Decade of progress in motor functional neurological disorder: continuing the momentum

David L Perez,1 Mark J Edwards,2 Glenn Nielsen,2 Kasia Kozlowska,3 Mark Hallett,4 W Curt LaFrance, Jr5

ABSTRACT
Functional neurological disorder (FND) is a prevalent, disabling and costly condition at the neurology–psychiatry intersection. After being marginalised in the late 20th century, there has been renewed interest in this field. In this article, we review advances that have occurred over the past decade (2011–2020) across diagnosis, mechanisms, aetiologies, treatments and stigma in patients with motor FND (mFND, that is, functional movement disorder and functional limb weakness). In each content area, we also discuss the implications of recent advances and suggest future directions that will help continue the momentum of the past decade. In diagnosis, a major advance has been the emphasis on rule-in physical signs that are specific for hyperkinetic and hypokinetic functional motor symptoms. Mechanistically, greater importance has been given to determining ‘how’ functional neurological symptoms develop, highlighting roles for redirected attention, expectation and self-agency, as well as abnormal influences of emotion/threat processing brain areas on motor control circuits. Aetiological, while roles for adverse life experiences remain of interest in mFND, there is recognition of other aetiological contributors, and efforts are needed to investigate links between aetiological factors and mechanisms. This decade has seen the first randomised controlled trials for physiotherapy, multidisciplinary rehabilitation and psychotherapy performed in the field, with consensus recommendations for physiotherapy, occupational therapy and outcome measures also published. Across patients, clinicians, healthcare systems and society, stigma remains a major concern. While challenges persist, a patient-centred integrated clinical neuroscience approach is primed to carry forward the momentum of the past decade into the future.

INTRODUCTION
Functional neurological disorder (FND), also known as conversion disorder, is a common, disabling and costly condition at the intersection of neurology and psychiatry.1,2 While of interest to founding leaders across the clinical neurosciences in the late 19th century, FND was largely abandoned by academics and researchers alike during the late 20th century.3-5 The rationale for these difficulties were based in part on a Cartesian dualism of the brain and mind, limited neuropathophysiologic understanding and few evidence-based treatments.6-8 In the 21st century, a resurgence of interest in FND has occurred, catalysed by improved diagnostic specificity, an expanding ‘toolbox’ of treatments and new pathophysiological models that embrace patient-centred biopsychosocial formulations. A newly formed professional society (www.fndsociety.org), authoritative FND textbooks9,10 and recent special journal issues on this topic have further energised clinical and research efforts in FND.

In this narrative review, we highlight important advancements and their implications for motor FND (mFND) over the past 10 years (2011–2020)—spanning functional movement disorder and functional limb/face weakness. We use a transdiagnostic approach across the range of functional motor symptoms given high phenotypic overlap across populations (eg, functional tremor with concurrent functional weakness in the same limb). Isolated functional (psychogenic non-epileptic/dissociative) seizures, functional speech/voice disorder, functional cognitive disorder, functional sensory deficits and the spectrum of functional somatic disorders are beyond the scope of this article and have been reviewed elsewhere.11,12 Sections here detail recent developments in diagnosis, mechanisms, aetiological factors, treatments and stigma in patients with mFND. In each content area, future directions are also suggested, aimed at continuing the momentum of the past decade.

DIAGNOSIS
New developments
Establishing the diagnosis of mFND has been made more practicable, as physical examination findings with diagnostic specificity have been identified (eg, Hoover’s sign with an estimated specificity of 95.7%–99.9%).13 Educational efforts have also made neurologists more confident in their ability to accurately diagnose patients with mFND, encouraging extensive laboratory testing unless a comorbid neurological disorder is suspected.7

The Diagnostic and Statistical Manual of Mental Disorders-Fifth Edition (DSM-5) criteria for FND include the diagnostic features of inconsistency and incongruity on examination, emphasising positive neurological features; identifying an underlying psychological trauma has been relegated to a discussion note and removed as a criterion.14 Inconsistency refers to changes in manifestation over time, such as variation in tremor frequency and amplitude or remissions and exacerbations. Incongruity refers to discordance with other known neurological disorders or human anatomy and physiology. Additional general diagnostic features
are distractibility and suggestibility. Most other diagnostic signs are phenotype specific and can be augmented by certain clinical neurophysiological tests. Clinical neurophysiological tests (eg, electromyography-assisted identification of tremor pause during ballistic movements) can either objectify bedside observations or identify features that clinicians cannot readily appreciate.

Positive signs on examination that characterise functional limb weakness (and functional sensory and gait disorders) have been analysed for their statistical properties (including sensitivities and specificities) and then subjected to prospective analysis, including inter-rater reliability. Reliable signs of functional limb weakness include give-way/collapsing weakness, drift without pronation, cocontraction, Hoover’s sign, hip abductor sign, Spinal Injuries Center Test and weakness of the sternocleidomastoid with hemiparesis.

Hyperkinetic mFND presentations include tremor, myoclonus and tics (jerky movements), dystonia, parkinsonism and gait disorders. Clinical and laboratory features are described in Table 1. Tremor and myoclonus are readily identifiable with established clinical features and excellent neurophysiological tests that can be used for confirmation in ambiguous cases. Differentiating functional versus neurogenic motor tics can be difficult if the presence of an urge or sensory tic is not present or uncertain. The presence of a Bereitschaftspotential (readiness potential) prior to the movement is common in functional jerky movements.

Tremor
- Entrainment (tremor takes on the rhythm of paced movements performed with another body part)
- Pause with quick movement of another limb
- Variability in frequency, amplitude
- Tonic contraction at onset
- Increase in amplitude with weighting
- Coherence of tremor between two limbs
- Whack-a-mole sign (restraint of tremor induces tremor in another body part)

Myoclonus
- Variability and long duration of the movement
- Complex movement
- Appearance of startle
- Long and variable latency of stimulus induced jerks
- Jerks when tendon hammer stops short of contact

Tic
- Lack of urge
- Lack of voluntary control (suppressibility)

Dystonia
- Certain patterns such as fixed dystonia or pulling lip to one side

Parkinsonism
- Marked slowness or incoordination in examination but not with normal movements
- Gegenhalten (variable resistance during passive movement)
- Lack of sequence effect (slowness without amplitude decrement during repetitive movements)
- Huffing and puffing sign (fatigue with minimal effort)

Gait disorders
- Specific patterns including knee buckling, dragging a monoplegic leg, astasia-abasia, excessive slowness and atypical limping
- Better balance than claimed, including improvement with distraction
- Either no falls, controlled falls or falling toward support
- Chair test (can use legs to move a chair better than walking)

Intermittent lip deviation to one side is commonly identified functional patterns. Functional facial spasm, a common stroke mimic, is characterised by platysma hyperactivation, jaw deviation and ipsilateral eyebrow depression.

