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Nicola Glennie, Fiona M. Harris & Emma F. France

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Perceptions and experiences of control among people living with motor neurone disease: a systematic review and thematic synthesis

Nicola Glennie, Fiona M. Harris, and Emma F. France

Motor Neurone Disease (MND) is a neurodegenerative disorder of unknown aetiology where the motor neurones no longer function effectively leading to muscle wasting and the associated decline in physical function [1]. The disease pattern can vary individually but the progressive nature of the disease means that, although the timing is unpredictable [2], the decline in the ability to complete activities of daily living and maintain a social life is inevitable. The most common physical problem is muscle weakness, which leads to respiratory failure, which is usually the cause of death [1,3]. Although rare, MND has a large socioeconomic cost [4] and the effects on daily life are profound for both people living with MND (plwMND) and their families and carers [5,6].

There is as yet no cure for MND, and thus medical care involves symptom treatment or the provision of life-sustaining treatments such as Non-invasive Ventilation (NIV) to aid breathing and placement of a feeding tube when nutritional intake is impaired. Although thought to have the potential to improve quality of life or prolong survival [7], neither of these life-sustaining treatments offer any respite in the progression of the disease [8] and 50% of patients die within 3 years of symptom onset [9].

There is no definitive diagnostic indicator for MND and due to the diversity of initial symptoms plwMND may have gone through a period of many months of referrals to various specialists before being "stunned" by the terminal diagnosis [10,p.82]. After this initial shock "stunned" by the terminal diagnosis [10,p.82]. After this initial shock, plwMND must learn how to live with the unpredictable progression of this disease. Research shows a great diversity in this experience [5,11] but one factor which seems to be consistent is a perception of a loss of control within their lives. Within health research many concepts and theoretical models of control have been researched: for a comprehensive discussion of control in relation to neurodegenerative...
diseases see Eccles and Simpson [12]. This review will focus on the
concept of perceived control which has been defined as the belief
that a particular situation is under control or can be controlled [12].
Research on other neurodegenerative illnesses has shown that per-
ceived control is an important factor in the quality of life [13,14].
Increasing a person’s perception of control in a given situation is
thought to lead to an increased sense of well-being [15].

The onset of a life-limiting disease is known to be a source of
stress that can lead to feelings of loss of control, helplessness and
anxiety [16]; more specifically feelings of a lack of control have
been highlighted as a major cause of anxiety and stress for
plwMND and their carers [17]. The importance of attempting to
maintain a sense of control in the lives of plwMND has been
found in a previous qualitative thematic synthesis [18] which
investigated in general terms people’s experiences of living with
MND. A previous quantitative review [12] highlighted a lack of
research measuring perceived control in MND and also the lack of
consistency in methods of assessing perceived control but there is
an increasing body of qualitative research exploring people’s
experiences of MND [18]. However, there have been no qualitative
evidence syntheses focusing specifically on how plwMND perceive
and experience control; such a synthesis could lead to a richer
interpretation and deeper understanding of the phenomena.

The aim of this systematic review and synthesis of qualitative
studies is to explore plwMND’s perception and experiences of
control within their daily lives, the potential reasons for a per-
ceived lack of control and how plwMND adapt to this. This is
important because research suggests that a person’s quality of life
and ability to cope with MND may be related to their perception
of how much control they have within their life [19,20]. An in-
depth exploration of perceptions and experiences of control or
lack of control during life with MND may suggest ways to
improve quality of life and inform health and support services
for plwMND.

This review explores the following questions:

What factors contribute towards perceptions of control
within plwMND?

Table 1. Inclusion and exclusion criteria.

| Inclusion criteria | Exclusion criteria |
|--------------------|--------------------|
| **Sample**         | Studies investigating adults with MND of any type. |
| **Phenomenon of interest** | Articles exploring plwMND’s perceptions or experiences of control or any domains related to control in their day to day lives. |
| Control includes any mention of control or autonomy or agency or any control constructs such as self-efficacy or mastery. |
| **Design**         | Primary studies of any type of qualitative method or a systematic review of qualitative methodologies. |
| Qualitative data or mixed methods if qualitative data clearly separated. |
| Any type of data collection e.g., interviews, focus groups however data must come directly from the plwMND. |
| Systematic reviews of qualitative or mixed methods studies. |
| **Research type**  | Peer-reviewed journal articles. |
| Articles which are not about MND. |
| Articles which include MND but the data for those with MND cannot be separated from other illnesses in the study. |
| Articles that do not focus on daily living e.g., those that focus solely on diagnostic process or end of life care. |
| Articles which make no mention of control, or any aspects related to control in their report. |
| Articles that focus on a particular institution or intervention or development of a scale. |
| Quantitative research or mixed methods if qualitative data cannot be isolated. |
| Studies where data came from carers or health professionals. |
| Books, theses, conference proceedings, grey literature, case studies, editorials, letters and other non-peer-reviewed articles |
| Any primary research articles already included in a relevant systematic review. |
| Articles not in English. |
| Articles published before 1999. |

How do people living with MND attempt to maintain control
in their daily lives?

