Review

The adult multidisciplinary respiratory neuromuscular clinic

Many neuromuscular disorders (NMD) are complicated by respiratory failure. These patients are best managed in a multidisciplinary outpatient clinic to provide timely access to the various disciplines they require. The key mainstay of treatment of respiratory failure in patients with NMD is noninvasive ventilation, supported by secretion clearance, speech and language therapy, optimisation of nutrition and the maintenance of mobility. Patients with specific conditions may also require cardiology, neurology, orthopaedics, urology and psychological services. The respiratory NMD multidisciplinary team should also provide access to palliative care, and caregiver health and wellbeing should also be reviewed at clinical reviews. The future of care for the respiratory NMD patient will increasingly involve home services and telehealth and the clinic should be equipped and resourced to deliver these. Although not all health systems will be able to provide all elements of the multidisciplinary team discussed here, this review provides the “ideal” recipe for the adult multidisciplinary team and the evidence base underpinning this from which a clinic can be developed.

Educational aims

- To provide an overview of the care of an adult neuromuscular disorder patient presenting to the multidisciplinary respiratory clinic.
- To provide the evidence base for establishing the different elements of the multidisciplinary respiratory clinic.

Introduction

Neuromuscular disorders (NMD) include a broad range of neurological diseases that can affect the respiratory system (figure 1). They can include rapidly progressive conditions such as amyotrophic lateral sclerosis (ALS) as well as more slowly progressive or stable conditions such as the inherited muscular dystrophies (table 1). Many forms of NMD can lead to the development of respiratory failure, which...
The adult multidisciplinary respiratory neuromuscular clinic contributes to significant impairment of quality of life, morbidity and is often the cause of death [1]. With improving management, the life expectancy of these patients continues to rise, and with increased survival, the prevalence of patients with respiratory morbidity is likely to increase [2]. Monitoring and optimising the respiratory function in these patients is therefore of utmost importance to prolong life and reduce suffering.

The management of NMD patients suffering from respiratory failure is best conducted in a multidisciplinary team (MDT) context, and several disease-specific guidelines, statements and consensus reports recommend that NMD patients should be managed in clinics encompassing all members of the MDT [3–5]. The MDT care model fosters collaborative working across multiple medical specialities and the allied health professions, allowing a patient with multiple complex issues to be supported in a coordinated manner, rather than attending multiple outpatient visits to see different specialists. In fact, studies have demonstrated that not only is the MDT model preferable for patients [6, 7], but it also results in improved clinical outcomes when compared to management in a traditional isolated department model [8–10]. A recent publication on patients’ experiences of a newly established MDT clinic for ALS reported on the benefits of having patients’ care centralised and the confidence that they are seeing clinicians who specialise in their specific disease area. Although some challenges were highlighted (such as the extended travel to a centralised clinic), these were not insurmountable, and the patients in this cohort were generally quite supportive of a transition to the MDT model [11]. This paper will review the make-up of the state-of-the-art multidisciplinary respiratory NMD clinic, the rationale for this make-up and logistical issues that should be considered when developing this model of care.

**The multidisciplinary respiratory clinic**

The MDT clinic should consist of a core group of professions that almost all NMD patients will require support from, accompanied by support from other professions who provide care to patients with certain conditions.