In support of the stability of an mFND diagnosis based on examination signs, a 14-year prospective study in 76 patients with functional limb weakness showed only a 1% misdiagnosis rate. Notably, some patients have both functional neurological signs and other neurological conditions, such as associations with multiple sclerosis, Parkinson’s disease and other neurodegenerative disorders.

Implications
Improved diagnostic specificity has made it easier for neurologists to present the diagnosis to patients, which is the first step in treatment. It is generally necessary for the patient to agree with the diagnosis or, at least, allow for the possibility of such before moving onto additional treatments. Additionally, it has been suggested that it is helpful to demonstrate positive signs to the patient to show how the diagnosis was made. Increased diagnostic specificity also permits identification of cohorts with content validity for research studies.

Future directions
There is a need to further test the specificity, sensitivities and inter-rater reliability of the growing range of positive functional signs compared with other neurological populations, particularly given that statistical properties for some signs have been only tested in a single cohort. Additionally, functional dystonia remains among the most challenging mFND diagnoses. The overlap of clinical features and neurophysiological tests with other dystonia subtypes remains obscure and needs to be explained. However, there are some promising tests (eg, blink reflex recovery and paired-associative stimulation induced plasticity) that need further validation.

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**Table 1** Positive clinical features (phenotype specific) and useful adjunctive laboratory tests for the diagnosis of functional movement disorder

<table>
<thead>
<tr>
<th>Functional motor symptom</th>
<th>Clinical features</th>
<th>Laboratory tests</th>
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<tbody>
<tr>
<td>Tremor</td>
<td>Entrainment</td>
<td>Clinical neurophysiological measurements can quantify entrainment, pause with quick movement, variability, tonic contraction at onset, increase amplitude with weighting and coherence between limbs</td>
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<tr>
<td></td>
<td>Pause with quick movement of another limb</td>
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<td></td>
<td>Variability in frequency, amplitude</td>
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<td>Certain patterns such as fixed dystonia or pulling lip to one side</td>
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<td></td>
<td>Normal blink reflex potential</td>
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<td></td>
<td>Normal plasticity with paired associative conditioning</td>
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<tr>
<td>Parkinsonism</td>
<td>Marked slowness or incoordination in examination but not with normal movements</td>
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<td>Lack of sequence effect (slowness without amplitude decrement during repetitive movements)</td>
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<td></td>
<td>Chair test (can use legs to move a chair better than walking)</td>
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Although relevant to only a small minority of cases, there is also a need to better distinguish mFND (where symptoms are experienced as involuntary) from factitious disorder and malingering; in both factitious disorder and malingering, there is conscious feigning of symptoms. The latter two presentations are rare but unfortunately influence physician attitudes toward patients, and they should be addressed differently.

Another challenge is to identify adjunctive diagnostic biomarkers. Quantitative neuroimaging alone in its current form may not provide the answer. Neuroimaging findings are valuable in understanding the neuropathophysiology of mFND, but are sufficiently subtle (and heterogeneous) that they will likely not have high sensitivity and specificity on an individual basis. While neurophysiological testing shows encouraging value in some circumstances, it will be important to show high specificity when used in relevant uncertain clinical circumstances. Composite diagnostic biomarkers across multiple neurobiological data points (electrophysiology, neuroimaging, autonomies, etc) also warrant future investigation.

Lastly, the field needs to better contextualise the overlap between mFND and other FND subtypes (eg, functional seizures), as well as to explore optimal approaches to contextualise other bodily symptoms frequently present in patients with mFND that closely relate to quality of life (eg, pain, fatigue, cognitive symptoms). Furthermore, the intersection of FND, functional somatic disorders (eg, fibromyalgia) and other neuropsychiatric conditions (anxiety and trauma-related disorders, somatic symptom disorders, mild traumatic brain injury, etc) requires clarification.

**MECHANISMS**

**New developments**

An important recent focus of mechanistic theorising has been to shift the typical viewpoint from which mFND has been studied. Traditional ‘Freudian’ and related viewpoints have been, arguably, ones that prioritise aetiological factors over mechanisms. In other words, the precise mechanics of how a particular functional motor symptom arises has not been of high concern, and instead emphasis had been almost exclusively on the influence of hypothesised stressors and psychological factors.

Emphasis on mFND mechanisms has drawn on a broad neuroscientific knowledge base, including from the fields of motor control (eg, the underpinnings of sense of agency), cognitive-affective neuroscience (eg, attention and emotion processing) and computational neuroscience. One of the important questions such work has sought to answer is: if functional neurological symptoms are truly involuntary, what are the implicated brain mechanisms of these unconscious processes?

Research has started to coalesce around the idea that there is a mechanism (or set of inter-related mechanisms) which mediates the relationship between conscious experience of movement control and the neural networks that enable movement and related sensory experiences to occur. In one expression of this idea grounded in the computational theory of active inference, perception and movement control rely on a dynamic relationship between actual sensory data and predictions about these data. The relative weighting of these ‘bottom up’ and ‘top down’ sources of information (known as precision) can be influenced by attentional focus (including modulation via limbic/salience networks). The suggestion in mFND is that abnormally strong predictions, relevant to symptoms such as weakness, tremor and gait difficulties, develop and are made more precise by body-focused attention. This drives symptom production in line with abnormal predictions and overwhelms contradictory sensory evidence. Notably, efforts are underway to test these theories, such as the recent use of the ‘broken escalator’ paradigms probing non-conscious and conscious forms of motor learning in patients with functional gait disorder to identify persistence of a locomotor after effect (representing a failure of deadaptation) (see figure 1). This mechanism can also lead to motor symptoms without a sense of agency, with functional neuroimaging studies in mFND implicating the right temporoparietal junction/inferior parietal lobule in deficits in action authorship perceptions.