**Methods**

This qualitative systematic review follows the methods for
themtic synthesis [21]. The inductive approach of thematic synthesis
was chosen to enable the data to be explored without *a priori*
algorithmic frameworks, generating initial themes from the data
and developing these into new interpretations or analytical
themes. In reporting this research the ENTREQ Enhancing trans-
parency in reporting the synthesis of qualitative research: report-
guidance [22] have been used and can be seen in
Supplementary File 2.

**Inclusion and exclusion criteria**

The inclusion and exclusion criteria were formulated by NG with
support from FH (a senior qualitative researcher) and are docu-
mented in Table 1. Peer-reviewed journal articles, both primary stud-
ies and qualitative evidence syntheses, containing qualitative data
which focused upon the daily lives of plwMND of any type were
included. As there may be a discrepancy between plwMND’s views
of their life and the views of their carers or health professionals
[23,24], only articles where the data came directly from plwMND
were included. The phenomenon of interest was plwMND’s perceptions
or experiences of control. The term control can be subjective
therefore other terminologies such as autonomy, agency, adaptation
and any domains potentially related to control were included;
articles that made no mention of control, or any aspects related to
control in the full text were excluded. Quantitative research was
excluded but articles reporting mixed methods studies were
included if the qualitative data could be separated and extracted.
Articles pre-1999 were excluded as were those not in English.

**Systematic search and screening process**

The SPIDER tool [25] for qualitative evidence synthesis offers a sys-
tematic strategy for searching for qualitative (and mixed-methods)
research studies and was used to help identify the key search concepts: (1) Sample – Adults with MND of any type (2) Phenomenon of interest – plwMND’s perceptions or experiences of control or any domains related to control in their day to day lives (3) Design – qualitative research. The search terms were mapped for each key search concept and the strategy tailored for each database; an example search can be found in the Supplementary Files. Six relevant databases CINAHL, Medline, PsycINFO, ASSIA, Embase, and AMED were searched up to January 2022. Full search histories are available on request from the corresponding author.

In total, the searches yielded 6755 citations. The results and screening process are detailed fully in the PRISMA flow chart [26] in Figure 1. Retrieved references were screened by title and abstract then by full text by the lead author, 20% of titles and abstracts were independently screened by the third author and another PhD student. Any disagreements in screening were discussed between screeners and an agreement was sought.

One of the articles found was a qualitative thematic synthesis of experiences of living with MND [18]. As the review article was a thematic synthesis with data extraction and analysis processes very similar to those intended for this paper and the review article met the inclusion criteria it was decided to include the review article rather than all the individual primary studies included in this review. In a qualitative analysis reanalysing these same primary studies may not yield any more significant results than those already found in the existing review.

Quality appraisal
All primary articles included in the final selection of this review were subject to quality assessment using the Joanna Briggs Institute Critical Appraisal Checklist for Qualitative Research [27]. Of these articles five were unclear if there was congruity between the stated philosophical perspective and the research methodology, ten had no statement locating the researcher culturally or theoretically and ten did not address the influence of the researcher on the research and vice-versa. No primary articles were excluded during the quality appraisal process as all 19 articles adequately portrayed participants’ voices.

