Assessment and management of respiratory function in patients with Duchenne muscular dystrophy: current and emerging options

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Abstract: Duchenne muscular dystrophy (DMD) is an X-linked myopathy resulting in progressive weakness and wasting of all the striated muscles including the respiratory muscles. The consequences are loss of ambulation before teenage years, cardiac involvement and breathing difficulties, the main cause of death. A cure for DMD is not currently available. In the last decades the survival of patients with DMD has improved because the natural history of the disease can be changed thanks to a more comprehensive therapeutic approach. This comprises interventions targeted to the manifestations and complications of the disease, particularly in the respiratory care. These include: 1) pharmacological intervention, namely corticosteroids and idebenone that significantly reduce the decline of spirometric parameters; 2) rehabilitative intervention, namely lung volume recruitment techniques that help prevent atelectasis and slow the rate of decline of pulmonary function; 3) scoliosis treatment, namely steroid therapy that is used to reduce muscle inflammation/regeneration and prolong ambulation in order to delay the onset of scoliosis, being an additional contribution to the restrictive lung pattern; 4) cough assisted devices that improve airway clearance thus reducing the risk of pulmonary infections; and 5) non-invasive mechanical ventilation that is essential to treat nocturnal hypoventilation, sleep disordered breathing, and ultimately respiratory failure. Without any intervention death occurs within the first 2 decades, however, thanks to this multidisciplinary therapeutic approach life expectancy of a newborn with DMD nowadays can be significantly prolonged up to his fourth decade. This review is aimed at providing state-of-the-art methods and techniques for the assessment and management of respiratory function in DMD patients.

Keywords: DMD, spirometry, respiratory muscles, cough device, NIV, steroids, idebenone, respiratory function, lung volume recruitment, scoliosis, spinal fusion, mdx, GRMD

Introduction

Duchenne muscular dystrophy (DMD) is an X-linked myopathy that affects one out of 3,600–6,000 male births.¹-³ It is caused by mutations in the dystrophin gene leading to a defect in the protein dystrophin which results in progressive weakness and wasting of all the striated muscles. At the beginning, the proximal lower limb muscles are affected followed by the shoulder muscles, by the distal limb muscle, and ultimately by the respiratory muscles. The consequences are progressive impairment of motor skills, loss of ambulation before teenage years, orthopedic complications, cardiac involvement, and breathing difficulties. Without any intervention, death occurs around the age of 19 due to cardiac and respiratory failure.⁴-⁷ At the moment therapy is not available, but in the last decades the survival of these patients improved and a newborn with DMD now has a life expectancy into the fourth decade.³,⁸-¹⁰ The natural history of the disease can now be changed thanks to a more comprehensive therapeutic approach.
comprising interventions targeted to the manifestations and complications of the disease, particularly in respiratory care.\textsuperscript{7,11–14} ACE-inhibitions are used to prevent or improve cardiomyopathy. Corticosteroids are used to reduce muscle inflammation, minimize later complications, and prolong ambulation.\textsuperscript{3,15,16} When the patient becomes non-ambulatory, treatments aim to reduce the risk of progressive scoliosis and stabilizing pulmonary function. According to the internationally recognized guidelines, a structured approach to respiratory management of DMD during the progression of the disease comprises: 1) the monitoring of respiratory function to deal with respiratory complications timely; 2) lung volume recruitment (LVR) techniques to open the lung and reduce atelectasis; 3) the use of a cough assisted device to improve airway clearance and to reduce the risk of pulmonary infections; and 4) the use of non-invasive ventilation (NIV) to deal with nocturnal hypoventilation, sleep disordered breathing, and ultimately respiratory failure.\textsuperscript{8,11,13,14}

The genetic therapy represents hope for the future to develop a definitive cure for DMD. Many promising researches on skeletal muscles are currently under development although not specifically addressed to respiratory muscles.\textsuperscript{17,18}

The present review article is thought to provide a state-of-the-art methods and techniques for the assessment and management of respiratory function in DMD patients. In particular, the techniques currently used for respiratory functional evaluation are illustrated, indicating limits, advantages, and usefulness for assessing the effects of different possible kinds of intervention.

