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The experience of informal caregivers of patients with Motor Neurone Disease: A thematic synthesis

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Abstract

Objectives: Research is required in order to illustrate and detail the experiences of informal caregivers of patients with motor neurone disease (pwMND) to further advance the research base and to inform the development of future support structures and services. Due to the heterogenous nature of caregiving for pwMND, one way in which this can be achieved is through a qualitative review. A qualitative thematic analysis of existing qualitative studies has not, to the best of the authors’ knowledge, been previously undertaken. Thus, this synthesis aims to identify caregivers’ experiences and to suggest factors that contribute to these experiences in order to fulfil the required research need.

Methods: A thematic synthesis of qualitative literature was conducted. AMED, Medline, SPORTDiscus, CINAHL, and PubMed were electronically searched from inception until September 2015. Studies were eligible if they included qualitative literature reporting on first-hand experience of informal caregivers of patients with MND, were published in English, and contained verbatim quotations. Critical appraisal was undertaken using a 13-item COREQ checklist.

Results: Ten studies met the inclusion criteria, with 148 (50 male) current or previous informal caregivers of pwMND identified. Critical appraisal demonstrated study design and reflexivity were under-reported. The synthesis derived three themes: 1) loss of control, 2) inability to choose, 3) isolation.

Significance of results: The synthesis highlighted factors which contribute to both positive and negative caregiving experiences. Through these experiences suggestions for service provisions such as improving communication with healthcare professionals and having a single point of contact have emerged. However, the outcome of such suggestions on the caregivers’ experience is beyond the scope of this synthesis and hence further research is required.

Key words Caregivers, motor neurone disease, qualitative research, experience.
Introduction

Motor neurone disease (MND) is an adult onset neurodegenerative disorder represented by four subtypes; the most common being amyotrophic lateral sclerosis (ALS) (Aoun et al., 2013; Nageshwaran, 2014). The terms ALS and MND are used inconsistently; the UK uses MND as the umbrella term whereas other countries use ALS (Mitchell and Borasio, 2007; Aoun et al., 2013). Therefore, they will be used synonymously in this study. Around 4,500 people in the UK are living with MND at any time and peak onset is between 50-70 years (MND Association, 2015). Although symptoms and rate of progression vary between the four subtypes (Herz et al., 2006; Nageshwaran, 2014), the most common symptoms include muscle wasting, weakness, fasciculations, and dysarthria (Mitchell and Borasio, 2007; Nageshwaran, 2014). Currently there is no cure, and 50% of patients do not live beyond three years from onset of symptoms (Mitchell and Borasio, 2007).

In the period between symptom onset and death, care of the patient often falls upon family members or friends (Mockford et al., 2006). The time spent caregiving is variable on the patients’ disease progression with Chio et al., (2006) reporting for patients with a mild disability (ALS-FRS score >30) a mean of 5 hours/day and for patients with severe impairments (ALS-FRS score ≤10) a mean of 15 hours/day. Caregivers’ quality of life (QoL) is significantly negatively impacted by undertaking the role (Peters et al., 2013) and many caregivers experience burden associated with their role, attributed to physical, emotional, social, financial, and psychological factors (Sorrell, 2014).
Although research focusing upon caregivers’ experience is expanding, it remains sparse (Mockford et al., 2006). Two relevant reviews have been published. Mockford et al., (2006) focused on published literature from 1994-2004; the main findings discussed caregivers experiencing ill health when unsupported and the positive impact of maintaining social support and activities on reducing the chances of ill health. Aoun et al., (2013) conducted a more recent review for the period 2000-2011 which indicated that, although research has highlighted factors such as diagnosis process and access to information and services which contribute to burden and negatively impact QoL, there is a need to develop literature focusing on improving caregivers’ experience. As further literature has been published since 2011, this study aims to update existing reviews.