Characteristics of included studies
Characteristics of included studies are shown in Table 2. The 20 included articles were 19 primary qualitative research articles.
| Study                                    | Location      | Study aims                                                                 | Sample details | MND Type                | Point in MND journey | Methodology and data collection |
|-----------------------------------------|---------------|----------------------------------------------------------------------------|----------------|-------------------------|----------------------|--------------------------------|
| Ando et al. [28]                        | UK            | To explore patients' perceptions and use of NIV over time                   | 5 People (m = 4) | 3 bulbar onset          | Illness duration 23–237 months | Phenomenology               |
| Ando et al. [29]                        | UK            | To explore why some plwMND refuse or withdraw from NIV                     | 9 People (m = 7) | 3 bulbar onset          | Unable to ascertain   | Phenomenology               |
| Ando et al. [30]                        | UK            | To understand factors important to QOL and explore how plwMND maintain and/or improve QOL | 26 People (m = 14) | 13 limb onset          | Illness duration 1–110 months | Phenomenology               |
| Cave and Bloch [31]                     | England       | To investigate what plwMND consider when deciding whether or not to bank their voice | 21 people      | Not specified           | Time since diagnosis 1–12 months | Semi structured interviews |
| Cipolletta et al. [32]                  | Italy         | To identify illness trajectories in pwMND                                  | 18 people (m = 8) | 12 ALS spinal form | Time since diagnosis range a few days to 4 years | Grounded Theory for qualitative section |
| Foley et al. [33]                       | Ireland       | To investigate how and why plwMND engage with healthcare services          | 34 people (m = 17) | Not specified           | Duration since onset of symptoms 4 months — 13 years | Mixed methods |
| Foley et al. [34]                       | Ireland       | To identify key psychosocial processes that underpin how plwMND engage with healthcare services | 34 people (m = 17) | 26 spinal onset | Disease duration 4–169 months | Grounded Theory |
| Greenaway et al. [35]                   | UK            | To investigate the decision-making processes involved in accepting or declining NIV | 21 people (m = 13) | 2 respiratory onset | Duration of disease 6–60 months | Hermeneutic phenomenology |
| Gysels and Higginson [36]               | UK            | To explore and compare the lived experience of breathlessness in patients with COPD, heart failure, cancer and MND | 48 people. plwMND = 10 (m = 9) | Not specified | Not specified | Method not specified used an inductive thematic analysis approach. |
| Harris [37]                             | UK            | Aims to answer the question 'What does it mean to be a person living through the illness trajectory of MND' | 4 people (m = 3) | 3 ALS | Time since diagnosis 14 months — 10 years | Hermeneutic phenomenology |
| Harris et al. [38]                      | UK            | To explore the meaning of living with uncertainty in pwMND                 | 4 people gender not specified | Not specified | Had been receiving treatment for MND for 3–6 months | Hermeneutic phenomenology |
| Hocking et al. [39]                     | New Zealand   | To shed light on the experience of living with MND                         | 7 people (m = 3) | Not specified           | Time since diagnosis 8 months 4–4 years | Phenomenology |
| Hodgden et al. [40]                     | Australia     | To identify factors influencing decision making in MND                     | 14 people (m = 7) | Not specified           | Time since diagnosis 2–93 months | Hermeneutic phenomenology |
| Mistry and Simpson [41]                 | UK            | Exploring the transitional process from diagnosis to living with MND       | 7 people (m = 3) | Not specified           | All diagnosed within previous 6 months | Hermeneutic phenomenology |
| Ozanne et al. [42]                      | Sweden        | To investigate how plwMND create meaning despite their diagnosis           | 14 people (m = 7) | Not specified           | Duration of disease 2–13 years | Semi structured interviews |
| Pavely et al. [43]                      | UK and Australia | Investigating technology and response to bodily limitation in MND          | 42 people (m = 31) | Not specified           | Not specified | Phenomenology 10 face to face interviews others conducted |

(continued)
Table 2. Continued.

| Study                  | Location          | Study aims                                                                 | Sample details       | MND Type       | Point in MND journey | Methodology and data collection                  |
|------------------------|-------------------|----------------------------------------------------------------------------|----------------------|-----------------|----------------------|-------------------------------------------------|
| Pinto et al. [44]      | UK                | To understand the emotional impact of living with MND and explore what impacts experiences of emotional distress and wellbeing | 35 people 25 plwMND (n = 15) Time since diagnosis 2 months | ALS limb = 18   | Time since diagnosis 2 months | Semi structured interviews                     |
| Sakelleriou et al. [18]| Includes studies from UK, Sweden, USA, New Zealand, Ireland, Australia Canada | To answer the question “What is known about people’s experiences of living with MND?” | 257 people gender not specified Time since diagnosis 1 month – 3 years | Not specified | Followed CRD systematic review procedures for search, quality appraisal, data extraction, analysis and synthesis |
| Weeks et al. [45]      | UK                | To explore needs and preferences of plwMND about psychological interventions and how these may be adapted for this population | 37 people 15 plwMND (n = 9) Age range 56–75 years | ALS = 10        | Not specified | Semi structured topic guides                   |
| Yuan et al. [46]       | China             | To gain understanding of the illness experience of als and meaning attached to these | 20 people(n = 13) Age range 28–72 years | ALS =20         | Time since diagnosis 6–60 months | Semi structured interviews                     |
Sub theme - Control beliefs about health

The studies suggested that prior to diagnosis most participants believed they were in control of their lives and their health. Several studies demonstrated that participants were working or caring for family, had active hobbies and took a fully active role in their communities [29,32,34,38,41].

The first symptom mentioned by participants was often a small loss of physical control over the body [36,39]. Harris [38,p.2065] reports “Charles was out walking his dog when he first noticed his foot drop.” Being unable to explain these symptoms led to the first doubts about the belief that they were in control of their health, Hocking [39,p.25] described this as a “wobble” in participants’ lives.

Participants sought the help of the medical profession [39,41] in the belief that they would be able to take a quick diagnosis and suggest treatment and thus they would regain control over their health. One participant (Bert) said:

So I’m thinking to myself, it could be this, that or the other, it’ll be something… I even thought it might be a tumour, something in my head that needs a quick zap, go in with a drill and pull it and the job’s done [41,p.944].

However, the lack of diagnostic markers in MND makes a quick diagnosis unlikely and most participants had a long and unsatisfactory process of diagnosis [32,34,36,37,39]. The diagnostic process itself contributed to the erosion of the belief that the medical profession would be able to help regain control over health [38,39,41,45,46]. In some cases, this led to mistrust as one participant put it: “The whole thing was disjointed and the worst of it, I was being kept in the dark… I’d lost confidence in the whole [swears] thing” [34,p.322]. When the diagnosis was finally disclosed, and it was explained that there is no real treatment or cure, participants’ previous beliefs that they or medical professionals were able to control their health appeared to be profoundly altered [36,37]. Phrases such as “bombshell” [41,p.943] and “a blow” [32,p.5036] were used to highlight the devastating shock of a terminal diagnosis. Yuan et al. [46] suggest that some people believed their doctor had misdiagnosed them and were in denial about their diagnosis.