Recent work, using neuroimaging and other experimental approaches, has also begun to contextualise the role of emotion/threat processing in the pathophysiology of mFND. This research has sought to elucidate the way in which networks relevant to voluntary movement might be abnormally influenced ('hijacked') by networks serving affective and threat processing. Noteworthy findings include: (1) a direct effect of recall of relevant traumatic life events on supplementary motor area activation; (2) abnormal connectivity between motor control areas and amygdala/insula brain areas during rest and affective provocation and (3) altered temporoparietal junction and insula cortex connectivity in the resting state.

**Implications**

An important implication of this work has been to support mFND as a brain-based condition. In this sense, it is simply a process of the mFND field catching up with the rest of neuro-psychiatry, benefiting from recent neuroscientific advances, building on historical concepts, which further bridges neurology and psychiatry. However, this change has a danger of creating a solely neurocentric view of mFND, ending up swapping one extreme viewpoint (psychology only) for another (neurology only)—a sentiment that we caution against.

**Future directions**

The path forward is one that continues building an integrated, mechanistic framework for mFND that is neither exclusively psychological nor neurological. A key future goal is to attempt to unpack, at an individual level, the different influences on symptoms, aetiologies, treatment response and prognosis. For example, efforts to integrate active inference principles not only for sensorimotor percepts but also pertaining to interoception and ‘emotion making’ based on the theory of constructed emotion may provide additional mechanistic advances in mFND. How mFND neural mechanisms relate to treatment mechanisms and clinical outcomes is critically important and under-researched. Additionally, it remains unclear if outward presenting phenotypes (eg, functional tremor) are driven by the same set of mechanisms across all patients or if there are a range of biological mechanisms that may lead to the same clinical phenotype. If neural mechanisms differ across patients with similar phenotypes, it will be important to understand if biologically informed subtypes are linked to specific treatment response and prognostic profiles. Such observations, if robustly elucidated, would facilitate the use of precision medicine in mFND care. Relatedly, research is needed to investigate if there are common neural mechanisms across FND and the spectrum of functional disorders across medicine. To
AETIOLOGICAL FACTORS

New developments

Over the past decade, aetiological research in mFND contextualising predisposing vulnerabilities have identified the presence of a number of potential putative contributing factors, while at the same time, acknowledging the importance of individual differences. A systematic review and meta-analysis showed that the odds of being diagnosed with FND was 3.9 times higher given childhood physical abuse compared with controls and 3.3 times higher given childhood sexual abuse.27 In a separate systematic analysis, the odds of being diagnosed with FND was 3.9 times higher given childhood sexual abuse compared with a range of mood, anxiety, personality and pain-related disorders.28 A systematic review and meta-analysis have also demonstrated the inter-relatedness between adverse life experiences and other predisposing vulnerabilities for the development of mFND, such as fearful attachment styles independently correlating with childhood abuse burden, alexithymia and depression scores.29 These findings highlight the importance of considering the relevance of adverse life experiences in mFND populations using stress-diathesis and neurodevelopmental perspectives, emphasising the interplay between biological (genetic/epigenetic) risk, life events and precipitating (triggering) factors.28 29 While in early stages, pathophysiology studies have started to contextualise the neurobiological importance of childhood maltreatment in promoting the development of mFND.24 Two examples from the functional MRI literature include: (1) the observation that resting-state connectivity strength between salience/limbic network brain areas (amygdala, insula) and the precentral gyrus correlated with the magnitude of previously experienced childhood physical abuse in patients with mFND and (2) the finding that the G-703T polymorphism (rs4570625) in the tryptophan hydroxylase-2 (TPH2) gene moderated the relationship between childhood trauma and functional movement symptom severity; differential amygdala-prefrontal connectivity profiles were also identified in patients with mFND based on TPH2 genotype (see figure 2).31

Acknowledging that not all patients with mFND endorse adverse life experiences, risk factors for mFND extend beyond these considerations. A heightened bodily attentional focus, at times to the decrement of perceptual accuracy, has been characterised in patients with mFND.32 33 Altered bodily attention and increased arousal may also help explain associations between physical injury and the subsequent development of mFND,34 given that physical injury promotes heightened attention to the self and activation of bodily arousal systems.35 The traditional conceptualisation of several demographic and psychosocial factors has also been challenged, including the increased appreciation of mFND symptoms in older populations (eg, Parkinson’s disease)36 and findings that patients with mFND and neurological controls have similar histories of employment in healthcare related fields.37 Psychiatric diagnosis are common (eg, 1/3 of patients with mFND meeting criteria for major depressive disorder), yet are not universally present
in all patients with mFND. Trait psychological constructs remain important, such as the finding that alexithymia, independent of depression scores, was elevated in patients with mFND compared with neurological and healthy controls; patients with mFND and prominent alexithymia also exhibited higher rates of obsessive-compulsive personality disorder. Novel risk factors for the development of mFND have also been identified, such as aberrant sensory and information processing.

Implications
Identification of a broad array of relevant, yet non-deterministic, risk factors for developing mFND suggests that links between aetiological factors and disease mechanism remain incompletely understood. More specifically, while adverse life experiences remain important vulnerabilities for developing mFND and are linked to other predisposing and perpetuating factors, the presence or absence of these events neither helps rule in nor rule out a diagnosis of mFND.

Future directions
Additional research is needed to understand the intersection of disease mechanisms, aetiological factors and treatment response within the context of the biopsychosocial framework (including spiritual and cultural influences). A precision medicine approach may be needed to not only link psychosocial risk factors to brain circuits but also to contextualise a range of relevant mediating and modulating factors including genetic/epigenetic information. The importance of developmental trajectories (including critical periods), gene–environment interactions and sex differences are also underexplored factors that may help better explain connections between risk factors and the later-life development of mFND. Given significant child maltreatment in a subset of patients with mFND, future research may also inquire if there is a ‘trauma subtype’ of mFND, while also clarifying important risk factors in patients who lack such a history. The relevance of aetiological factors to treatment selection and response (eg, targeting concurrently present PTSD symptoms as an approach to treat mFND) also requires more research inquiry.

TREATMENTS AND PROGNOSIS
New developments
The late 20th century’s lack of interest and investment in mFND by healthcare systems is reflected in exceedingly few care programmes for this population, which in turn is reflected in patients feeling marginalised and unable to access treatments.