Researchers have called for further consideration of caregivers’ experiences and in particular qualitative based reviews (Goldstein et al., 2006; Murphy et al., 2009; Atkins et al., 2010). Due to the expanding literature base and the risk of single qualitative studies being overlooked in practice if not synthesised (Malpass et al., 2009) it was considered important to utilise a qualitative review based approach to address these calls. Both Mockford et al., 2006, and Aoun et al., (2013) included quantitative and qualitative literature and hence, as this study aims to focus solely on a qualitative approach, it has the potential to offer alternative interpretation and results as the caring experience can be considered as heterogeneous and subject to personal or unique interpretations (Crellin et al., 2014). Thus, this study aims to provide a thematic synthesis of existing qualitative literature regarding experiences of informal caregivers of pwMND. The study aims to encompass the spectrum of experiences through not focusing on a specific stage of the disease progression.
Additionally, the study aims to highlight areas for service development to reduce caregiver burden in the future through discussing positive caregiver experiences and suggestions made in existing literature.
Methods

The methods section has been reported in accordance with the enhancing transparency in reporting the synthesis of qualitative research guidelines (ENTREQ) (Tong et al., 2012a).

Search strategy

Electronic databases were searched from inception until September 2015; Allied and Complementary Medicine Database (AMED), Medline, SPORTDiscus, Cumulative Index to Nursing and Allied Health Literature (CINAHL), and PubMed. Multiple databases were utilised to maximise search yield (Wu et al., 2012). The selected databases were chosen as they focus on medical, allied-health, and social science disciplines (Shaw et al., 2004; Hewitt, 2007) and all but PubMed have been used in published reviews and synthesis concerning MND (Aoun et al., 2013; Soundy and Condon, 2015).

Combinations of the following key terms were utilised: “care*” OR “caregiv*” AND “motor neuron* disease” OR “amyotrophic lateral sclerosis” OR “MND” AND “experience” AND “qualitative” OR “mixed method” AND “interview” OR “focus group”. Although highlighted as potential keywords, “family” and “perceptions” were not utilised as they retrieved large volumes of irrelevant studies during scoping search. Additional search strategies included citation chasing, reviewing reference lists, and contacting researchers (n=1).

Eligibility criteria
Included studies needed to fulfil the eligibility criteria which utilised the SPIDER (sample, phenomenon of interest, design, evaluation, research type) search tool (Cooke et al., 2012);

- **Sample**: Informal primary caregivers were defined as individuals who previously or currently provide the majority of support and assistance to a pwMND at any disease stage, without financial reward.

- **Phenomenon of interest**: Studies reporting informal caregivers’ first-hand experience with MND were included.

- **Design**: Any type of qualitative design was considered. Excluded designs: were quantitative research, mixed-method research not containing clear qualitative data, grey literature; unpublished or protocol studies, thesis, conference proceedings, and case studies.

- **Evaluation**: Any qualitative method of data collection; interview, focus group or mixed-method articles containing clear qualitative research were included.

- **Research type**: Studies including clear qualitative data from a qualitative or mixed-method study. Studies published in languages other than English were excluded.

**Critical Appraisal**

This review used an adapted 13-item version of Tong et al., (2007) Consolidated Criteria for Reporting Qualitative Studies (COREQ), which has three domains:

(i) research team and reflexivity which increases transparency of personal characteristics of the research team as well as their relationship with participants which could influence bias within data collection and interpretation of results,
(ii) study design which appraises participant retention and the data collection process,

(iii) data analysis and reporting which appraises the process of coding and derivation of themes.

The adapted version of COREQ was devised by Soundy et al., (2016) following critique of the larger 32 item checklist (Tong et al., 2007) where items that were not sensitive were removed.

Studies were independently appraised scoring one for reporting a checklist item and zero if unclear or unreported (Tong et al., 2007). A total score of 13 was possible and, based on Soundy et al., (2016) previous research, a score of 4 was identified as a threshold where further consideration of the study was made. Any studies scoring less than 4 were discussed to identify if there was a methodological weakness which compromised the results and warranted exclusion of the article. No articles were excluded following this process.

Synthesis

An adapted four-stage approach was taken for this synthesis (Thomas and Harden, 2008; Aria et al., 2007; Rodgers et al., 2009): Stage one involved textual description, extraction and tabulation of results following which a summary of individual extracted quotations via line-by-line coding was used to facilitate comparison. Stage two involved idea webbing to visualise and connect concepts between studies (Rodgers et al., 2009). Stage three involved translation; interpreting and converting similar concepts between studies (Aria et al., 2007). Stage four involved synthesis of
translations to combine the analysed themes and form a new interpretation of the studies involved. See supplementary file for the audit trail.
Results

Systematic search

The full search process is demonstrated via Prisma (Moher et al., 2009) flow diagram (Figure 1). Ten articles (Brown, 2003; Akiyama et al., 2006; Herz et al., 2006; Ray and Street, 2007; Aoun et al., 2012; O'Brien et al., 2012; Whitehead et al., 2012; Hyunjin and Schepp, 2013; Oyebode et al., 2013; Weisser et al., 2015) met the inclusion criteria and were analysed within the thematic synthesis. Characteristics of included participants are presented in table 1.
Critical Appraisal

The results of the 13-item COREQ ranged from four (Akiyama et al., 2006; Whitehead et al., 2012) to eight (Ray and Street, 2007) with a mean total score of six. Hence, no studies were excluded nor discussed for exclusion. For full COREQ results see the supplementary file.