**Table 1** Classification of NMD that most commonly affect the respiratory system

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<thead>
<tr>
<th>Rapidly progressive</th>
<th>Variable progression</th>
<th>Slowly progressive or stable</th>
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<tbody>
<tr>
<td>Amyotrophic lateral sclerosis</td>
<td>Duchenne muscular dystrophy</td>
<td>Spinal muscular atrophy</td>
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<td>Limb girdle muscular dystrophy</td>
<td>Myotonic dystrophy</td>
<td>Post-polio syndrome</td>
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<td>Myofibrillar myopathy</td>
<td>Facio-scapulohumeral muscular dystrophy</td>
<td>Becker muscular dystrophy</td>
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<td>Danon disease</td>
<td>Kennedy disease</td>
<td>Inclusion body myositis</td>
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<tr>
<td>Mitochondrial myopathies</td>
<td>Myasthenia gravis</td>
<td>McArdle disease</td>
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conditions or those with specific issues (figure 2). The link with community practitioners and services such as palliative care is crucial, and is likely to foster longer periods out of hospital and for the patient to remain at home for as long as possible. The best example of an MDT model of care is arguably in oncology. Oncology MDTs invariably involve an MDT coordinator, responsible for bridging the communication gap between the service provider and the patients [12]. A respiratory NMD MDT would likely benefit from a centralised coordinator to ensure patients are seeing the appropriate services, and only those that actually need to be seen. They would also have a crucial role in managing the contact between patients at home and the clinic, to assure that all remain informed of developments in the patients' care. An alternative approach would be to establish a lead physician and a case manager for each patient, who would ensure that the patient can access the necessary services they need at that time point in the course of their illness [13].

Patients may be referred into the adult NMD respiratory clinic from a number of sources. Neurologists frequently make the diagnosis of neuromuscular conditions and lead on the management of their peripheral nerve dysfunction, as well as administration of disease-modifying agents. Their input in terms of medical management and prognosis is essential. In many cases, they remain the primary physician managing the care of the patient but are often not co-located with the respiratory clinic; however, usually both neurologists and respiratory physicians jointly manage these patients. Paediatricians will manage all cases of congenital NMD until the time of transition to adult care. Both neurologists and paediatricians will refer to the adult NMD respiratory clinic when their patient exhibits features of respiratory failure for investigation and ongoing management. Some patients will be referred as incident cases of hypercapnic respiratory failure, or respiratory muscle dysfunction of unknown origin for investigation. Others may be referred as part of pregnancy planning [14] or pre-operative assessment [15], if they are considered to be at risk of respiratory muscle weakness during the perioperative period. Any patient referred in to the adult NMD clinic will benefit from a core set of baseline investigations (table 2) [16].

Patients referred from paediatric services must receive specific consideration for the transition to adult care. Barriers to successful transition to adult care include access to specialist adult providers of neuromuscular respiratory care [17], a break in the bond with their long-term paediatric care team [18].

Figure 2 Elements of the neuromuscular respiratory MDT. The inner ring illustrates the specialities that are likely to be physically present in the neuromuscular clinic and considered core to the respiratory neuromuscular MDT, while the outer ring displays specialities that will be consulted and will reach-in to services dependent on specific disease processes.
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and changes to financial remuneration policies from paediatrics to adult care. In particular, children receiving ventilation are likely to require a large input from the MDT to ensure a smooth transition to adult care [19]. The US Department of Health and Human Service’s Maternal and Child Health Bureau’s six core elements of healthcare transition might be a useful model when designing a pathway for paediatric neuromuscular patients to transition to adult services [20].

**Ventilation**

A large proportion of NMD patients will require home mechanical ventilation (HMV) at some point during the course of their illness. The initiation and monitoring of HMV are key roles of the respiratory NMD MDT. Key issues that need to be addressed when organising the MDT include: when long-term ventilation should be started, where it should be initiated, and how it should be monitored. An additional consideration is the assessment of the need to convert to invasive ventilation via tracheostomy.

Patients should be monitored for signs and symptoms of respiratory weakness, prior to initiating HMV. The primary indications for initiating HMV are: 1) respiratory muscle weakness, 2) overnight hypoventilation, and/or 3) symptoms of respiratory muscle weakness during the daytime and/or at night (sleep-disordered breathing) [21]. Daytime hypercapnia (>45 mmHg) is a sign of advanced respiratory muscle weakness and patients referred with this should be treated as urgent cases for initiation of HMV. Indirect