With the DSM-5 modifications, the neurologist’s (and other clinician’s) role in mFND has now expanded to making a positive ‘rule-in’ diagnosis, communicating the diagnosis effectively and facilitating access to additional treatments. ‘How to’ articles have disseminated good clinical practices on the delivery of the diagnosis and longitudinal care. Specialist FND clinics, often led jointly by neurologists and psychiatrists, have also been developed in some countries for complex cases (eg, those with diagnostic uncertainty, multiple comorbidities). With the time required to adequately manage this population, challenges have been raised regarding clinical bandwidth.

The website neurosymptoms.org has become a valuable educational resource for patients and clinicians. Other information websites have been created, including from patient support charities (eg, fndhope.org, fndaction.org.uk). The efficacy of online information and self-help used in isolation was assessed in a randomised controlled trial (RCT). At 3 months, there was no difference in improvement on self-rated health or in secondary outcomes between groups. This suggests that online education, while generally rated favourably, is inadequate as a stand-alone treatment.
There has been a rise in physical rehabilitation and multidisciplinary research. Since 2010, no less than 17 rehabilitation [38] studies have been published in the treatment arm in measures of physical health that have compared participant improvements compared with 18% community neurophysiotherapy, with 72% of the intervention [2] published in 2014, compared a 3-month feasibility [8] of two RCTs, 7 prospective and 8 retrospective. The first RCT, cohort studies of patients with mFND have been published [41].

<table>
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<tr>
<th>Study</th>
<th>n</th>
<th>Description</th>
<th>Points</th>
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<tr>
<td>Physiotherapy</td>
<td></td>
<td>Randomised feasibility study of specialist physiotherapy versus usual care</td>
<td>The intervention was delivered by physiotherapists and included education, treatment retraining and self-management. The control was standard community neurophysiotherapy. At 6-month follow-up, 72% of the intervention group reported symptom improvement, compared with 18% of the controls. Significant improvement was seen in a range of physical and quality-of-life outcome measures. The intervention was associated with a gain in quality adjusted life years and an incremental cost effectiveness ratio that was suggestive of a cost-effective intervention.</td>
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<tr>
<td>Nielsen et al[38]</td>
<td>60</td>
<td>Duration and setting: 5 days in an intensive outpatient/day programme</td>
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<td>Outcome measures: Feasibility (recruitment and retention rates, intervention fidelity, acceptability); SF-36; WoAS; EQ-SD-5L; DASH; CGI-patient rated; FMS, BBS</td>
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<tr>
<td>Multidisciplinary rehab</td>
<td></td>
<td>Randomised study of multidisciplinary rehabilitation versus a wait list</td>
<td>This is the first and currently only randomised study of multidisciplinary rehabilitation (described by the authors as ‘adapted physical activity with a cognitive behavioural framework’). Post treatment, there was a significant difference between groups in physical and quality-of-life outcome assessments. Treatment gains were, for the most part, maintained at 12-month follow-up.</td>
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<tr>
<td>Jordbru et al[39]</td>
<td>60</td>
<td>Duration and setting: 3-week inpatient programme</td>
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<td>Outcome measures: FIM, FMS, SF-12</td>
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<tr>
<td>Cognitive behaviour</td>
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<td>A pilot, single-blinded randomised study, comparing CBT alone versus CBT</td>
<td>The two CBT containing interventions (with and without physical activity) showed reduction in PMDRS over 12 weeks. SMC showed no improvements.</td>
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<tr>
<td>Dalocchio et al[40]</td>
<td>29</td>
<td>plus physical activity versus SMC</td>
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<td></td>
<td></td>
<td>Duration and setting: 12 weeks of outpatient 90-minute CBT with or without</td>
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<td>adjunctive outpatient physical activity (60 min, two times per week)</td>
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<td>Outcome measures: PMDRS, PHQ-15</td>
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<tr>
<td>Sharpe et al[41]</td>
<td>127</td>
<td>Randomised controlled trial of CBT-based guided self-help plus usual care</td>
<td>Participants allocated to self-help CBT reported greater improvement on the primary outcome of self-rated health (CGI). At 6 months, the treatment effect was no longer statistically significant.</td>
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<td>versus usual care alone</td>
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<td>Duration and setting: self-guided outpatient CBT plus four 30-minute</td>
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<td>face-to-face guidance sessions over 3 months</td>
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<td></td>
<td>Outcome measures: CGI-patient rated; SF-12; PHQ-13</td>
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<tr>
<td>Botulinum neurotoxin</td>
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<td>Randomised, double-blinded controlled trial of botulinum neurotoxin</td>
<td>At 4 months, there were no statistically significant differences between treatment arms across primary and secondary outcomes. However, improvement was observed across both treatment arms (56%–64%), suggesting a notable placebo effect. Across the length of the entire trial, 81% improved from baseline.</td>
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<tr>
<td>Dreissen et al[42]</td>
<td>49</td>
<td>versus placebo (sterile saline) injections</td>
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<td>Duration and setting: 2 outpatient injections 3 months apart, followed by a</td>
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<td>10-month open-label extension</td>
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<td>Outcome measures: CGI-clinician and CGI-patient rated; PMDRS; SF-36; AMC</td>
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<td></td>
<td>Linear Disability Score</td>
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<tr>
<td>Vizcarr et al[43]</td>
<td>14</td>
<td>Randomised controlled trial of botulinum neurotoxin versus placebo</td>
<td>There were no differences in clinical outcomes at 12 weeks. While both</td>
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<td>(sterile saline) injections, followed by CBT</td>
<td>treatment arms showed a tendency toward improvement, a statistically</td>
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<td></td>
<td></td>
<td>Duration and setting: outpatient injection followed by 12 weeks of psychotherapy</td>
<td>significant change from baseline was only observed in the placebo-CBT group.</td>
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<tr>
<td>Online education and self-help</td>
<td></td>
<td>A randomised controlled trial of internet-based education and self-help plus</td>
<td>No additional treatment benefit was found in the intervention group; however, the participants valued online information. This suggests online self-help is not on its own an effective treatment.</td>
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<td>Gelauff et al[44]</td>
<td>186</td>
<td>versus usual care and self care alone</td>
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<td></td>
<td></td>
<td>Duration and setting: online access with outcomes evaluated at 3 and 6 months</td>
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<td></td>
<td>Outcome measures: CGI-patient rated; RAND36; WSAS</td>
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</table>

* Of the 127 participants, only approximately one-third had motor symptoms (weakness or tremor); the exact number of participants with motor symptoms is not specified. Additionally, outcome measures listed in this table focus on the selected physical functioning and quality-of-life instruments used in each study. Two small sample size randomised controlled trials using psychodynamic psychotherapy approaches are not shown here, but are detailed in online supplemental table 1.