Although Akiyama et al., (2006) and Whitehead et al., (2012) scored the minimum score of four, they were the only included studies to score 0/5 within domain one of the COREQ. This impacts credibility of these studies’ results as there is a lack of transparency of the researchers’ credentials and relationship with participants thus there is an inability to distinguish the degree to which personal bias may be influencing the interpretation of the findings (Tong et al., 2007). Both Akiyama et al., (2006) and Whitehead et al., (2012), failed to report on the number of, or reason for, participant dropouts resulting in a possibility of selective reporting (Tong et al., 2007; Cote and Turgeon 2009). Furthermore, although both studies described a coding process, Akiyama et al., (2006) failed to report on the derivation of themes. Therefore, there is a lack of transparency regarding the study design and the researchers’ rationale for their interpretation and analysis of the data (Fossey et al., 2002). Hence, due to lack of transparency limiting the ability to interpret results, novel data from Akiyama et al., (2006) and Whitehead et al., (2012) was interpreted with caution and with consideration of findings from the other included studies.
Thematic synthesis

The thematic synthesis highlighted that, although the caregivers experience is non-homologous, shared key themes exist. These themes are explored below and are grouped as follows: 1) factors contributing to the experience of loss of control, 2) factors contributing to the sense of choice, 3) factors contributing to the experience of isolation. Studies supporting each theme are demonstrated in table 2.

INSERT TABLE 2 HERE

Loss of control

Progressive nature of MND

The progression of the cared for patients’ MND was considered in relation to the idea of loss of control within the caregiving role. Several studies highlighted that caregivers were unsure of how their role would evolve alongside the progression of the patients’ MND (Aoun et al., 2012; Brown, 2003; O’Brien et al., 2012; Ray and Street, 2007; Whitehead et al., 2012). Caregivers were aware symptoms would progress but were unable to highlight how symptoms would change and how this would impact their caregiving role (O’Brien et al., 2012). Furthermore, caregivers were uncertain of the rate of the patients’ MND progression (Aoun et al., 2012; O’Brien et al., 2012; Oyebode et al., 2013; Ray and Street, 2007; Whitehead et al., 2010). These factors contributed to caregivers’ inability to plan and prepare for the future within their caregiving role (Aoun et al., 2012; Brown, 2003; O’Brien et al., 2010; Ray and Street, 2007; Whitehead et al., 2012). One caregiver epitomises this through the quote; “This is so undetermined you just don’t know it from one moment
to the other. Whether it’s going to stop in its tracks or plummet downhill you just don’t know and that’s very difficult” (Ray and Street, 2007). Furthermore, this inability to prepare for continual progression resulted in a sense of progressive overwhelming responsibility and a struggle to stay in control (Herz et al., 2007). For some, uncertainty resulted in fear and anxiety (Aoun et al., 2012; O’Brien et al., 2012), conversely other caregivers could embrace uncertainty and ‘live in the moment’ as a coping mechanism (Weisser et al., 2015; Oyebode et al., 2013).

One suggestion for uncertainty is that the breadth of symptoms and life expectancy of pwMND are not fully understood and therefore cannot be accurately predicted (Ray and Street, 2007). However, other caregivers’ experience suggests a breakdown in communication between healthcare professionals and caregivers’ knowledge exists with relevant information not being shared with caregivers (O’Brien et al., 2012; Ray and Street, 2007). A suggested reason for lack of clear communication is professionals having insufficient time for discussion with both the pwMND and their caregivers (Herz et al., 2006; O’Brien et al., 2012). Furthermore, insufficient time was associated with professionals’ lack of up-to-date MND specific knowledge (Herz et al., 2006) which subsequently limited caregivers’ knowledge and expectations of future caregiving intensity.