**Table 2 Assessments in the adult multidisciplinary respiratory neuromuscular clinic**

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<tr>
<th>History</th>
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<td></td>
<td>Immersion dyspnoea</td>
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<td></td>
<td>Daytime somnolence</td>
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<td></td>
<td>Impaired cognitive function</td>
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<td></td>
<td>Aspiration pneumonia</td>
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<td>Physical examination</td>
<td>Use of accessory respiratory muscles</td>
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<td>Paradoxical breathing</td>
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<td></td>
<td>Tachypnoea (rapid shallow breathing)</td>
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<td>Gas exchange assessment</td>
<td>Pulse oximetry</td>
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<td>Arterial/capillary blood gases</td>
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<td></td>
<td>Transcutaneous capnometry</td>
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<td>Radiological</td>
<td>Chest radiograph</td>
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<td>Diaphragmatic ultrasound</td>
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<td>Questionnaires</td>
<td><em>Epworth Sleepiness Score (ESS)</em></td>
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<td></td>
<td><em>Sleep-Disordered Breathing in Neuromuscular Disease Questionnaire (SiNQ-5)</em></td>
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<td></td>
<td><em>Severe Respiratory Insufficiency (SRI)</em></td>
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<tr>
<td>Respiratory muscle function assessment</td>
<td>Spirometry (seated/supine)</td>
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<td></td>
<td>Slow vital capacity</td>
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<td></td>
<td>Peak cough flow</td>
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<td></td>
<td>Maximal mouth inspiratory and expiratory pressure (MIP and MEP)</td>
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<td></td>
<td>Sniff nasal inspiratory pressure (SNIP)</td>
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<tr>
<td>Advanced respiratory muscle function assessment</td>
<td>Invasive respiratory muscle assessment <em>(Sniff P&lt;sub&gt;oes&lt;/sub&gt; and P&lt;sub&gt;a&lt;/sub&gt;)</em></td>
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<td></td>
<td>Phrenic nerve stimulation <em>(twitch P&lt;sub&gt;a&lt;/sub&gt;)</em></td>
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<td></td>
<td>Phrenic nerve conduction studies</td>
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<td>Nocturnal respiratory function</td>
<td>Overnight pulse oximetry</td>
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<td>Transcutaneous capnometry (monitoring)</td>
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<td>Respiratory polygraphy</td>
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<td>Polysomnography</td>
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Investigations that may provide additional information, but may not be available at all centres are listed in italics.
causes of respiratory muscle weakness, such as congestive cardiac failure, should also be optimised prior to initiation of HMV. Although strong evidence supporting the initiation of HMV prior to the onset of daytime hypercapnia is yet to be published [22, 23], and its impact is likely to be different in different disease groups [24, 25], there are increasing reports of its benefit [26, 27], including prolonging tracheostomy-free survival [28]. Conversely, the presence of bulbar symptoms has been proposed as a reason to delay initiation of noninvasive ventilation (NIV) until the respiratory failure has worsened [29]. There are a number of national and international guidelines that advise on the initiation of HMV in specific disease groups [27, 30–33].

To investigate for evidence of respiratory muscle weakness, the MDT must have access to a range of investigations (table 2). While respiratory muscle strength is formally assessed using invasive measures of diaphragm weakness, these investigations are invasive, highly specialised and not widely available. There is, therefore, a need for surrogate measures of diaphragmatic weakness. The MDT should be able to conduct spirometry (seated and supine), as well as sniff nasal inspiratory pressure (SNIP), maximal inspiratory (MIP) and maximal expiratory (MEP) pressure. To investigate for daytime hypercapnia and nocturnal hypoventilation, the MDT must have the ability to assess daytime blood gases and organise overnight studies, including overnight oximetry and polysomnography [34]. Screening tools, such as the SiNQ-5 questionnaire can be used during clinic to identify sleep-disordered breathing [35]. Although the ESS is widely used to monitor daytime sleepiness in patients with sleep-disordered breathing, it has low predictive value in patients with neuromuscular conditions [35, 36]. Important disease-specific questionnaires may also be performed in the respiratory NMD clinic. For example, the Amyotrophic Lateral Sclerosis Functional Rating Scale-Revised (ALS-FRS-R) is a useful tool in monitoring the decline in respiratory function in patients with ALS [37].