Sub theme: Control beliefs about fate

Articles highlighted participant beliefs that through being a “good” person or leading a healthy lifestyle they were able to have some control over their fate, in the context of a long and healthy life [41,42]. Hogden et al. [40,p.832] offered a participant quote: “Because I’m such a healthy person, and to get something like this, you think ‘Ok what did I do?’ It’s not right.”

Studies [41,42,46] demonstrated how diagnosis presented a significant challenge to beliefs in the control of fate. Participants in these studies spoke of feeling that they were being punished and of losing their belief in God post-diagnosis. This quote from one participant illustrates his struggle with the belief that if he led a good life, he would have a long and healthy life:

I have never done anybody any harm… never been in trouble with the police… lead a good life… why me and I thought ‘it’s not fair’ yet people are robbers, the muggers, all the rapists; they live till forever and I got this and I’m not going to live my full term [41,p.942].

“Looking for a reason - why me?” emerged as an inductive code during the synthesis process and this search for causal understanding was a theme throughout the articles [32,36,38,39,41]. This seemed to be something plwMND struggled with soon after diagnosis [39,41]. However, current medical understanding of MND cannot offer any explanations for the cause of the disease nor clear potential risk factors. This lack of causal explanation presented a challenge to participants’ beliefs that they (or others) had some control over their fate, and this appeared to be dealt with in differing ways.

Some people seemed to go through a process of coming to terms with this challenge to their control beliefs concluding that MND was not their fault and the cause was something they had not had any control over [36,41]. The participant quoted above, who initially thought his diagnosis unfair as he had led a good life, over time changed his view to believe that life events were random and that his diagnosis was not connected with whether he had led a good life or not. The authors of this study suggested this to be “successfully navigating these central concerns of maintaining control and reaching causal understanding.” [41,p.945].

Other participants seemed to struggle with the challenge to their control beliefs and were unable to accept that no action or person or divine being had control over the cause [32,40–42]. This could lead to feelings of bitterness, resentment, anger, frustration and despair as demonstrated by this participant’s quote “…neurology controls are useless if you do not know the cause and there isn’t a cure, it’s just a waste of time” [32,p.5039]. Those who were able to accept that their previous control beliefs about their fate needed adaptation post-diagnosis appeared better able to cope with their new life with MND [30,41]:

I have reflected on my granddaughter. Each time I see her – she can do more. And each time she sees me – maybe I can do less. And that’s an interesting balance. I’m not bitter. I could be but I chose not to be [30,p.6]

Sub theme: Control beliefs about the body

The accumulating physical losses caused by MND led to some study participants doubting the belief that they were in control of their bodies [18,28,33,35,39,41,44–46]. It was found that plwMND expressed that their bodies felt and moved differently and no longer reacted to commands [39]. The sense that their body was now in control of them is demonstrated in this quote:

Personally I think it’s [that] my own body is withdrawing from my old life… I’m not doing it deliberately. My body is saying you know pull back from that and that’s what’s happening [33,p.115].
This reduced sense of command over the body was a feature expressed in several studies [28,29,32,41,46] but it was not just the inability to give the body individual commands that were lost. The growing realisation that nothing could control the rate of physical decline [29,40] some plwMND expressed as feeling like their body had been taken over by something else “this alien that has taken over my body and I have not too much defence against it” [33,p.116].

This lack of belief that they were able to control their bodies along with the knowledge that MND can affect almost all areas of the body led to fears of losing all physical control [18,33,42,46]. Some plwMND feared they would become “trapped” [43,p.450] or “prisoners in their own bodies” [42,p.2144], able to understand the world around them but powerless to move or communicate their needs or wishes:

“… I don’t want things keeping me alive when frankly I’d be better off dead” [35,p.1005].

Potentially influenced by traditional culture, participants in China spoke about making the most of their remaining days but seldom discussed death [46].

Those for whom identity was strongly related to their role in paid employment seemed to find the challenge to their control beliefs particularly difficult [29]. Self-image appeared to be strongly related to work identity and for participants having to rely on carers for washing and dressing meant a loss of control over identity which could lead to frustration and resentment, as the following quotes demonstrate:

In business it’s important, you know, dress the part- not like this, I hate this now that I’m wearing” Simon [29,p.349]

And as well she didn’t shave me so I have no shave now and I hate that … I think that might be the worst thing about these motor neurones. Matthew [29,p.347].

Mistry and Simpson [41] suggested that whilst the level of impairment and thus dependence are important, it is perhaps other changes, such as loss of self-image or identity, which affect psychosocial wellbeing. This seems to be reflected in Ando et al. [29] as the above quotes indicate it was not the loss of independence that was an issue but the impact on self-image, so important to these people’s sense of identity, was not being maintained by their carer.