Assessment of respiratory function

The causes of death in DMD patients are both respiratory and cardiac, with the former being the main cause until the 1980s to 1990s. More recently, the structured approach to respiratory problems and, in particular, the institution of nocturnal NIV, had a major effect on survival. Since respiratory failure, however, is still among the main causes of death in patients affected by DMD,\textsuperscript{3,5–7,12} it is extremely important to measure lung and respiratory muscle function in order to monitor the progression of the disease, identify early signs of respiratory failure, and therefore plan optimal interventions to improve the quality of life and longevity in these patients. The assessment of respiratory function is also important to quantify the possible effects of innovative therapeutic trials.

According to their invasiveness and to the collaboration required to the patients, lung and respiratory muscle function tests are classified as non-invasive/non-volitional, non-invasive/volitional, invasive/non-volitional, and invasive/volitional. These tests can be “specific” or “global” according to whether they allow to distinguish between the contribution and the relative involvement of the different respiratory muscles, namely the diaphragm, the intercostal muscles, the abdominal muscles, and the accessory inspiratory muscles. These tests can assess: i) the level of muscle activation by measuring the electrical activity of the muscle (electromyography [EMG]); ii) the force developed by the muscles by measuring the pressure developed; iii) the resulting action of the contraction of the muscles by measuring flow and/or volume; or iv) the morphology of the muscles with different imaging techniques.\textsuperscript{19}

Non-invasive/non-volitional tests

The electrical activity of each respiratory muscle can be non-invasively measured by transcutaneous EMG using surface electrodes. Unfortunately, at rest the signal/noise ratio is very poor, particularly in weak muscles, because of the “cross-talk” of other muscles, including cardiac activity.\textsuperscript{20,21}

The ventilatory pattern during quiet breathing at rest can be measured at the mouth on a breath-by-breath base by a pneumotachograph preferable using a mask rather than a mouthpiece in order to avoid possible leakage due to weakness of cheek muscle and macroglossia, which are common characteristics in the advanced stages of the disease. The face mask connection to the lung function device is a valid alternative; however, it is important to choose the appropriate prediction equations, depending on subject-instrument interface.\textsuperscript{22} From the integration of the flow signal, tidal volume ($V_T$) and breathing frequency ($f_R$) are measured and minute ventilation ($V_E = V_T \times f_R$) and rapid and shallow breathing index ($RSBi = f_R/V_T$) derived.\textsuperscript{23} These are global respiratory indexes that can be performed at any age and in different postures: standing, seated, and supine.

Measurements of thoraco-abdominal motion do not require the use of mouthpiece or mask and can be divided into three groups:

- those recording the anteroposterior variations of chest wall diameters, such as the magnetometers (ie, coils placed anteriorly and posteriorly on the ribcage and the abdomen);\textsuperscript{24}
- those measuring variations of ribcage and abdominal cross-sectional areas, such as respiratory inductive plethysmography (ie, coils sewn in belts or wearable garments placed below the axillary line and above the umbilicus);\textsuperscript{19,25}
those measuring total and compartmental chest wall volumes, such as opto-electronic plethysmography that detects the 3D displacement of a large set of reflective markers placed on the thoraco-abdominal surface of the subject.26–29

Although the shape of the diaphragm can be reconstructed with an excellent resolution by using magnetic resonance imaging (MRI), its clinical use is still limited because of the high cost. Moreover, patients are not always able to maintain the supine posture for the prolonged time required for the acquisition.30

On the other hand, ultrasound (US) imaging is increasingly used since it allows to non-invasively study the diaphragm at rest from three different views: 1) the displacement of the diaphragm dome by positioning the US probe on the anterior abdominal wall; 2) the length of apposition of the diaphragm by positioning the US probe on the lateral ribcage; and 3) the thickness of the diaphragm by positioning the US probe in the intercostal space around the eighth rib.31–33

Non-invasive/volitional tests

All non-invasive/volitional tests require repetition of maximal efforts to get reliable results that can fatigue the weak muscles of DMD patients. They also require full cooperation and are difficult for children to perform. Cooperative children over the age of 7 are considered able to perform reproducible and therefore reliable maneuvers.