Traditionally, patients are set-up on NIV during a multi-night inpatient stay, where the NIV is titrated according to polysomnography results [38]. Set-up in outpatient settings has resulted in equivalent adherence to therapy as nocturnal set-up [39] and can avoid the need for admission to hospital. With the advent of auto-titrating devices and telemedicine, there is increasing interest in the validity of remote set-up of NIV. Auto-titrating devices result in similar outcomes to standard devices [40] and home-initiation of NIV is both feasible [41] and noninferior to inpatient-initiation [42]. As devices are improved and the impact of artificial intelligence arrives in respiratory medicine, we are likely to see increasing evidence for the home set-up of NIV [43]. This will add a logistical telemedicine burden to a respiratory NMD MDT, with skilled telemedicine practitioners required to facilitate treatment. It is also important to note that even with remote titration and set-up of the device, an in-person encounter would still be required to determine the optimal interface and provide training on the use of the ventilator; however, this could be done at the patient’s home.

Monitoring patients on NIV is important to ensure good adherence and effective treatment. Frequency of monitoring has to be balanced between frequent reviews at the time of initiation, to achieve optimal care, and the need for a disabled individual to come into hospital repeatedly. Once the slowly progressive NMD patient is on stable treatment, annual reviews with daytime gases and nocturnal oximetry are sufficient [44]. Rapidly progressive NMD patients are likely to need more frequent reviews, and with progressive disease, these may require home assessments, emphasising the need for a comprehensive home assessment team as part of the NMD clinic.

The interface used with NIV has considerable impact on outcome. Although the majority of studies investigating the impact of interface have focused on patients with acute respiratory failure [45], two randomised studies in patients with chronic hypercapnic respiratory failure have demonstrated that interface can impact on tolerance to therapy as well as physiological markers of therapy success [46, 47]. A recent systematic review on NIV in ALS patients has demonstrated the importance of a patient-tailored approach to selection and fitting of the interface to optimise delivery of NIV [48]. Specialists such as respiratory nurses, physiologists and technicians with experience of ventilation are an essential element of the respiratory NMD MDT. They are responsible for ensuring the correct interface is selected to suit the patient’s needs and troubleshoot issues with the interface. They will also deliver education and training to patients and carers on applying the NIV mask and using the ventilator. In addition, they would be able to support the detailed interrogation of the data download from the ventilator, which can provide information on adherence, leak, tidal volume and breathing pattern. This information can help optimise treatment at clinic reviews [49, 50], and going forward, will allow remote assessment of ventilation efficacy [51].

Some patients receiving NIV will benefit from conversion to invasive ventilation via tracheostomy [52]. This may include patients with copious secretions despite optimal medical therapy or patients unable to tolerate or achieve satisfactory ventilation. Despite its invasive nature, patients have reported improved breathlessness and fatigue, and better speech and oral intake following tracheostomy [53]. Thus, the MDT will need access to physicians with skills to manage long-term tracheostomised patients.
**Secretion management**

Patients with NMD often suffer from an ineffective cough [54]. This can be due to damage to the innervation of the cough reflex (any of the vagus, phrenic, intercostal or recurrent laryngeal nerves), weakness of inspiratory or expiratory respiratory muscles, inability to close the glottis due to bulbar dysfunction, or any combination of these. Ineffective cough results in inadequate secretion clearance, which can lead to respiratory infections, morbidity and death [55–57]. It is therefore important to support these patients with adequate secretion clearance techniques. Indeed, secretion management is a priority for NMD patients [58].