Loss of control over daily activities

The idea that caregivers had a sense of loss of control over their own lifestyle emerged in several studies (Akiyama et al., 2006; Aoun et al., 2012; Brown, 2003; Herz et al., 2006; Hyunjin and Schepp, 2013; O’Brien et al., 2012; Oyebode et al., 2013; Ray and Street, 2007). A key idea for the mechanism for this developed from
the sense of role captivity (Akiyama et al., 2006; Aoun et al., 2012; Herz et al., 2006; Hyunjin and Schepp, 2013; O’Brien et al., 2012; Oyebode et al., 2013; Ray and Street, 2007). “You can’t go out when you want to go out, you can’t go where you wish to go or you used to be able to go” (Oyebode et al., 2013). Caregivers highlighted the sense of being solely responsible for the pwMND daily care and needs without the support of others to share the role and allow previous activities and social opportunities to be continued (Akiyama et al., 2006; Aoun et al., 2012; Herz et al., 2006; Hyunjin and Schepp, 2013; O’Brien et al., 2012; Oyebode et al., 2013; Ray and Street, 2007). Hence, the sense of captivity and subsequent loss of control could differ between caregivers depending on the level of support they receive (Oyebode et al., 2013; Weisser et al., 2015). However, formal support may have a peak intensity which prevents the feeling of loss of control with intensity above this threshold contributing to the caregivers’ experience of loss of control due to loss of privacy and increased dependence on others (Weisser et al., 2015). However, this idea did not consistently emerge throughout the included studies so it is plausible the contribution of support to loss of control may be caregiver specific.

Alternatively, one study explored the idea that the loss of ability to choose daily activities arose from the inability to complete activities previously shared with the pwMND due to their functional decline (Ray and Street, 2007). This demonstrates how others influence the sense of control over decision making for one’s own life.

For caregivers, one consequence of the experience of loss of control over activities was the sense of burden (Hyunjin and Schepp, 2013; O’Brien et al., 2012). Another was the sense of living for others dehumanising their own existence (Akiyama et al.,
For some, this resulted in a change in the individuals’ self-perception and a sense of being a different version of themselves (Herz et al., 2006; Ray and Street, 2007) “You’re not yourself, you’re certainly not yourself…It’s, you feel more of a carer than anything” (Ray and Street, 2007).

Choice

Choice regarding uptake of the caregiver role

Caregivers experienced a sense of obligation resulting in an inability to choose whether to become a caregiver (Herz et al., 2006; O’Brien et al., 2012; Oyebode et al., 2013; Whitehead et al., 2012). One emerging idea was the sense that the wishes of the pwMND to be cared for by informal caregivers came before the caregivers’ needs or desire (Akiyama et al., 2006; O’Brien et al., 2012; Whitehead et al., 2012) due to the sense of needing to alleviate suffering within a limited lifespan (Whitehead et al., 2012). Furthermore, to protect the pwMND from further suffering, there is a sense of caregivers concealing difficulties and struggles from the pwMND and hence their own needs going unnoticed and neglected (Oyebode et al., 2013). However, for others the obligation was engrained in the expectation to care for loved ones (Herz et al., 2006) and demonstrate commitment to marriage (Oyebode et al., 2013).

Access to alternatives

Caregivers experienced a lack of alternative options to caring fulltime for the pwMND (Aoun et al., 2012; Brown, 2003; Herz et al., 2006; O’Brien et al., 2012; Oyebode et al., 2013; Weisser et al., 2015; Whitehead et al., 2012). Although unable to access
support, some caregivers could identify the need for specialist support (Herz et al., 2006) whereas others suggested insufficient time and captivity of the role did not allow for the consideration of the need for support (Aoun et al., 2012).

The synthesis highlighted the financial implications of accessing support for example equipment or paid for caregivers being a luxury not all could access (Herz et al., 2006; Oyebode et al., 2013; Whitehead et al., 2012). Furthermore, a lack of haste in processing applications for financial support prevented eligible pwMND receiving funding due to passing away prior to the reviewers coming to a decision (Whitehead et al., 2012). Although this may relate to the difficulty in predicting progression of MND, it highlights the need for more rapid consideration of applications.