Maximal static inspiratory (MIP) and expiratory (MEP) pressures measured at the mouth provide a global index of inspiratory and expiratory muscle strength, respectively. Being MIP and MEP influenced by respiratory muscle length that in turn depends on lung volume, it is crucially important to control the lung volume at which the maneuver is performed. MIP has to be performed at residual volume (RV) and MEP at total lung capacity (TLC), respectively when inspiratory and expiratory muscles are maximally stretched.19,34

Sniff nasal inspiratory pressure (SNIP) measured at the nostril is generally easier to perform, since the requested maneuvers is more natural. If performed correctly (ie, with an outward movement of the belly), it can be considered a specific index of the strength of the diaphragm, although in presence of nasal obstruction it underestimates the real pressure.19,35,36

Inspiratory muscle fatigue can be assessed by the tension time index (TTI) that is calculated as \( \frac{P_{DI}}{MIP} \times \text{duty cycle} \), where \( P_{DI} \) is the pressure measured during the first 100 milliseconds of occlusion while breathing quietly and the duty cycle is the ratio between inspiratory time and the total time duration of the breath. High values of TTI indicate that inspiratory muscles are at risk of fatigue.37–40

Peak expiratory flow (PEF) and cough peak flow (CPF) are measured at the mouth respectively during a maximal exhalation and cough. They provide information on the expiratory muscles.41

Traditional spirometric parameters such as forced vital capacity (FVC) and forced expiratory volume in 1 second (FEV₁) are global indexes of the respiratory function. Values above 80% of their ratio (FEV₁/FVC), the Tiffenau index, can indicate the presence of restrictive lung disease. The diagnosis has to be confirmed by reduced values of TLC. A drop in FVC (>20%), when the patient goes from sitting to supine position, is a more specific sign of diaphragmatic weakness.42

TLC, functional residual capacity, and RV are absolute lung volumes that cannot be measured by simple spirometry. They require either nitrogen washout technique or body plethysmography. Wheel-chair bound DMD patients can meet great difficulties in entering a body plethysmograph.19,43–46

Invasive/non-volitional tests

Invasive/non-volitional tests imply the use of esophageal and/or gastric unpleasant balloon-catheter systems. They provide specific indexes of the different respiratory muscles. The use of the catheters is not recommended in very severe DMD patients with difficulties in swallowing to avoid regurgitation and aspiration.19 An additional caution of pressure measurements by esophageal and gastric balloons is represented by the possible bradycardia mediated by the stimulation of the vagal nerve. They can be used during quiet breathing at rest without requiring any maneuvers.47

The measurement of esophageal pressure (PGES), based on transducer-catheter-balloon system placed in the lower third of the esophagus, is considered a useful surrogate of pleural pressure while abdominal gastric pressure (PGA) is measured by placing the transducer-catheter-balloon system in the stomach. The trans-diaphragmatic pressure (PDI) is calculated as \( \frac{P_{GA}}{P_{GES}} \) and represents the force generated by the diaphragm.19 The ratio between swings of PGA and PDI (\( \Delta P_{GA} / \Delta P_{DI} \)) during spontaneous quiet breathing is able to provide an assessment of paradoxical breathing.48

Peak PDI during magnetic stimulation of the phrenic nerve tests the strength of the diaphragm in non-cooperative patients.49

Volumetric computed tomography imaging can be used to obtain 3D reconstruction of the diaphragm with an excellent
resolution, but the radiation exposure limits the clinical use of this technique in DMD patients.60

Invasive/volitional tests
Invasive/volitional tests include the measurements of $P_{\text{OES}}$, $P_{\text{GA}}$, and $P_{\text{Di}}$ during maximal maneuvers like sniff and cough.51,52 Maximal $P_{\text{OES}}$ and $P_{\text{Di}}$ during sniff maneuvers represent reliable indexes of the strength of global inspiratory muscles and of the diaphragm, respectively;36,53,54 while $P_{\text{GA}}$ during maximal cough is an index of the strength of expiratory muscles.51,55 Because cough efficacy depends on the inspiratory volume preceding the maneuver, lung volume is a source of variability and should be controlled.56

Table 1 summarizes the different techniques, categorized into invasive/non-invasive, and volitional/non-volitional, which are used to evaluate different ventilatory functional parameters in DMD patients, as reported in the literature.