There was also evidence in the studies of some participants attempting to retain the belief that they controlled their identity by finding new roles or activities post-diagnosis [30,32,39,42,44,46]. These included helping the family with manageable tasks, inspiring others with MND and working out practical ways to continue normal activities despite MND. The use of technology to maintain control of identity was highlighted in Pavey et al. where one participant in the study stated “I come across as ‘me’ in email.” [43,p.449].

**Theme 2 – plwMND use a range of control strategies to attempt to retain control in their lives**

Sakellariou et al. [18] suggested that plwMND do not have control over the disease and therefore try to have control over their lives. This theme explores how plwMND attempted to retain control over their lives; evidence of a range of different control strategies used by plwMND was found and will be elaborated upon here.

The significant disruption of long-held control beliefs and the changes to daily life brought about by MND meant that in order to “move beyond the confines of the MND body” [43,p.447] and maintain a life worth living, participants used differing control strategies to attempt to retain some form of control within this new life. Control strategies are what people do to retain control in situations where this has been challenged [47]. The control strategies documented below were all derived from this synthesis of studies.

Although described separately for ease of explanation, these control strategies were not necessarily used separately by plwMND but intertwined with each other and could be adopted or abandoned along the illness trajectory according to their usefulness at the time [28,29,34,35,38,40]. When changes in control strategy were forced due to the speed of MND progression this led to frustration [44]. Due to the unpredictable length of time, people can live with MND there is the opportunity for a change in plwMND’s views over time as the disease progresses [28,38].

What was at one time perceived as an unacceptable loss of control may over time be reconsidered [30,34,35,40]. There is an example of this in Pavey et al. [43,p.447] where participant ‘Andrew’ explained how incidental use of a wheelchair led to a change in his control strategy. After years of struggling with a walking stick, he realised that using a wheelchair could enhance

Sub theme: Control beliefs about identity

Identity appeared to be closely bonded to the roles people played within their lives, whether this was mainly through work, family or community activities, all roles required physical ability to carry them out. The physical losses caused by MND often meant these activities were no longer possible therefore, for plwMND, their role within their lives changed and thus their identity was now no longer within their control [18,29–31,33,39,43,44]. Increasing dependence often led to substantial role changes within family dynamics, with the carer of the family sometimes becoming the cared for [32,46]. Many participants found this change devastating [18,30,32,33,39,41,46]. Cultural identity may also play a role here, Yuan et al. [46] in their study in China found younger plwMND felt guilt that their elderly parents had to care for them because the cultural expectation is that they should care for their parents.

**Pavey et al.** where one participant in the study stated "I come across as 'me' in email." [43,p.449].
the level of perceived control in his life rather than reducing it, as this quote demonstrates:

> Over the years my attitude has changed considerably. Originally, I resisted the need to use a wheelchair and stuck (stubbornly) to using a stick … … Looking back using a walking stick was a ludicrous situation but had anyone talked to me about it, I would have ignored them.

Three control strategies were identified in this synthesis: developing their own illness beliefs, focusing on the present, and maintaining personal values. Each are explained in more detail below.

**Sub theme: Retaining control through developing their own illness beliefs**

The lack of certainty in the available information on illness progression along with conflicting information from healthcare professionals and seeking information from the internet [35,39,40,44] led some participants in studies in the United Kingdom, New Zealand and Australia to feel frustrated with their situation. There was evidence of supportive health professionals [44] but there was also evidence of poor provision of healthcare services in studies set in both the United Kingdom and Ireland [33–35,44].

For example, in Ando et al [29] six of the nine participants expressed problems with their interactions with hospitals, two of these participants felt so negatively that they would not consider a trial of NIV treatment.

This uncertainty of information and in some cases negative experiences with healthcare services could lead to plwMND lacking trust in healthcare services and contribute to an erosion of the belief that healthcare professionals would be able to help them maintain control [18,29,31–33,35,38–40,44].

These issues and uncertainties may have led to some participants feeling that they could not rely upon medical or healthcare professionals and thus they developed their own illness beliefs to attempt to retain a level of control over their illness [18,28,43,44] as explained by “Fay”: “Well I thought the more I walk, the more I'll be able to walk” [43,p.448]. These personal beliefs in illness progression could be stronger than beliefs in treatments recommended by health professionals as Stephen illustrates:

> If I'm not breathing for myself and letting a machine do it I'll get worse quicker than if I just keep breathing on my own [28,p.52].

This could lead to plwMND questioning the need for the treatments recommended by health professionals or seeking alternative therapies [31,44]. Other studies suggested that due to the progressive nature of MND personal beliefs in illness progression may only be a short-term method of retaining control. "Fay," quoted above, initially believed she could control her ability to keep walking but as time progressed, she realised her personal illness belief was not working:

> and now I know I must've looked like a drunken old bag walking down the road. But to me, you know, I thought the more I did the stronger I would become but it wasn't to be … [43,p.448].