ADL, activities of daily living; AMC, Academic Medical Center; BBS, Berg Balance Scale; CBT, cognitive behavioural therapy; CGI, Clinical Global Improvement Scale; DASH, Disabilities of the Arm, Shoulder and Hand; FIM, Functional independence Measure; FMS, Functional Mobility Scale; IADL, Instrumental ADL; Katz Index of Independence in ADLs; Lawton instrumental ADL; PMDRS, Patient-Mediated Disability Rating Scale; RAND36, Dutch equivalent of SF-36; SF-12, Short Form 12; SF-36, Short Form Health Survey 36; SMC, standard medical care; WSAS, Work and Social Adjustment Scale.
self-directed attention is usually emphasised as a factor exacerbating symptoms. Physical therapies aim to retrain movement with diverted attention, and physical interventions are informed by a psychological understanding of symptoms (eg, addressing fear-avoidance behaviours using graded exposure).

As well summarised in a recent systematic review, the major advance for psychotherapy in mFND is that initial RCTs have been conducted and published—after a dearth of controlled data in decades prior. Examples of the interventions are described below and summarised in table 2 and online supplemental table 1.

Psychotherapy trials for mFND include a pilot single-blind RCT of 29 patients with mFND (mostly functional tremor) randomised to receive 12 weeks of conventional CBT alone (90-minute session, once a week) versus CBT+adjunctive physical activity (APA) (60-minute session, two times per week of low-intensity/moderate-intensity walking). The control group consisted of eight patients receiving standard medical care (SMC). The CBT intervention focused on the interplay of somatic misinterpretations, negative thoughts, illness beliefs and low mood or anxiety, along with use of distraction, relaxation and other problem-solving techniques. The two CBT containing interventions (with and without APA) showed improvements in functional motor symptoms, depression and anxiety scores at 12 weeks, while the SMC arm showed no significant improvements. A prospective single-arm study in 15 patients with functional tremor also demonstrated the efficacy for CBT in reducing tremor severity. Furthermore, an RCT of self-guided CBT in 127 patients with mixed FND randomised to CBT+usual care (n=64) versus usual care alone (n=63) showed a statistically significant improvement in patient-rated global improvement at 3 months for those receiving CBT; reductions in somatic symptom burden and health anxiety were also observed. These gains were no longer significant at 6-month follow-up. See online supplemental table 1 for details regarding two small psychodynamic psychotherapy RCTs, as well as other psychotherapy cohort studies in mFND populations.

Regarding paediatric mFND—while there are ethical and practical challenges to performing RCTs in this population—efficacious multidisciplinary programmes generally combine psychotherapy, physiotherapy, occupational therapy and family work targeting focus of attention and pertinent stressors and school attendance/reintegration (see online supplemental table 1).

There remains little evidence for pharmacological therapy in the direct treatment of mFND symptoms, yet medications have a role in managing concurrently present anxiety, depression, migraine and insomnia. Regarding other treatments, a recent randomised placebo-controlled trial of botulinum neurotoxin (BoNT) for jerky and tremulous functional movement disorder (n=48) found no benefit compared with placebo. Here, approximately two-thirds of patients in both groups improved, demonstrating a large placebo effect. A similar positive placebo response was observed in a pilot randomised trial of BoNT followed by 12 weeks of CBT in patients with functional dystonia (n=14). While placebo effects are important considerations, there is an argument to be made for the use of BoNT in patients with chronic symptoms that have not benefited from other treatments. Transcranial magnetic stimulation (TMS) also continues to be investigated as a promising therapeutic, although disentangling circuit-level neuromodulatory effects from placebo remains challenging. When placebo is considered the ‘active ingredient’, there remains debate regarding how transparent to be with patients (we favour an open and transparent stance).

Given that non-specialist clinicians may feel ill prepared to assess and manage patients with mFND, expert opinion-based recommendations and practical advice are welcomed additions. These include:

- Assessment and diagnosis of mFND symptoms.
- Neuropsychiatric assessment.
- Delivering the diagnosis (including providing clear, empathic communication with a cautiously optimistic stance for improvement).
- mFND presenting to stroke services.
- Physiotherapy.
- Occupational therapy.

Consensus recommendations to standardise outcome measures for clinical trial research in FND have also been published, emphasising patient-reported data.

Regarding prognosis, a systemic review of long-term follow-up studies from 10 to 491 individuals reported that 39% of patients across the spectrum of FND were the same or worse and the majority (approximately 80%) remained symptomatic. The same research group recently published a 14-year follow-up study in 76 adults with weakness, identifying that 20% had symptom resolution, 31% improved, 23% were the same and 26% were worse. In terms of discrete prognostic factors in adults, findings have been inconsistent and understudied in more recently developed care models. Outcomes from specialist paediatric multidisciplinary programmes are more optimistic with approximately three quarters of children returning to full health and full-time school attendance. Outcomes are less favourable for children with chronic mFND symptoms at presentation; those with cognitive vulnerabilities, whose comorbid mental health disorders or other (comorbid) functional somatic symptoms do not resolve and those who subsequently develop chronic mental health problems.

Implications

While delivery of the diagnosis is the first step in treatment, online self-help information alone is insufficient for symptom reduction and should not be considered definitive treatment. Likewise, self-help psychotherapy approaches appear to lack durability in maintaining improvement. Careful assessment is needed to triage patients towards the most suitable treatment based on available options, including physiotherapy, skills based psychotherapy and/or multidisciplinary interventions. Given evolving care models and lack of robust predictors of prognosis, those with chronic symptoms, formerly considered refractory, should not be excluded from evidence-based treatments.

Future directions

To further advance mFND treatments, future research should continue to pursue fully powered RCTs across rehabilitative and psychological interventions. Studies examining optimal treatment setting(s) are also needed.