Table 1 Assessment of ventilatory function in DMD

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**Abbreviations:** DMD, Duchenne muscular dystrophy; $P_{\text{OES}}$, esophageal pressure; $P_{\text{GA}}$, abdominal gastric pressure; $P_{\text{DI}}$, trans-diaphragmatic pressure.

Respiratory function in DMD
Respiratory function assessed by non-invasive non-volitional tests
During spontaneous quiet breathing at rest in supine position, $V_E$ decreases because of the reduction of $V_T$ with preserved respiratory rate, particularly in older DMD patients.57 In seated position $V_E$ does not change significantly because respiratory rate increases to compensate for the fall of $V_T$.37,58,59

The abdominal contribution to $V_E$ in supine position during inspiration is an index of the action of the diaphragm. It progressively decreases with age and it is an important predictor of nocturnal hypoexemia57,60 and inefficient cough.61

Quantitative muscle US is a non-invasive child-friendly tool able to measure DMD muscle changes in terms of both echo intensity and muscle thickness.62 DMD patients are characterized by poor spirometry, reduced inspiratory
muscle force (lower MIP), and higher diaphragmatic thickness, measured at its appositional zone by B-mode ultrasonography, compared to healthy controls. This may indicate pseudo-hypertrophy of the diaphragm secondary to infiltration of connective tissue and fat deposition as occurs in limb muscles.

Recent studies investigating MRI measurements of T2 relaxation time (MRI-T2) on lower limb muscles in patients and on canine model confirm the feasibility of MRI to objectively and non-invasively measure muscle pathology in DMD. MRI-T2 of the sternocleidomastoid muscle of DMD patients are significantly lower than an age-matched control group indicating altered tissue composition. MRI, therefore, represents a potentially promising technique for early detection of disease progression not only of limb muscles but also of the diaphragm and other respiratory muscles in DMD. It can also be used to assess the efficacy of therapeutic interventions on the muscles.

**Respiratory function assessed by non-invasive volitional tests**

The absolute value of FVC follows a pathognomonic pattern characterized by an ascending phase, a peak approximately at 12–14 years, and a plateau followed by a descending phase. The absolute value reached at the plateau correlates with life expectancy in these patients. Values of FVC lower than 2.1 L are associated to lower CPF and represent important thresholds to start with techniques that improve airway clearance. FVC below 1 L is a suitable indicator to consider starting more intensive patient support, being a good predictor of severe oxygen desaturation, uncontrolled hypercapnia, and impending death within 12 months. The decline in FVC has recently been shown to be associated with ventilation inhomogeneity assessed by lung clearance index measured with the multiple-breath washout technique. The maximal percentage postural difference of FVC in DMD patients is approximately 9%. It tends to decrease with age like MIP values. This can indicate that a global weakness of both inspiratory and expiratory muscles is not able to compensate for the diaphragmatic impairment.

In DMD, FVC expressed as percentage of the predicted values (FVC%) starts approximately 80% at the age of 10 and then it linearly declines with age as shown by different authors indicating a progressive respiratory muscles weakness. FVC% and its decline are prognostic markers in DMD.

Inspiratory flow reserve volume is defined as the ratio between the maximal inspiratory flow during tidal breathing and during FVC. This ratio is higher in DMD compared to controls indicating a reduction in inspiratory flow reserve in DMD.