**Sub theme: Retaining control through focusing upon the present**

Living day by day and only dealing with issues when necessary, appeared in several studies [18,30,32,35–37,40,42,44]. This strategy appeared to be a way of maintaining some control over an overwhelming situation, as this quote demonstrates:

> to a large extent I keep my head pretty well buried in the sand as far as the reality of it all is concerned. Because as soon as you lift your head up and start to think about these things - you have to every now and then - it all gets a bit overwhelming [40,p.833].

It appeared that for some, it was not just the disease and its consequences that caused them to feel overwhelmed but also the variety and number of healthcare services involved in their care [29,34,38]. For other participants, contemplating their future life with MND was not thought to be helpful [31–33,35,36]. In fact, some participants felt they had “no future” [33,p.115]. The certainty of knowing that they were going to die but the uncertainty of not knowing when that would be was at the forefront of participants’ minds [28,30,34,42,44] as James explained: “It’s a funny situation … .It’s like being on death row without knowing when the executioner is going to pop in” [37,p.1585]. Therefore, as a way of attempting to retain some form of control within their lives, they focused only on the present circumstances [31,35,36,40], for example, not wishing to think about a gastros-tomy until unable to take food orally or only planning a few weeks ahead [44]. The effort involved in just surviving day to day is shown in this quote:

> Until you get to that stage you don't actually focus on it because you're actually focusing on the stage where you're on. And that kind of consumes all your energy and all your time. So, until you actually reach that point you don't deal with it [35,p.1009].

**Sub theme: Retaining control by focusing on maintaining their own personal values**

Some plwMND when faced with the terminality of their diagnosis went through a process of evaluation of their lives and personal values [18,30,40–42]. Supportive relationships became more important and people spoke of strengthening relationships with those who meant most to them [30,36,42,46]. As demonstrated by this quote: “What do I think is important? Having someone who cares about you, that we stand up for each other. These are things I took for granted before” [42,p.2145].

Activities that met the most important values were maintained and those which reduced in importance now life had been limited were discarded [30,42]. Some chose to give up work and focus on family life:

> After I was told that diagnosis and that I would be in a chair within 12 months, I decided to leave work … .I wanted to spend more time with my family. My work wasn’t that important to me [40,p.833].

This focus only on what was important to the individual enabled a feeling of control within their lives and personal measures of disease progression were based upon the ability to continue doing the things which they valued most. Thus, plwMND developed their own personal outcome measures based on their values [18,29,30,44].

If the disease progressed to a stage where tools, equipment or services were required to enable the continuation of the activities which met these personal values, then they were accepted, and daily life adapted accordingly [29–31,43]. For example, Oliver accepted NIV as it enabled him to still go on family holidays, “It’s been good for me using that ventilator. I mean if, for instance going up to his caravan … I don’t think I would have been able to go” [28,p.53].

Frustration was evident if participants could no longer take part in or no longer felt enjoyment from activities that met their personal values [30,44]. If tools, equipment or services did not enable the continuation of an activity that was valued or if they were perceived as taking too much time or effort to master [31], then plwMND did not perceive the need for them [29,33]. Ando et al [29] gave the example of “Catherine” whose breathing had been medically determined as poor but as she was still able to be
in control of the things she valued, she did not perceive that NIV would be of benefit to her. As she said “Actually my own breathing is quite good … I’m usually just sat here during the day or I’m in bed and it doesn’t cause me any problems” [29,p.353]. This shows she was still able to do the activities she valued and hence although her breathing tests may have indicated NIV would be of benefit she did not perceive there to be a need for this.

Discussion

The synthesis identified two analytical themes and seven sub-themes which offer insights into which factors contribute toward perceptions of control and suggest a range of control strategies that people use to maintain control in their daily lives. Here we summarise how this synthesis answered these questions and how the findings bear relation to other literature in the field.

Factors which contribute towards feelings of lack of control

The suggestion in this synthesis that MND can begin to affect personal control beliefs prior to diagnosis appears to be new. However, it relates to research that suggests that the diagnostic process and delivery of the diagnosis is a key stage in the journey of plwMND and can significantly impact both plwMND and their carers [10,48,49]. This synthesis suggests that the dissolving of previously held control beliefs may be involved in plwMND’s ability to come to terms with their diagnosis. Some participant’s beliefs around control of fate quoted in the results have some resonance with the moral model of disability [50], which attributes disability to a moral failing or punishment of past deeds. Research considering MND from the lens of disability theories is lacking but this may be a relevant future area of research. The alterations in control beliefs found in this synthesis significantly affect two key areas involved in adaptation to life with MND - the sense of self [51] and dealing with existential issues [52].