An important future direction could be to develop specific interventions that are tailored both towards the mFND phenotype (eg, weakness, tremor, dystonia) and the wider clinical syndrome. For example, in addition to motor symptoms, common comorbidities and other health-related problems could be considered within a single-treatment package (eg, PTSD, anxiety, chronic pain, migraine, joint hypermobility, social difficulties, etc). The timing of the different treatment elements may also be important, and the value of a modular approach to treatment could be explored, where the focus of whole-person treatment can be personalised and evolve according to the patient’s
biopsychosocial clinical formulation. As an example, a patient with a functional gait disorder and major depression with suicidality may benefit from physical therapy at some point, but it is probably more important initially to treat aggressively their depression and existential concerns. Conversely, a patient with sudden onset disabling physical symptoms may need to make some initial progress with physical therapy (eg, to regain sitting balance) before engaging well in psychotherapy.

Additional research should explore the development of technology-based adjuncts (informed by advances in elucidating mFND pathophysiology) and innovations to improve access to specialist treatment (eg, tele/remote health as has been used in other FND subtypes, virtual reality, wearable technology, biofeedback, TMS, etc). More aggressive focus on psychosocial factors might also be useful. Additional work is needed to define the most suitable clinical outcome measures and to also determine if the creation of new FND-specific outcome measures may be beneficial. Further clarifying neural mechanisms and predictors of treatment response will also be important, offering the potential to develop novel psychologically and biologically informed treatment interventions. Finally, it is crucial that more treatment programmes are developed; it will not do any good to find optimal treatment strategies if they will not be available to the majority of patients.

STIGMA
New developments
Stigma pertaining to the diagnosis of mFND is increasingly recognised as an important, multifaceted issue requiring clinical and research attention. In mFND, stigma represents a complex interplay between patients, clinician–patient relationships, healthcare systems and sociocultural factors. The very fact that mFND sits at the intersection of neurology and psychiatry challenges deeply rooted medical and societal norms of health and disease. Furthermore, the variability/distractibility seen in many individuals mistakenly perpetuates a framing that symptoms are voluntary (when perceived nonetheless as involuntary by the patient). These and other nuanced issues related to stigma can be discussed across three levels—public stigma, personal self-stigma and patient label avoidance—and we use these categories to frame our discussion (see box 1).

Box 1 Three different aspects of stigma

Stigma category and definition

► Public stigma occurs when the general population—or certain subsets of the population—endorses negative beliefs pertaining to a certain illness and acts on these beliefs in a discriminatory manner, often by avoidance and withdrawal.

► Personal stigma occurs when the individual person—child or adult or the family—becomes aware of the negative beliefs about a certain illness, internalises these beliefs and applies them to the self.

► Patient label avoidance refers to the patient’s reluctance and efforts to distance himself or herself from a label—in the case of motor functional neurological disorder, a diagnosis—because the label is perceived as being socially unacceptable. This type of stigma is more commonly perceived in relation to mental health disorders, because such disorders are commonly misperceived as being the result of personal weakness or poor character.

With the growth of scientifically based medicine and the complexities in how to understand functional neurological symptoms, mFND was sidelined and publicly stigmatised into the category of ‘medically unexplained’ disorders in the late 20th century. Interest in mFND waned reflected in mFND largely disappearing from medical textbooks, educational curricula and bedside teaching. Relatedly, many physicians avoided the diagnosis—or used terminology that was offensive (eg, confusing mFND with malingering)—leaving patients perplexed or angry and setting up a negative process whereby patients sought help from multiple physicians, were subjected to repeated unnecessary diagnostic tests and treatment was delayed or not provided.

Public stigma regarding mFND has unfortunately continued. Some physicians caring for patients with mFND may manifest their discomfort in non-verbal communication patterns that convey uncertainty, negative treatment expectations and/or their own perceptions that the diagnosis is a delicate (stigmatised) matter that needs to be managed cautiously (or at least not by a neurologist). In doing so, physicians undermine their own capacity to use positive suggestion to influence belief—relevant in the treatment of mFND—and core to the art of healing in all of medicine. Patients can also have negative interactions with other healthcare professionals (eg, nurses, administrative staff), coworkers, friends and family—driven in part by the conceptual misunderstandings outlined above. The general lack of dedicated services in hospitals and the failure to include FND in national research priorities also communicate a powerful message that mFND is seemingly unimportant.

Patients with mFND may internalise stigma, make it personal and report being harmed by it. Internalisation of stigma promotes feelings of vulnerability, helplessness, hopelessness, frustration and anger. Stigma contributes to the shaping of patients’ own internal beliefs and expectations, which may limit their ability to improve. These factors also contribute to negative doctor–patient interactions.

As a consequence, some patients find it difficult to accept an mFN diagnosis and others reject the diagnosis altogether or litigate the physician. This label avoidance contributes both to clinician anxiety and potentially to endless doctor shopping (and medical procedures) in hopes of receiving a different ‘medical’ diagnosis. Sometimes label avoidance even propels patients to accept explanations that are outlandish or that conceptualise their mFND as a ‘medical mystery’.

Changes in how the diagnosis is communicated to patients are occurring, including more clinicians using the term ‘functional’. Educational efforts are underway to help physicians avoid the pitfalls of oversimplified explanations to patients that ‘it is all stress-related’ and moving towards describing ‘stressors’ in the context of life events. The Multidisciplinary FND Society, several authoritative FND textbooks, growth of specialised treatment programmes and high impact publications advocating for change and research funding indicate that mFND is re-entering mainstream medicine.

Implications
Public stigma, personal self-stigma and patient label avoidance in mFND remain major concerns, and efforts to mitigate stigma need to be driven by clinicians, researchers, patients, advocacy groups and policy-makers. A helpful transition that has occurred in recent years is the framing of mFND at the intersection of neurology and psychiatry (synonymous with a core neuropsychiatric disorder), and rigorous pathophysiology research in mFND is also decreasing stigma by advancing a brain-based, mechanistic
understanding of the condition. Unfortunately, mental health is stigmatized in many societies and healthcare systems in comparison to medical/neurological conditions. This is a notable issue, given that the multidisciplinary approach to mFND patient care includes important roles for mental health clinicians. Treatment implications that emerge from this body of work highlight the importance of the biopsychosocial model—the body, mind and family and social context—to address each patient’s particular presentation and issues. The mFND field needs to be cautious so as to strike a good balance between neurological and psychological/psychiatric conceptualisations of the disorder. Moreover, mental health professionals need to be empowered at the same level of engagement as neurologists, rehabilitation specialists and research scientists, as we work collaboratively for the benefit of patients.