When listed as a function of age, the absolute values of FEV1, functional residual capacity, TLC, RV, inspiratory and expiratory reserve volumes reveal ascending, plateau, and descending phases like FVC. Between the ages of 8 and 12 the restrictive pattern is mainly due to reduced expiratory reserve volumes and increased RV. This can indicate an early deterioration of expiratory muscles. After the age of 16, the decline of lung volumes is associated with reduced inspiratory reserve volumes and TLC and therefore with inspiratory muscles’ impairment.  

MIP and MEP in Duchenne patients are reduced compared to normal values and they decrease with age, with MEP being globally lower than MIP. Expiratory muscles, therefore, seem to be more compromised while inspiratory muscles seem to be relatively preserved. MIP predicts the extent of hypercapnia and starts to decline from the age of 10 years onward. Because FVC decay starts a bit later, MEP maneuver seems to be affected earlier than spirometry. The mean decline with age of MIP varies from 4% to 6.9%, 59,70,72

MEP is associated with cough efficiency: if MEP is higher than 60 cm H2O, DMD patient is expected to have an adequate cough; while MEP below 45 cm H2O is an indicator to start cough-assisting techniques.

SNIP increases until the age of 10.5 and decreases thereafter with a mean decline of 4.8 cm H2O (5.2% predicted) per year. Because it declines earlier than FVC, it can be considered an early sign of inspiratory muscle strength. Reliable values of SNIP can be obtained from the age of 5 and therefore its measure should be initiated early in the course of the disease.

With the progression of DMD, fatigue index (TTI) increases and consequently, without any intervention, older patients are likely to approach respiratory muscle fatigue.

**Respiratory function assessed by invasive non-volitional tests**

As indicated earlier, \( \Delta P_{GA}/\Delta P_{DI} \) ratio measured at rest during spontaneous quiet breathing is a non-volitional marker that reflects the relative contribution of the diaphragm to \( V_T \). In teenagers with DMD, \( \Delta P_{GA}/\Delta P_{DI} \) is negative indicating that the diaphragm is weaker than the other inspiratory muscles and it paradoxically moves upward. \( P_{DI} \) secondary to magnetic stimulation of the phrenic nerve is lower in
DMD patients than patients with spinal muscle atrophy and congenital myopathy indicating that Duchenne patients are characterized by the weakest diaphragm.31

Respiratory function assessed by invasive volitional tests

Trans-diaphragmatic and P_{OES} during sniff are low and progressively decrease after the age of 10 years. The evolution of gastric pressure during cough follows a similar decrement pattern, being significantly reduced by the age of 8 years.51,59

Figure 1 shows a schematic diagram summarizing the natural course of the different respiratory functional parameters over time according to what has been reported in the up-to-date literature.

Pharmacological intervention

There is no current curative treatment for DMD and the only medications available and used are glucocorticoid steroids.3,88 The main effects of corticosteroids on skeletal muscles are inflammation reduction, mass increment and deterioration delay. Steroid therapy delays the motor impairment of DMD children by improving muscular strength leading to prolonged ambulation for up to 2–3 years longer.16,89–95 At the beginning, steroid therapy was prescribed to improve motor function. Successively some authors suggested prolonging the treatment also after patients are wheelchair bound for its potential effect in preventing scoliosis and respiratory deterioration. There are studies showing no effect of steroid therapy on the respiratory function in treated DMD patients.30,83 A stabilization of FVC and FEV\textsubscript{1} after 2 years of treatment has been described86 while another study reports a significant increase of MEP and cough peak flow (CPF) after treatment without FVC improvement/stabilization.97 Many authors have found the effectiveness of steroid treatment in improving spirometry in their cohort of DMD patients.98–100 Two reasons can explain such a discrepancy in the role of steroid therapy on lung function in DMD: the lack of standardization in steroid administration and/or the binary subdivision (steroid vs non-steroid) according to the use of therapy. The non-steroid group includes not only young patients, but also old patients who never took the therapy (because its introduction in the clinical management of DMD is relatively new) together with patients who chronically took steroids but in a discontinuous way. As already pointed out by Mayer et al100 the authors believe that the heterogeneity of the two groups can introduce a bias effect on the results since old and therefore more compromised patients can worsen the overall lung function of the group, while the discontinuous-users can contribute in a positive way thanks to the possible beneficial effect of the therapy. The cooperative international neuromuscular research group Duchenne natural history study15,16 groups 340 DMD patients according to age and to the use of steroids as follows: 1) naïve patients: never used steroids or treated <1 month total and not currently under medication; 2) current recipients; and 3) past users: treated in the past for >1 month and not currently under medication. An overall improved lung function in terms of FVC, FEV\textsubscript{1}, PEF, MIP, and MEP has been found in the current steroid users across the age groups 10–12 and 13–15 years old.13,16 This demonstrates how crucial it is to properly subdivide patients while testing the efficacy of new therapy in order to avoid possible bias due to the patients’ heterogeneity in their clinical conditions because of the progressive nature of the disease.