Also highlighted was a lack of specialist caregiving services for the needs of pwMND (Aoun et al., 2012; Herz et al., 2006; Oyebode et al., 2013; Whitehead et al., 2012). This resulted in caregivers not being confident that the needs of the pwMND would be met and subsequent unwillingness to release the caregiving role to unspecialised support services such as paid for caregivers (Herz et al., 2006; Whitehead et al., 2012). However, this is not conclusive as other caregivers were able to access specialist support from services such as the MND association and hospice care (Herz et al., 2006). Hence, care quality may be inconsistent between providers.

Isolation

Communication with healthcare professionals
An idea which emerged from the synthesis was that communication between professionals and caregivers contributed to the caregivers’ sense of being unsupported and isolated (Aoun et al., 2012; O’Brien et al., 2012). One factor highlighted was the idea of high volumes of involved healthcare professions without clarity on each of their roles or a key member to contact (O’Brien et al., 2012). Hence it could be suggested that potential sources of support are not utilised due to poor clarity of communication which leads to caregivers being overwhelmed. Furthermore, for some, discussion with professionals lacked depth and personalisation with failure to attend to emotional aspects of caregiving for a pwMND (Brown, 2003) resulting in caregivers feeling unsupported (Brown, 2003; Herz et al., 2006). “I think it’s all talked about mainly in a business-like manner rather than in an emotional way” (Brown, 2003). This suggests a lack of services which attend to the caregivers’ needs and a requirement for professionals to shift their focus when communicating with caregivers. However, others used counselling services which allowed cathartic release (Herz et al., 2006; O’Brien et al., 2012) and for some caregivers, depth and effectiveness of communication was accessible however this appears to be profession dependent and may highlight a greater need for access to counselling services to meet caregivers’ needs.

Communication between family and friends

Changes in communication within pre-existing relationships with family and friends emerged in several studies (Herz et al., 2006; Hyunjin and Schepp, 2013; O’Brien et al., 2012; Oyebode et al., 2013; Weisser et al., 2015). A sense of inability to share was noted to avoid avoiding burdening others (Herz et al., 2006; Hyunjin and Schepp, 2013; O’Brien et al., 2012) which resulted in concealing emotions from
others and a sense of isolation; “I’d cry. I used to cry in the shower. The shower was my friend. I loved my shower. Because they don’t notice your tears in the shower” (Herz et al., 2006). Additionally, caregivers struggled to communicate with non-involved others due to their lack of understanding of the caregivers’ situation and burden (Hyunjin and Schepp, 2013) suggesting a lack of empathy and an alienating experience. However, other caregivers appreciated a sense of normality when communicating with friends (Akiyama et al., 2006; Ray and Street, 2007) indicating that communication within previous relationships was not impacted by the caregiving role. However, it is plausible that this sense of ‘normality’ is a denial coping mechanism where the caregiver avoids sharing their current experience or concerns. Thus it is arguable that communication is impacted as there may be a shift in topics which can be openly discussed when communicating with others.

Caregivers also experienced a decreased frequency of verbal communication with the pwMND, another pre-existing relationship (Hyunjin and Schepp, 2013; Weisser et al., 2015). One contributing factor which emerged was that MND symptoms were attributed to decreased quality of communication due to a need to simplify language (Oyebode et al., 2013). For some the frustration of the caregiving role led to tension and avoiding communication (Hyunjin and Schepp, 2013; O’ Brien et al., 2012). Alternatively, others noted an increase in non-verbal communication through activities involved with the caregiving role (Weisser et al., 2015), demonstrating a potential shift away from verbal language and hence altering the method of communication. Additionally, some caregivers found the ability to share more readily with the pwMND, increasing verbal communication (Weisser et al., 2015), however,
the motive is unclear, and could be due to differing factors such as the terminal prognosis rather than directly due to the caregiving role.

Changes to relationship dynamics

Caregivers experienced a change in the dynamics of the relationship with the pwMND (Aoun et al., 2012; Herz et al., 2006; Hyunjin and Schepp, 2013; Oyebode et al., 2013; Ray and Street, 2007; Weisser et al., 2015). Former spousal relationship identities became a nurse-patient or parent-child relationship identity due to the pwMND dependence and lack of reciprocity within the relationship (Aoun et al., 2012; Hyunjin and Schepp, 2013; Ray and Street, 2007). Caregivers explored the sense of disconnection from their partner and sense of being alone in their marriage (Hyunjin and Schepp, 2013; Oyebode et al., 2013). For some, this escalated to questioning their commitment to the marriage due to the pwMND dependence and intensity of the caregiving role (Herz et al., 2006; Hyunjin and Schepp, 2013).