Future directions

In addition to increasing advocacy to further decrease stigma pertaining to mFND and related conditions across medicine and society, mFND can serve as a model condition through which to challenge the inherently artificial divide between physical and mental health that is pervasive in medicine and society—a dualism exposed most directly by mFND. This condition also offers the potential to bring the disciplines of neurology and psychiatry, two specialties for the same organ system, increasingly together to leverage an integrated clinical neuroscience perspective.

CONCLUSIONS

Significant advances have occurred in the past decade in the diagnosis, conceptual understanding and treatment of mFND. This progress reflects a renaissance taking place across the field of FND, bringing neurology and psychiatry together again, as we are confronted by the need for multidisciplinary and collaborative care models for this complex population. With a biopsychosocial understanding of mFND and with better neuroscience tools to provide rigorous biological phenotyping and classification, we are poised to design well-constructed and patient-centred clinical trials to benefit patients with this condition.

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Competing interests DLP has received honoraria for continuing medical education lectures in functional neurological disorder and is on the editorial board of Epilepsy & Behavior. MH is an inventor of patents held by National Institutes of Health (NIH) for an immunotoxin for the treatment of focal movement disorders and the H-2c for magnetic stimulation; in relation to the latter, he has received license fee payments from the NIH (from Brainways). He is on the medical advisory boards of Cala Health and Brainways. He has research grants from Allergan for studies of methods to inject botulinum toxins, Medtronic, Inc. for a study of deep brain stimulation (DBS) for dystonia and Cala Health for studies of a device to suppress tremor. WCL receives editor’s royalties from the publication of Gates and Rowan’s Nonepileptic Seizures, 3rd edition (Cambridge University Press, 2010) and 4th edition (2018) and author’s royalties for Taking Control of Your Seizures: Workbook and Therapist Guide (Oxford University Press, 2015) and receives research support from the Department of Defense (DoD/W81XWH-17-0169).

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Supplementary References


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# Supplementary Table 1. Other notable cohort treatment studies in motor functional neurological disorder.

<table>
<thead>
<tr>
<th>Study</th>
<th>n</th>
<th>Description</th>
<th>Key Points</th>
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</thead>
<tbody>
<tr>
<td><strong>Multidisciplinary rehabilitation – cohort studies</strong></td>
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<tr>
<td>Theuer et al 2020 [S43]</td>
<td>129</td>
<td>Retrospective review of patients admitted to a rehabilitation unit over a 20-year period. 185 patients were identified, 129 received treatment (multidisciplinary rehabilitation).</td>
<td>Treatment involved physiotherapy, occupational activities, psychiatric, and psychological support. After treatment, 70% of patients improved (36.2% with complete remission). Younger patients and those with an acute onset had a better outcome.</td>
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<td></td>
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<td><strong>Duration and setting:</strong> length of inpatient stay not reported</td>
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<tr>
<td></td>
<td></td>
<td><strong>Outcome measures:</strong> Improvement defined as remission of symptoms, marked improvement (75%) or moderate improvement (50%)</td>
<td></td>
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<tr>
<td>Hebert et al 2020 [S35]</td>
<td>20</td>
<td>Prospective cohort study of multidisciplinary rehabilitation based on the MoRe protocol in patients with functional movement disorders.</td>
<td>17 of 20 patients completed inpatient rehabilitation. 93% of patients completing treatment rated themselves as much improved. While not statistically significant at 1-year follow-up, patient-rated improvement was noted in 10 of 13 patients with available date.</td>
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<tr>
<td></td>
<td></td>
<td><strong>Duration and setting:</strong> mean 7.5 inpatient days</td>
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<td></td>
<td></td>
<td><strong>Outcome measures:</strong> CGI-severity; MAS; FGA; BBS; TUG; FIST; FIM</td>
<td></td>
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<tr>
<td>Kozlowska et al. 2020 [S49]</td>
<td>57, 60, 25</td>
<td>Three prospective cohort studies of pediatric multidisciplinary rehabilitation.</td>
<td>67%, 53%, and 80% of children in the 3 cohorts had mFND. Treatment included physiotherapy, psychotherapy (individual and family), attendance at hospital school,</td>
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<tr>
<td></td>
<td></td>
<td><strong>Duration and setting:</strong> 1-3 weeks, inpatient</td>
<td></td>
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<tr>
<td>Study</td>
<td>Sample Size</td>
<td>Study Design</td>
<td>Outcome Measures</td>
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<tr>
<td>Butz et al 2019 [S48]</td>
<td>100</td>
<td>Prospective cohort study of pediatric multidisciplinary rehabilitation.</td>
<td>GAF, resolution of FND, return to school and reintegration to home school post discharge.</td>
</tr>
<tr>
<td>Jimenez et al 2019 [S37]</td>
<td>63</td>
<td>Retrospective review of patients with functional motor symptoms participating in an interdisciplinary chronic pain rehabilitation program over a 4-year period.</td>
<td>WeeFIM</td>
</tr>
<tr>
<td>Jacob et al 2018 [S36]</td>
<td>32</td>
<td>Retrospective cohort study of specialist multidisciplinary rehabilitation.</td>
<td></td>
</tr>
</tbody>
</table>

Outcome measures: GAF, resolution of FND, return to school.
<table>
<thead>
<tr>
<th>Study</th>
<th>Duration and Setting</th>
<th>Outcome Measures</th>
<th>Improvement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bolger et al 2018 [S47]</td>
<td>Duration and setting: 8.4 ± 4.2 days, inpatient</td>
<td>Outcome measures: WeeFIM</td>
<td>25/30 children had mFND as part of their clinical presentation. Treatment included physiotherapy, occupational therapy, recreational, and music therapy, and psychological support. WeeFIM score change of 30 ± 11.9 (P &lt;.001), maintained at 3 months.</td>
</tr>
<tr>
<td>Demartini et al 2014 [S33]</td>
<td>Duration and setting: 4-weeks, inpatient</td>
<td>Outcome measures: CGI-patient rated; Health of the Nation Outcome Scale; COPM; PHQ-15; The Common Neurological Symptom Questionnaire</td>
<td>Together with Saifee et al (2012) below, the outcomes of a 4-week multidisciplinary rehabilitation program for patients with chronic mFND symptoms are reported. Significant but modest improvements were seen in a range of assessments post treatment and at 12-month follow-up (55% retention at 12 months). This included two-thirds of individuals rating their general health as better or much better at discharge; similar though slightly less positive gains were reported at 12-months.</td>
</tr>
<tr>
<td>McCormack et al 2014 [S40]</td>
<td>Duration and setting: 101-day median length of stay, inpatient</td>
<td></td>
<td>Similar to Demartini et al (2014) above, the outcomes of patients admitted for rehabilitation on a neuropsychiatric unit are presented. Outcomes are reported at discharge, with no follow up data. As with Demartini</td>
</tr>
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</table>

87% of patients reported improvement at the end of treatment. At 6-month follow-up, this reduced to 69%. Improvements occurred despite the long average symptom duration of 7.4 years.
Outcome measures: qualitative mobility and ADL performance, MRS et al (2014), improvements are seen despite long symptom durations and complex psychiatric comorbidity.