Since the use of corticosteroids has well-known side effects, such as weight gain, cataract, osteopenia, behavioral changes, increased insulin resistance, and delayed puberty, there is a high medical need for new effective, safe, and well-tolerated pharmacological therapy able to slow the impairment or even ameliorate the respiratory function in DMD. In a pre-clinical placebo-controlled study in the mdx mouse idebenone,
a potent antioxidant shows cardio-protective effect and improvement in exercise performance. A 12-month Phase 2 randomized placebo-controlled trial on a small group of DMD patients shows a significant treatment effect not only on cardiac markers, but also on PEF, as absolute values and as percentage of the predicted values. The beneficial effect of idebenone on PEF is more pronounced in naïve steroid patients rather than in current users, suggesting an inhibitory effect of glucocorticoid steroids. Finally, the trial reaches Phase 3 by studying the efficacy of idebenone on respiratory function in DMD patients not on concomitant steroid therapy. Idebenone significantly reduces the decline of different parameters of the respiratory function, namely PEF, FVC, and FEV, and results have shown it to be safe and well tolerated by patients therefore representing a new therapeutic approach for DMD. The goal for future pharmacological therapy is to have at least equal efficacy than glucocorticosteroids but with lower side effect profile. Multiple new treatment approaches have emerged ranging from molecular based therapies aimed to reduce muscle fibrosis or to promote muscle blood flow. It seems more likely that a combination of different approaches, targeting more than one pathway, may be more satisfactory. Alternative approaches including mutation-specific therapy (gene replacement or repair and stem cell therapy) represent the future possibility to find a definite cure for DMD. There have been genetic-based treatment experiments in the last few years in clinical trials, ie, ataluren for non-sense mutation in the dystrophin gene and antisense oligonucleotide induced exon skipping. The promising results of some of these studies still have to face the lack of accurate clinical and biological outcome measures able to discriminate between and describe modifications in the disease history. The ongoing clinical trials are not specifically addressed to study new therapeutic effects on respiratory muscles and consequently do not have respiratory function measures as primary outcomes. The following trials consider respiratory variables as secondary outcome in addition to other clinical and functional measures: a) evaluating the long-term safety of ACE-031 in subjects with DMD; b) long-term safety trial of ataluren to treat Becker and Duchenne muscular dystrophies; c) NCT02286947; d) NCT00264888; e) NCT00759876; f) NCT01009294; g) NCT01239758; h) NCT01099761; and i) NCT01462292.

Respiratory muscle training (RMT) and physiotherapy intervention

Respiratory muscles are morphologically and functionally skeletal muscles and therefore RMT can be a possible treatment to improve their strength and endurance and to try to counteract the progressive weakness induced by DMD. The role of RMT in neuromuscular patients is still controversial being potentially dangerous since it may accelerate muscle fatigue by overwork. Beneficial effects of RMT have been shown in DMD. These comprise increased strength of expiratory muscle, improved muscle endurance, and augmented MIP. Another study demonstrates that inspiratory muscle training (IMT) through resistive breathing and/or efforts against a variable inspiratory resistance allows diaphragmatic training in terms of strength and endurance. Peak esophageal and PDI during sniff maneuvers have been measured before and after 6 months of training. In ten out of 15 patients, respiratory muscle function parameters improved and the effect has remained even 6 months after the end of training. RMT is not effective in patients with FVC lower than 25% predicted and/or PaCO₂ higher than 45 mmHg. It seems that RMT is not applicable in the most compromised patients, maybe because they do not have enough functional muscle mass left or because their respiratory muscles are so impaired and thus more susceptible to damage.