Loss of intimacy

A factor contributing to the changing dynamics of spousal relationships and subsequent experience of isolation was the loss of physical and emotional intimacy (Aoun et al., 2012; Hyunjin and Schepp, 2013; Oyebode et al., 2013; Ray and Street, 2007). Aoun et al., (2012) and Oyebode et al., (2013) discussed physical barriers due to MND symptoms which limited intimacy and resulted in the experience of isolation due to absence of physical contact and sense of connection with their spouse. Additionally, for some caregivers the requirement to attend to the pwMND personal care limited the desire for physical intimacy (Hyunjin and Schepp, 2013; Ray and Street, 2007). Caregivers’ decreased desire for physical intimacy resulted in
increased tension between spousal couples subsequently leading to a sense of isolation (Hyunjin and Schepp, 2013). However, as well as the direct relationship between attending to the pwMND care needs and loss of desire for intimacy, caregivers highlighted how attending to the pwMND personal care altered the relationship identity due to the loss of reciprocity in the relationship and it was the change in relationship identity which limited intimacy (Aoun et al., 2012; Ray and Street, 2007). This demonstrates that relationship identity can be the cause of loss of intimacy and a result of loss of intimacy.

**Opportunities for interaction**

Decreased opportunity for social interaction with others further contributes to caregivers’ experience of isolation (Akiyama et al., 2006; Aoun et al., 2012; Brown, 2003; Herz et al., 2006; O’Brien et al., 2012; Ray and Street, 2007; Weisser et al., 2015). Decreased opportunity for interaction related to role captivity of caregiving and subsequent insufficient time. One caregiver reported “I used to go weeks and sometimes months before I saw anybody.” (Brown, 2003) highlighting physical isolation from others limiting interaction. As well as limited time, fatigue associated with the caregiving role was associated with decreased interaction (Ray and Street, 2007). However, another caregiver highlighted the sense of isolation and loneliness despite maintaining work alongside the caregiving role (O’Brien et al., 2012) suggesting physical isolation is not the only factor contributing to limitations in interactions. For other caregivers, the alteration in relationship dynamics and the inability to complete social activities previously completed as a couple limited social opportunities leading to a sense of isolation both during the caregiving and bereavement phase (Ray and Street, 2007). Conversely Aoun et al., (2012) suggest
that isolation was only experienced in the bereavement phase as caregivers were supported in the caregiving phase “You go from having a whole army of people then it’s just you”. However, whether the caregiver received formal support in the form of paid caregivers or informal support from friends and family was not discussed which may impact the sense of isolation and loss of interaction. Hence, opportunity for interaction maybe caregiver specific and level of support may impact the caregivers’ experience of isolation.
Discussion

The aim of this thematic synthesis was to explore existing literature to increase depth of understanding of informal caregivers’ experience. Through synthesis of the existing qualitative literature three themes were identified which demonstrated how loss of control, absence of choice, and isolation are experienced by caregivers and identified factors which contribute to these experiences.

A key idea highlighted across themes was the sense of being solely responsible for the pwMND without help or support from others. This responsibility can stem from the sense of obligation to fulfil the caregiver role either to satisfy the patient’s wishes, demonstrate love and commitment, or due to the absence of alternatives or support. Access to support differed between participants included in the thematic synthesis with those individuals who were financially able in an advantageous position. However, as the synthesis included studies from any country, financial requirements may differ between countries and the support systems available (Sakellariou et al., 2013). Also highlighted was the lack of communication about available support between healthcare professionals and caregivers. Reasons for lack of communication included insufficient time, inadequate knowledge of MND, involved healthcare professionals lack of awareness of caregivers’ emotional difficulties, or having a multitude of involved healthcare professionals with lack of clarity over their roles. Highlighted was the need for healthcare professionals to have knowledge of, and ability to communicate, the likely future for the pwMND and how this subsequently impacts the caregiver. One idea which emerged was the need for a key worker to address caregivers’ questions, provide practical and emotional support and highlight or signpost to relevant services to increase access to support.
However, this idea is not used in any of the included studies and thus the outcome cannot be determined within the scope of this thematic synthesis. However, this idea has been reflected in the UK through the NICE guidelines’ recommendation to provide of a single point of contact within the MND-specific multidisciplinary team (NICE, 2016). This therefore supports the ideas emerging in the synthesis as they are recommended for implementation in clinical practice.