The sense of self and the belief in having control over identity for some participants in this synthesis appeared to be crucial to their ability to live more positively with MND and was also identified by Sakellariou [18] as a way of taking control of one’s life. This however did not appear to be an issue always considered by the authors of the studies included in this synthesis. For example, in Ando et al. [28,p.53], one of the participants “Stephen” expresses “I go to work and some days it’s killing me to go. But I have to keep going and not give in,” the authors interpret this as a resistance to MND itself but perhaps what “Stephen” was expressing was an inability to come to terms with the belief that he is no longer in full control of his identity. He was continuing to battle into work to maintain his identity thus his refusal of NIV may be because he could not see how it would help him retain control over his identity. Whereas the authors, focusing on the physical effects of breathing suggested NIV would have “the potential for the amelioration of his current state” [28,p.54]. In this case, it appears the psychosocial effects of MND are more important to the plwMND than the physical effects focused on by the authors and would suggest that clinical outcomes as assessed at medical consultations may not be the outcomes that matter to plwMND. This would have implications for evidence-based healthcare as the outcomes that matter to plwMND may not be the quantitative outcomes commonly used in quantitative studies and systematic reviews [7]. This suggests that further understanding of the effects of MND on the sense of self from the point of diagnosis is needed. Acknowledgement by medical professionals that for some plwMND psychosocial issues may be of more importance to them than their physical symptoms may aid communication and understanding between plwMND and medical professionals.

The belief that participants were no longer in control of their bodies appeared to lead to consideration of existential issues and the question of when a life is worth living. Due to the wide range of illness duration within the included studies, it is impossible to determine where in the disease trajectory this question became a concern for plwMND but if it is related to belief in control over the body that would suggest that these issues are perhaps considered by patients long before they reach the end of their lives. Terminal diseases are usually considered to be of short duration however for those diagnosed with MND, although they know the disease is terminal upon diagnosis, the time period of disease progression can vary considerably. Lerum et al. [53] suggest that MND inhabits an unusual place somewhere between chronic and terminal illness and thus is perhaps difficult to deal with in healthcare systems organised to treat an illness as chronic until it becomes terminal. This point was noted in this synthesis where these existential issues highlighted did not appear to be dealt with by current healthcare systems, in a range of high-income countries including the United Kingdom, Italy, Ireland and Sweden. Palliative care was not mentioned in the studies as being considered until near the end of life. Palliative care offers a holistic approach that would encompass these existential concerns, this review has highlighted the importance of these issues to plwMND and the provision of person-centred palliative care from the point of diagnosis may enable increased perceptions of control and improve quality of life. The findings of this synthesis would point towards an agreement with research that suggests that palliative care in terminal illness should begin from the point of diagnosis [54,55] and not be left towards the end of life as the participants in the included studies found.

Ways that people living with MND attempt to maintain control in their daily lives

The synthesis suggests that plwMND develop strategies to help them maintain control and that these strategies alter and change as the disease progresses. The impact these strategies may have on healthcare is discussed here.

Retaining control through focusing on the present

The use of strategies by plwMND to deal with losses of control was common throughout the synthesis. Many plwMND spoke of dealing with life day by day and not wanting to think about the future. However, this control strategy appears to conflict with the current professional opinion and guidelines. Recent care guidelines in the UK [56,57] encourage advance or anticipatory care planning perhaps even from the point of diagnosis. However, our findings reveal that plwMND are perhaps not ready to plan for the future and some plwMND feel they have no future or find the future overwhelming.

These differences of opinion appear to cause conflict between clinicians and plwMND, and this point is evident in Hogden et al. [40,p.829] who suggest “clinicians perceive that the patients’ ability to engage in timely decision making is extremely challenging.” This may explain the current interest in research into how plwMND make decisions about their care [58-60].

However, from the plwMNDs’ perspective, reducing their focus to the present time is probably quite effective in helping to control the overwhelming and disempowering effects of a future with MND. Greenaway et al. interpret this as a state of “active
denial or non-acceptance” [35,p.1006] but the findings from our synthesis would suggest that perhaps these people are not denying they have MND but are distancing themselves from it and the future as a method of controlling the situation. Denial can be an effective coping strategy. Indeed, research into Parkinson’s disease, a similarly chronic and progressive illness, has suggested that acceptance and denial may be intertwined, with denial not necessarily being maladaptive [13]. Therefore, it appears that further research is needed to bridge this gap between the clinician’s wishes to plan for future care and plwMND who perhaps use focusing upon the present as a control strategy that works for them.

**Retaining control through focusing on personal values**

Another strong theme through the synthesis was of plwMND focusing upon the activities that mattered to them most thus “deteriorating performance of familiar activities became the benchmark for progression of the disease” [39,p.25]. This focus on personal values appeared to be the basis of plwMND setting their own outcome measures which differed from clinicians. This difference in outcome measures has also been found in previous research by Brown [61] where she considered that professionals and lay people approach MND from differing value standpoints. Clinical care for MND often focuses on symptom management and enhancing quality of life. However, those outcome measures used by clinicians may not be of value to plwMND [62]. These studies appear to support the finding in this synthesis that outcome measures for plwMND are not objective measurements that can be measured with a standardised tool but are very subjective and specific to personal values. For clinicians, this can make treatment difficult as assessment processes tend to use standardised tools which are not patient-specific but for plwMND, these outcome measures used by clinicians may not be meeting their individual needs. This could lead to misunderstanding if the parties involved are working towards different outcome measures.