The presence of a respiratory physiotherapist specialising in neuromuscular diseases in the respiratory NMD MDT would be the gold standard, although it is acknowledged that this may not always be possible in all healthcare systems. Physiotherapy interventions can be divided into proximal and peripheral airway clearance techniques, and different conditions benefit from different techniques. At a minimum, the respiratory NMD MDT should be staffed and equipped to be able to offer manual assisted coughs, mechanical insufflation–exsufflation, lung volume recruitment, glossopharyngeal breathing, and chest wall percussion and oscillation [59]. It is important to note that while it may seem intuitively useful [60] and does not cause any harm [61], there is no convincing evidence for the use of respiratory muscle training to improve clinical outcomes or quality of life in NMD patients [62, 63].

Sialorrhoea is also common in certain NMDs, such as ALS and any condition causing bulbar dysfunction or facial muscle weakness [64, 65]. Management is important as it can cause skin maceration, psychological difficulties and contribute to intolerance to NIV [66]. Treatment ranges from conservative management, medical therapy with anticholinergics, to surgical therapy. In patients who are refractory to medical management, injection of botulinum toxin into the salivary gland is considered a safe and effective alternative [67–69], with a positive impact on quality of life [70]. Botulinum toxin injections can be safely administered in an outpatient setting, and so practitioners with this skillset would be a valuable addition to the respiratory NMD MDT. In fact, a UK survey of practice of sialorrhoea management identified that the main barrier to the use of botulinum toxin was the availability of trained staff [65]. If botulinum toxin is unsuccessful, contraindicated or unavailable, radiotherapy of the salivary gland is also an option. The literature includes reports of its use in ALS, where it is shown to be successful in effecting an improvement in symptoms for 4–6 months [71, 72]. There appears to be a preference for electron-based therapy compared to photon-based therapy [73]. Surgical options include ligation, denervation, ablation or resection of the salivary glands, but these procedures require a pulmonary reserve, making it unavailable to most NMD patients and infrequently utilised [21].

**Cardiovascular care**

Patients with NMD frequently suffer from cardiac disease [74]. The same pathophysiological process causing respiratory muscle weakness can also affect the cardiac myocytes. The prevalence and pattern of cardiac disease varies between different conditions. Some patients will suffer from a dilated cardiomyopathy (e.g. Duchenne muscular dystrophy, Becker muscular dystrophy, limb girdle muscular dystrophy), while others suffer from arrhythmias (e.g. myotonic dystrophy), which can lead to sudden cardiac death [75]. Input from cardiac services for both monitoring and treatment is therefore crucial. Any patient reporting cardiac symptoms, and all patients suffering from conditions that are associated with cardiac dysfunction (the hereditary muscular dystrophies, myotonic dystrophy, mitochondrial disorders and congenital myopathies [75, 76], should be screened regularly with electrocardiograms for arrhythmias [77] and echocardiograms for ventricular dysfunction [76]. It is important to note that the value of echocardiogram decreases with progressive scoliosis and obesity, and so the availability of cardiac magnetic resonance imaging [78] or real-time three-dimensional echocardiography [79] is an important consideration. The availability of a dedicated cardiologist with expertise in NMD in the respiratory NMD MDT would be the gold standard, so that patients can be evaluated and reviewed at the same time. It would also facilitate access to more advanced cardiac interventions such as implantable devices, should the need arise.

**Speech and language therapy**

There is a high prevalence of dysarthria and dysphagia in patients with NMD, stemming from respiratory muscle pump, airway muscle and bulbar dysfunction [80]. Although they may be obvious during clinical assessment, the presence of dysphagia correlates well with the presence of dysarthria [81], and so it is important to screen for both. Dysphagia can have a significant impact on nutrition [82] and there is an increased risk of aspiration pneumonia, which can be life-threatening [83, 84]. Dysarthria is likely to have a significant impact on an already decreased quality of life [85]. The input of speech and language therapists can have an important impact on the care of these patients. Although the quality of evidence for speech interventions in NMD patients is weak [86], there is no evidence of harm from speech interventions and so it remains a useful
service to feature in the respiratory NMD MDT. In progressive conditions, patients’ speech is likely to deteriorate to a point where verbal communication is not possible. Ensuring access to augmented communication therapies for these individuals is vital to optimise quality of life and health [87, 88]. In addition, different modes of ventilation impact on speech quality and so close coordination with the respiratory physician is important to ensure that speech is optimised with evolving ventilatory therapies [89].