<table>
<thead>
<tr>
<th>Study Authors</th>
<th>N</th>
<th>Study Design</th>
<th>Duration and Setting</th>
<th>Outcome Measures</th>
</tr>
</thead>
<tbody>
<tr>
<td>Czarnecki et al 2012 [S32]</td>
<td>60</td>
<td>Retrospective cohort study of a multidisciplinary rehabilitation vs. treatment-as-usual controls.</td>
<td>Duration and setting: 5 consecutive days, intensive outpatient</td>
<td>Outcome measures: physician-rated improvement; patient-rated improvement (5-pt Likert) scale at 25 months post-treatment</td>
</tr>
<tr>
<td>Saifee et al 2012 [S42]</td>
<td>26</td>
<td>Retrospective cohort study of multidisciplinary rehabilitation.</td>
<td>Duration and setting: 24-day median length of stay, inpatient</td>
<td>Outcome measures: WSAS, time bothered by symptoms via visual analogue scale</td>
</tr>
</tbody>
</table>

**Physiotherapy - cohort studies**

<table>
<thead>
<tr>
<th>Study Authors</th>
<th>N</th>
<th>Study Design</th>
<th>Duration and Setting</th>
<th>Outcome Measures</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maggio et al 2020 [S38]</td>
<td>50</td>
<td>Retrospective cohort study of outpatient physiotherapy in consecutive patients.</td>
<td>Duration and setting: 6-12, 60-minute outpatient sessions</td>
<td>This study found that physiotherapy delivered in an outpatient setting, in a less intensive manner than previous rehabilitation studies, has the potential to benefit patients with mFND.</td>
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<tr>
<td>Study</td>
<td>Reference</td>
<td>Duration and setting</td>
<td>Setting</td>
<td>Outcome measures</td>
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<tr>
<td>Demartini et al 2020 [S34]</td>
<td></td>
<td>24 sessions, including 21 weekly tele-sessions</td>
<td>Telehealth</td>
<td>PMDRS, SF-36, CGI-patient rated</td>
</tr>
<tr>
<td>Matthews et al 2016 [S39]</td>
<td></td>
<td>18 days (mean length of stay), inpatient</td>
<td>Inpatient hospital</td>
<td>MRIMI</td>
</tr>
<tr>
<td>Nielsen 2015 [S41]</td>
<td></td>
<td>5-day, intensive outpatient</td>
<td>Outpatient</td>
<td>SF-36, EQ-5D-5L, WSAS, BBS, 10-meter walk test</td>
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<tr>
<td>Cognitive behavioural therapy – cohort studies</td>
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<tr>
<td>O'Connell et al 2020 [S46]</td>
<td></td>
<td>Retrospective review of CBT for mFND.</td>
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<tr>
<td>Study</td>
<td>Duration and Setting</td>
<td>Outcome Measures</td>
<td>Findings</td>
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<tr>
<td>Espay et al 2019&lt;sup&gt;26&lt;/sup&gt;</td>
<td>12-15 sessions, outpatient</td>
<td>3-point scale of improvement based on clinical note review, CORE-OM</td>
<td>Functional tremor severity improved significantly after 12 weeks of CBT. The improvement was associated with changes in the anterior cingulate / paracingulate activity on fMRI.</td>
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<tr>
<td>Kompoliti et al 2014 [S45]</td>
<td>Prospective unblinded CBT for mFND.</td>
<td>PMDRS</td>
<td>While patients in both groups improved in terms of CGI-severity scores with time, there were no statistically significant group-level differences at 3-months.</td>
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<tr>
<td>Hubschmid et al 2015 [S44]</td>
<td>Randomized study of interdisciplinary psychotherapeutic intervention (psychodynamic interpersonal treatment) vs. standard care.</td>
<td>PMDRS</td>
<td>Outcome assessments occurred at 2, 6 and 12-months post intervention initiation. SDQ-20 and CGI scores showed statistically significant group x time effects favouring the group receiving the interdisciplinary psychotherapeutic intervention vs. standard care.</td>
<td></td>
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<tr>
<td>Outcome measures: SDQ-20, CGI, MRS, healthcare utilization, SF-36</td>
<td>The intervention group also showed reduced inpatient hospital use compared to standard care. There were, however, no treatment group differences in terms of reported quality of life.</td>
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</table>

*the study included 174 participants, of which only 98 had a diagnosis of mFND. Additionally, outcome measures listed in this table focus on the selected physical functioning and quality of life instruments used in each study. Abbreviations: ADL, activities of daily living; BBS, Berg Balance Scale; CBT, Cognitive Behavioural Therapy; CGI, Clinical Global Improvement scale; COPM, Canadian Occupational Performance Measure; CORE-OM, Clinical Outcomes in Routine Evaluation-Outcome Measure; FGA, Functional Gait Assessment; FIM, Functional Independence Measures; FIST, Function in Sitting Test; GAF, Global Assessment of Function; MAS, motor assessment scale; mFND, motor functional neurologic disorder; MRIMI, modified Rivermead Mobility Index, MRS, modified Rankin scale; PHQ-15, Patient Health Questionnaire 15; PHQ-9, Patient Health Questionnaire 9; PMDRS, Psychogenic Movement Disorder Rating Scale; RCT, randomized controlled trial; SF-36, Short Form Health Survey – 36; SDQ-20, Somatoform Dissociation Questionnaire – 20; TUG, Timed Up and Go; WeeFIM, Functional Independence Measure for Children; WSAS, Work and Social Adjustment Scale.