There has been researching that suggests that patient-perceived outcomes are important in MND and may give a better measure of the quality of life than some standardised tools [63]. However, it is encouraging to note that the recent General Standards for Neurological Care and Support for Scotland [56] have person-centred care as Standard 7 encouraging the management of risk to support personal outcomes thus improving quality of life. These standards were only published in 2020, therefore it will be interesting to observe their impact on care for plwMND.

**Strengths and limitations**

The strengths of this review include the conduct of comprehensive literature searching of six databases using a robust search strategy. The literature screening process involved independent screening of 20% of retrieved references by two reviewers, with strong agreement. The involvement of two senior experienced qualitative researchers to support the junior researcher strengthened all aspects of the review. The included articles represent the views of 578 plwMND aged 20 to 90, from across nine countries. The similarity of perspectives on control found in the synthesis across these nine high-income countries suggests an element of cultural commonalities.

The review was undertaken as an unfunded Masters’s dissertation. Some steps of the rapid review methodology [64–66] were adopted, for example, no grey literature was included and no citation searching was undertaken. Therefore it is possible that relevant studies may have been omitted, however, the omission of a few relevant studies in a qualitative review is not problematic because the purpose is to develop conceptual understanding rather than to draw firm conclusions about the effectiveness of an intervention [67,68]. The studies in this synthesis covered a range of high-income European, Scandinavian and Australasian countries and China, reflecting differing cultural contexts and healthcare systems. Other cultures may have different perceptions of control that were not apparent in this synthesis; and the issues discussed here may not be applicable to other healthcare systems such as those in lower-income countries. Another potential limitation is that only nine of the 19 included primary studies specified the type of MND that participants had and illness duration ranged from one month to 19 years, therefore no conclusions could be drawn about the impact of different types of MND which have different severity levels and duration. This may be important, particularly in research into perceptions of control which may be related to MND severity and duration.

One important issue not acknowledged in this review or in the primary studies included in the review is that of the potential effects of cognitive impairment among plwMND. None of the included studies mentioned any form of cognitive appraisal of participants; it is increasingly being recognised that cognitive impairment may be present therefore cognitive appraisal should be considered in future MND research.

**Conclusion**

This thematic synthesis has explored in-depth the perceptions and experiences of control for plwMND. It has expanded upon the findings of the included systematic review [18] by adding an extra 19 studies to the body of evidence and contributing further depth and detail to their finding that plwMND attempt to regain some control within their lives, revealing the strategies that people actually use.

It is also proposed that control beliefs about health, fate, identity and bodily control are significantly altered by a diagnosis of MND. Due to the rarity of the disease and the diversity of the symptoms, diagnosis can often be a long process and this synthesis found that some alteration of control beliefs may occur prior to diagnosis. This suggests that upon diagnosis, MND may have already had a significant psychosocial impact and this impact should be assessed and considered in care planning. For plwMND, the psychological impacts of a terminal diagnosis are immediate. If the focus of MND care is on treating physical symptoms and not dealing with the extensive psychosocial impact caused by a terminal diagnosis this may not be meeting plwMND’s needs. The factors found which contribute towards a perceived lack of control, such as a lack of palliative care input from diagnosis and the differing viewpoints on the significance of outcome measures would suggest further research is needed into ways that the organisation of healthcare may be contributing towards a perceived lack of control. The finding that some plwMND are perhaps not ready to plan for the future would suggest further research is required to understand plwMND’s immediate needs as well as anticipated needs due to illness progression.

The ability to adapt to alterations in control beliefs appears to affect how plwMND come to terms with their diagnosis and therefore their quality of life. It seems that plwMND are not only dealing with the physical symptoms of MND from an early point in the disease journey but also the psychological, existential and
social impacts which in turn can affect the quality of life. This suggests that the holistic approach of palliative care is required for some plwMND from the point of diagnosis but also that the individual nature of people’s experiences highlights a need for future research to listen to these experiences. The first author is currently undertaking further research using narrative methods to identify issues that may be unaddressed by current health care systems.

This synthesis found some control strategies used by plwMND that conflict with current care guidelines and outcome measures. Focusing on the present, setting personal goals or outcome measures and developing their own illness beliefs all conflicted with health professionals’ standardised systems of care, leading to frustration on both sides. None of the included articles had perceived control as the main focus of their research but the synthesis of these articles indicates perceived control to be a major factor in plwMND’s decision-making and wellbeing. Further research into control beliefs and how plwMND select control strategies would explore the conflicts highlighted in this synthesis and hopefully lead to a more satisfactory healthcare system and a relationship which meets the needs of both the plwMND and the healthcare professionals.

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ORCID

Nicola Glennie http://orcid.org/0000-0002-8857-4417
Fiona M. Harris http://orcid.org/0000-0003-3258-5624
Emma F. France http://orcid.org/0000-0003-0876-7030

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