**Nutrition**

Both undernutrition and overnutrition are common issues in the care of patients with NMD. Adequate management of nutrition results in improved quality of life for both patients and their families [3, 90]. Decreased muscle strength causes slow chewing and an increased time to consume food, resulting in reduced oral intake and undernutrition. Patients frequently suffer from delayed gastric emptying (due to gastrointestinal muscle hypotonia), gastrointestinal reflux and constipation [91], which leads to abdominal discomfort and reduced oral intake. In addition, the use of ventilators can lead to swallowing of air, contributing to abdominal discomfort [3]. Overnutrition is a consequence of normal oral intake with reduced caloric need and physical activity or excessive oral intake secondary to appetite-increasing medications, and caregiver compassion preventing caloric restriction [91]. In addition, the introduction of artificial feeding via gastrostomy can result in iatrogenic overnutrition, if not carefully assessed and guided by specialist diéticians. Resulting obesity has implications for longer term cardiovascular and metabolic health, as well as a direct impact on respiratory physiology [92]. It is therefore imperative that the management of NMD patients includes input from dietetic services, ideally in the same clinic location, so that advice can be provided when issues are identified. A study of clinicians caring for ALS patients has highlighted that nutrition can also be impacted by psychological factors, such as distress around the time of diagnosis, the need to maintain control over their own nutritional intake and the loss of social interaction from mealtimes [93]. Clinicians should be mindful of these factors and facilitate access to psychological interventions to support the patient to accept their new nutritional needs and accessibility.

Undernutrition, particularly when secondary to impaired swallowing, can be effectively managed with gastrostomy, inserted surgically, percutaneously or radiologically. This has been demonstrated to reduce the risk of respiratory infections and hospitalisations [94]. Timing of gastrostomy is key; if implemented before the onset of severe respiratory failure, the potential complications of the procedure itself can be minimised [95]. Conversely, there may be psychological issues around maintenance of independence without gastrostomy-nutrition to be considered and so close communication between the gastroenterology team, dietetics and the respiratory team are required to provide optimal nutritional support to the patient.

**Physical and occupational therapy, and rehabilitation**

In NMDs there is an element of muscle degradation and so there is long-standing interest in the value of strength-building and rehabilitative interventions on improving quality of life, slowing progression and reducing mortality. Reviews of the existing literature on ALS [96, 97] and muscle disorders in general [98] were unable to make conclusive recommendations about the use of physical therapy and rehabilitation interventions. This is primarily due to the heterogeneity of the studies that have been conducted and poor study design. Importantly, no adverse effects of physical therapy have been identified. Furthermore, a synthesis of qualitative data revealed that ALS patients feel rehabilitation interventions result in an improvement in quality of life [99]. As there do not appear to be any negative sequelae, and there are perceived patient-orientated benefits, the MDT should aspire to be able to offer physical therapy to NMD patients. It is acknowledged that in some healthcare systems, the lack of conclusive evidence may result in a difficulty in funding allocation towards this. Physical capability may be augmented with other interventions such as mobility aids, orthotics and abdominal binders. In progressive conditions, patients’ mobility may eventually deteriorate such that independent mobilisation is not possible. A dedicated wheelchair service is crucial at this point, particularly if electric powered wheelchairs can be provided, as this will help to maintain some level of independence and reduce the burden on caregivers.

