included 41 papers in the review
• Included the views of 358 people with MND and 369 carers

<table>
<thead>
<tr>
<th>Profile</th>
<th>People with MND</th>
<th>Carers</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age range</td>
<td>21-85</td>
<td>25-86</td>
</tr>
<tr>
<td>% female</td>
<td>45</td>
<td>65</td>
</tr>
<tr>
<td>Time to diagnosis</td>
<td>3-60 months</td>
<td>-</td>
</tr>
<tr>
<td>Type of carer</td>
<td>-</td>
<td>78% partner</td>
</tr>
<tr>
<td></td>
<td></td>
<td>16% child</td>
</tr>
<tr>
<td></td>
<td></td>
<td>6% other (eg parent)</td>
</tr>
<tr>
<td>Time as a carer</td>
<td>-</td>
<td>6-96 months</td>
</tr>
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</table>
## International pool of papers

<table>
<thead>
<tr>
<th>Country</th>
<th>Number of studies</th>
</tr>
</thead>
<tbody>
<tr>
<td>UK</td>
<td>18</td>
</tr>
<tr>
<td>Australia</td>
<td>6</td>
</tr>
<tr>
<td>Sweden</td>
<td>4</td>
</tr>
<tr>
<td>Ireland</td>
<td>3</td>
</tr>
<tr>
<td>USA</td>
<td>3</td>
</tr>
<tr>
<td>Canada, Germany, Italy, Japan,</td>
<td>1 each</td>
</tr>
<tr>
<td>the Netherlands, Norway</td>
<td></td>
</tr>
</tbody>
</table>
What we found

A series of seven analytical themes that represent patients’ and carers’ experiences of living with MND, across the course of its trajectory

The focus of the review was palliative care needs. Very few reports of palliative care in the papers. Where it was talked about it was within the last days of life.
• Delays of up to 60 months
• For some diagnosis came as a shock, for others it confirmed their expectations
• Negative responses, including bewilderment, denial, a sense of loss and sadness

“you don’t understand the news, you deny it, become very despairing”

(Bolmsjo 2001)

• Diagnosis was the time-point where poor communication from HCPs was most noted
• Individuals and carers stated a clear need for follow-up support immediately after the diagnosis, which was recalled as a very vulnerable time
Maintaining control

• People sought to maintain control of their lives, keeping a sense of normality and purpose.

• As the disease progressed, both patients and carers tried to find ways to counteract feelings of loss and uncertainty, and to find new meaning in their life.

“Determination to do what I can for as long as I can, desire for independence [and a] resilient temperament.” (patient)

(Hogden et al 2012)
Deterioration & decision making

• Deterioration and its physical, emotional and social sequelae had a significant effect on the lives of both patients and carers.

• Physical difficulties experienced during periods of significant deterioration included:
  • speech/communication
  • breathing
  • eating/swallowing
  • decline in mobility from increasing muscle weakness

During deterioration decision making became key, supporting (or not) the introduction of interventions to palliate symptoms.....but....
Interventions ......

...life-sustaining treatments were not always seen as interventions to ease potential suffering, but as a way of being kept alive:

Participant:  *It would be staying alive under what conditions. Like you know, if they were just keeping you alive ... that Hawkins fella, sure he has it since he was 22 and he is 75 now ... But he’s all curled up in a chair. Ah I would hate that ...*

Interviewer:  *So that wouldn’t be acceptable to you?*

Participant:  *No, even if I had his brain.*

Individuals reported valuing autonomy in decisions, wanting to be involved and respected in the decision-making process, and wanting to have the final say in what interventions they would try
End of life

- Palliative care was often associated with the **dying phase only**, with life-sustaining interventions in MND not being related to a palliative care approach.
- It was associated with treatments that would alleviate suffering at end of life.
- Assisted suicide and euthanasia were freely talked about by both patients and carers as potential options.
- Such considerations were connected with a wish to die before the ‘final stage’ of the illness; both to maintain a sense of control over its timing and the extent of the suffering that would be experienced.
There is a need for information to help people make decisions regarding end-of-life care. This was commonly perceived to be lacking:

‘I would like to know how it will be managed and what my choices are’

‘I really do not want to die in hospital unless it is absolutely necessary’

(Whitehead et al 2011)

Discussions concerning dying or death were not common between patient and carer

‘I think we were in denial for quite some while you know, we knew it was coming, but we didn’t plan anything about it’

(Ray et al 2014)
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If conversations about dying and death had occurred they were generally positive. If plans had been made caregivers were able to achieve some sense of comfort.

‘He told us everything he wanted; he always had done from the end of last year. He said you know I want this, I don’t want that’

(Ray et al 2014)
Family caregivers

- Serious concerns re **lack of knowledge** among health care professionals influences willingness to engage with such services and ‘hand over’ care

- Impacts on care at the end of life - inexperience or lack knowledge from palliative care services for the needs of people with MND

  ‘The last 3 weeks of my mother’s life... and they made it terrible absolutely terrible, well it was horrific...through their incompetence and through their not knowing, the lack of knowledge of motor neurone...’

    *(O’Brien & Preston 2015)*

- Family carers struggled to find answers about the dying process; dealing with a multitude of service providers added to the confusion.

  ‘I would have thought that it would be better for the whole MND process to have a single liaison person..’

    *(Bentley & O’Connor 2016)*
• Despite these efforts to plan for death, problems arose when plans were not communicated effectively eg when ‘not for resuscitation’ orders and the caregiver’s wishes were ignored:

‘I couldn’t [make them stop] ‘stop it, don’t give her artificial resuscitation’ but...... That’s the one disappointment; I would have loved her to go peacefully.’

• Living with a neurodegenerative illness raised expectations that the person would eventually die of respiratory failure and this would be anticipated. When people with MND died, it was often sudden and traumatic for their families.

‘I didn’t understand that her life was in danger. Therefore, even when she was making these dreadful noises.I did not realize that this was near the end.’

(Ray et al 2014, Bentley & O’Connor 2016)
Some recommendations

• Re-framing palliative care to the point of diagnosis and ensuring prompt access
• Encouraging health professionals to initiate advance care planning early in the disease
• Preparation for the next stage of the disease trajectory should take place during the phase before, particularly in relation to the introduction of interventions
• Single point of access or a ‘key worker’ to the multi-disciplinary team and focus for all care
• The lack of knowledge among health care professionals in acute and community services of the needs of people with MND needs addressing – against the challenge of the lack of exposure to caring for people with the condition
Thank you for coming along to listen