Caregivers’ sense of being solely responsible for the pwMND resulted in insufficient time to maintain social interactions and previously completed activities due to being captive within the caregiver role. This was associated with a sense of burden, isolation, and living for others. This is supported by previous research regarding informal family caregivers for non-specific chronic conditions which highlights how insufficient time for activities which previously contributed to personal identity results in a sense of living for others and loss of self (Eifert et al., 2015). Highlighted in this thematic synthesis was the need for support or respite to allow maintenance of activities however frequency, type, or feasibility of support was not discussed and hence it is beyond the scope of the synthesis to make conclusive recommendations. Despite this, the synthesis highlights the need for formal caregivers to have sufficient knowledge and skills to care for the pwMND to allow informal caregivers to readily accept support, as insufficient ability is a key identified reason for underutilisation of support. This is supported by Aoun et al., (2013) review which suggested the need to train involved service providers to improve their understanding of working with pwMND however this was not discussed in the context of poor uptake of support services. Conversely another barrier to social interaction is the sense of avoiding burdening others, and lack of understanding or empathy from non-involved others limiting communication and interaction within previous relationships. Hence time for
social interactions through formal support alone would not address these issues. Further research is therefore indicated to understand the complexities in caregivers’ social interaction to make conclusive recommendations to address the issues. Caregivers experienced a sense of isolation with one contributing factor being a shift in spousal relationship dynamics and identity due to the pwMND dependence. Previous research amongst stroke patients and their caregivers suggested that dependence can result in the experience of inequality in the relationship due to increased contribution compared to benefits received and is associated with caregivers’ perception of burden (McPherson et al., 2011). Furthermore, Ybema et al., (2002) suggested receiving support from others maintained relationship equity and reduced caregivers’ sense of burden. Hence, formal support could reduce the pwMNDs’ dependence on the informal caregiver maintaining relationship equity and subsequently preserving relationship identity and dynamics. However, analysis of the outcome of this suggestion is beyond the scope of this review.

Limitations
Study limitations exist as, although systematic and thorough, it is possible that relevant studies may have been neglected due to variable indexing of qualitative research. The synthesis included caregivers of patients with any type of, and at any stage of MND, thus the findings were not specific to any stage or symptoms of MND. Subsequently this could be explored in future research. Female caregivers were over-represented in this synthesis, although, as more men are affected by MND, this may not be problematic (Goldstein et al., 2006). Demographical factors were not considered for example disposable income or culture; this may affect the ability of
caregivers to relate to the findings. Furthermore, the synthesis included studies from any country and thus consideration of different healthcare systems was not made.

Conclusion

This synthesis explores the experiences and challenges faced by non-formal caregivers. Although suggestions for service development have emerged from existing literature, there is a lack of information regarding the implementation and outcome of these suggestions upon the caregivers’ experience.

Conflict of Interest

The authors declare there are no conflicts of interest.

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References


Figure 1. PRISMA flow diagram.
<table>
<thead>
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<th>Participant Characteristics</th>
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<td><strong>Country of residence</strong></td>
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<tr>
<td>Japan</td>
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<td>South Korea</td>
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<td><strong>Caregiver type</strong></td>
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<tr>
<td><strong>Caregiver status</strong></td>
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<tr>
<td>Current</td>
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<td><strong>Gender</strong></td>
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</tr>
<tr>
<td>Male</td>
<td>50</td>
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Table 2. Overview of themes and study support at the subtheme unit

<table>
<thead>
<tr>
<th>Theme</th>
<th>Subtheme</th>
<th>Supporting studies</th>
</tr>
</thead>
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<tr>
<td>Loss of control</td>
<td>Due to the progressive nature of MND</td>
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<td>Over daily activities</td>
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<td>Choice</td>
<td>Uptake of caregiver role</td>
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<td>Access to alternatives</td>
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<td>Isolation</td>
<td>Communication with healthcare professionals</td>
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<td>Communication between family and friends</td>
<td>1, 2, 4, 5, 6, 7, 8, 9</td>
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<td>Changes to relationship dynamic</td>
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<td>Loss of intimacy</td>
<td>2, 5, 7, 8</td>
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<td>Opportunity for interaction</td>
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