**End-of-life considerations**

Palliative care input can provide positive support to patients with NMD, such as the perception of control over their illness, management of troublesome symptoms and future planning [100–102]. Qualitative assessments of patients [103] and bereaved relatives [104] have reported that patients would welcome the input of palliative services, and that otherwise death can involve unmanaged respiratory and psychological symptoms. Despite this, the provision of palliative services is not widespread. In younger people in particular, there appears to be a reluctance from both caregivers and clinicians to initiate conversations about end-of-life care with the...
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The evidence base to support the inclusion of palliative care is sparse [107, 108], restricting access to resources. This is partly because, unlike in oncology or COPD, “end-stage” NMD has not been categorically defined [109], leading to clinical challenges about when to introduce palliative care. The perception that the introduction of palliative care leads to cessation of life-prolonging interventions also acts as a barrier, and future models of NMD care should consider palliative care and advanced planning as complementary to life-prolonging therapies, rather than dichotomous [109]. A retrospective review of patients who died within a year of diagnosis of NMD reported that older age at diagnosis, early-onset breathlessness, dysphagia, constipation and impaired mobility are all significantly associated with early mortality [110], and these criteria could serve as indicators for the introduction of palliative services to an individual’s care. Recently, a prognostic model to identify ALS patients at risk of death within 6 months has been published and may be of use in other NMDs [111]. The first report of a joint palliative and NMD service in the UK demonstrated that patients were receptive to this, but that challenges remain around the timing of introducing this service over a patient’s disease course [112].

Psychological support

The presence of psychological disturbance in patients with NMD is well established across many conditions [113–118]. Given that mental health conditions are associated with poorer health outcomes [119], it is likely that they have an important impact on outcomes in NMD patients too. A systematic review on adults with NMD was unable to provide conclusive support for the provision of psychological interventions to improve quality of life; however, this was primarily due to a lack of high-quality evidence [120]. The studies they did identify all reported short-term benefits on quality of life following the introduction of psychological interventions. Indeed, an audit of patient satisfaction after the introduction of a clinical psychologist to a neuromuscular clinic reported not only that patients were supportive and found this useful but also that quantitative measures of wellbeing and satisfaction improved [121]. The provision of psychological support within the respiratory NMD MDT is therefore likely to be beneficial and an important factor in patient-orientated outcomes. There is no evidence-based specialised approach for NMD, particularly because of the varying nature of different NMDs, and so it is important that this provision is staffed by psychologists specialised in the care of NMD patients, who can use their experience to provide a specialised service for this unique cohort [122].

Outreach team and telehealth

The respiratory NMD MDT is often located at a regional centre, providing care to individuals who come from a large area. Travel to the clinic can be troublesome for the patient, particularly given their mobility limitations and ventilation requirements [11]. A home service, visiting patients in their home, can offset these challenges. A home care team can deliver various functions including clinical review, ventilator servicing, regular tracheostomy tube change, and training of caregivers on equipment use. The MDT should therefore include individuals able to travel to patients’ homes or care facilities with the necessary skills-set and equipment to deliver these functions. A recent qualitative study has highlighted the importance of the support a home care team provides, particularly in patients requiring long-term tracheostomy ventilation [123]. The logistics that underpin this service should not be underestimated and financial remuneration for this service will likely require a bespoke system, as “ad hoc” costs are more likely. With centralised care and the increasing survival of NMD patients, the demands on this service are likely to burgeon over the coming years.

The incorporation of telehealth into the MDT model is a potential mutually convenient solution to the travel burden on patients to a centralised clinic and the resource burden of a home service on the MDT. To date, investigations have focused on using telehealth to manage patients ventilated at home. Studies have demonstrated that remote monitoring and adjustment of ventilator settings is feasible [124] and acceptable to patients [125]. When comparing telehealth to usual care, patients receiving telehealth appear to suffer from less emergency room visits and hospitalisations [126, 127] and a lower risk of disease progression [128], although this may reflect a patient selection bias, as those deemed suitable for telehealth may be a healthier cohort. Most of the literature supporting the use of telehealth in NMD reports on cohorts of patients who have been successfully supported via telehealth, without a “usual care” comparator [129–131]. Although not able to demonstrate superiority or equivalence with usual care, they do provide useful descriptions of how telehealth can be implemented in this patient population. An American telehealth service for ALS reported a significant cost reduction compared with standard care [132]; although this will be different in individual health systems, it is likely that once a telehealth service is established, the per-patient cost will be lower by avoiding visits to hospital and possibly reducing the number of investigations. The true value of telehealth is difficult to ascertain without randomised studies; a recent study reported that the feasibility of conducting a randomised trial about telehealth in NMD patients is low [133]. In addition, a review of the evidence supporting the use of telehealth in respiratory illnesses, including NMD, was unable to provide any conclusive findings about its utility [134], even though it appears to...
be acceptable to patients [135]. Nevertheless, as technology advances and patient preferences for interaction with healthcare professionals evolve, particularly with the development of app-based consultations, telehealth is likely to become an important feature of any respiratory NMD MDT.

**Burden on caregivers**

Studies in various conditions have reported a significant impact on caregivers’ mental health, with a high prevalence of anxiety and depression symptoms [136, 137]. These are often related to: the amount of time spent providing care [136, 138]; the severity of the condition, as well as the emotional health of the patient [139, 140]; isolation due to care commitments [138, 141]; and physical difficulties such as the introduction of a wheelchair [142]. The introduction of HMV adds further burden on caregivers [143], including negatively impacting on carers’ sleep quality [144]. Caregivers have reported that they experience a feeling of loss while the patient is still alive, as well as after they have passed away [145]. Psychological support specialised for caregivers of NMD patients would be a useful addition to the MDT to provide interventions when caregivers accompany the patient to clinic. In addition to the impact on caregiver quality of life, the financial burden of caring for NMD patients on families must also be considered [146, 147]. When providing an holistic approach to these patients, it is important to consider the impact on their carers and suggest options to relieve the burden, such as respite care, further education about the care they are providing and signposting to financial assistance, where available. The need for input from social services in the multidisciplinary NMD clinic is clear.

**Summary**

The optimal care of a patient with NMD suffering from respiratory failure involves all members of the MDT; the provision of care from each member of the MDT is ideally delivered at the same time in an outpatient clinic where each specialty is co-located. The future of the respiratory NMD MDT will probably rely heavily on telemedicine and home services, and the logistical burden these bring must be considered. Although it is recognised that not all healthcare systems will be able to deliver all of these specialties, and that the requirement will be dependent upon the disease make-up of the clinic population, this review serves as a recipe for the “ideal” multidisciplinary respiratory NMD clinic, from which specific requirements for specific situations can be selected.

**Key points**

- Patients with neuromuscular disorders who suffer from respiratory failure require complex care that is best delivered in a multidisciplinary context.

- The core disciplines that most patients will require and that should be available at every clinic include respiratory physicians, physiotherapy, occupational therapy, dietetics, and speech and language therapy.

- The future of the care of the respiratory neuromuscular patient will increasingly involve home services and telehealth, and their burden need to be considered when establishing a service.

- Provision should also be provided for palliative care and support for caregivers.

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**Self-evaluation questions**

1. In which of the following settings can NIV be initiated in patients with NMD?
   - a. inpatient admission with polysomnography
   - b. outpatient clinic with capnometry
   - c. home with remote monitoring

2. Which of the following factors have a negative impact on caregiver quality of life?
   - a. severity of the condition
   - b. emotional health of the patient
   - c. introduction of home mechanical ventilation
   - d. introduction of a wheelchair

3. What cardiovascular complications commonly occur in patients with NMD?
   - a. cardiomyopathy
   - b. ischaemic heart disease
   - c. arrhythmias
   - d. valvular disorders

4. Which of the following investigations are not part of the assessment of a respiratory NMD patient?
   - a. respiratory muscle testing
   - b. overnight oximetry
   - c. cough assessment
   - d. cardiopulmonary exercise testing
Suggested answers
1. a–c.
2. a–d.
3. a and c.
4. d.

References


1. Davidson ZE, Truby H. A review of nutrition in Duchenne

2. Ramelli GP, Aloysius A, King C, et al. The role of physical therapy and