IMT is strongly recommended preoperatively in case of spinal surgery in order to prevent pulmonary complication secondary to anesthesia. IMT improves the functionality of respiratory muscles by increasing FVC to 26.2% before surgery and by reducing postoperative risks. RMT can be simply implemented within video games with the possibility to control all the training processes. Improvements in respiratory performance correlate positively with the actual amount of training and negatively with baseline lung function test. This implies that RMT seems to be more effective in DMD patients who have moderate respiratory impairment.

The main limitations of these studies are the reduced number of patients and the lack of standardized protocols. RMT therapeutic effects rely on two factors: the amount of training and the relatively good condition of the patient.

LVR, also known as “breath stacking”, is a method to inflate and recruit the lung. It consists of an inspiratory pressure applied to the airways, the hold of the delivered volume of air by glottic closure, followed by a spontaneous or assisted forced expiratory maneuver. In DMD patients, the progressive impairment of the ventilatory pump reduces the range of motion of the chest wall and therefore the ribcage becomes stiff with reduced compliance of the chest wall and of the lung. The main consequence is the formation of areas of atelectasis. These lead to reduced oxygenation of the blood and to impaired CO₂ elimination due to the imbalance between alveolar ventilation and blood perfusion.
of the capillary. LVR helps to prevent atelectasis, to limit contractures of the thoracic muscles, to maintain the motion of the chest wall particularly of the costo-vertebral joints, and to improve lung compliance. It can be used also in uncooperative and in very weak patients who cannot hold a mouthpiece and therefore it is applicable to DMD patients. LVR has been shown to slow the rate of decline of pulmonary function compared to the decline before the start. The values of FVC and peak cough flow improve after LVR. Because LVR can reach higher pressure compared to in-exsufflator, great care should be taken to avoid barotrauma. Failure to successfully stack the breath may occur in very sick patients whose $V_t$ is lower than the dead space of the mask and whose inspiratory effort is not enough to trigger the valve. In such a case, it is recommendable to use a supported breath stacking device in which a resuscitator bag helps to augment $V_t$ and to open the valve.

Glossopharyngeal breathing (GPB) represents an autonomous lung-inflation maneuver that these patients should be taught, practiced and encouraged particularly in the last stage of the disease. GPB has been proven to increase lung volume by inflating one gulp at a time and to delay the onset of daytime ventilator use. It can be performed in the event of ventilator failure and during the inspiratory cough phase, since cough flow correlates with pre-cough volume.

Recently, a 10-month program of yoga breathing exercises has been shown to ameliorate the pulmonary function in DMD patients in terms of significant improvement of FVC and FEV1.

Finally, very severe DMD patients use a spontaneous movement called the “row-a-boat phenomenon”, ie, swing of the torso back and front rhythmically synchronized with $f_r$ to compensate for the atrophied respiratory muscles. By simply allowing the row-a-boat movement, FVC increases up to 50.8% compared to the values reached when the movement is prevented.

**Management of scoliosis and spinal fusion**

In the natural history of DMD, the development of scoliosis is a frequent complication that impacts on respiratory function, seating, and comfort. Ninety percent of untreated DMD boys develop scoliosis, secondary to muscle weakness, pelvic imbalance, and after becoming wheelchair bound full-time. For this reason, prolonging ambulation in DMD patients is one of the main strategies to delay the onset of scoliosis. Other strategies include physiotherapy (passive stretching) to maintain lower limb range of motion, night splints or serial casting. Moreover, surgery can release contractures of lower limbs and then rehabilitation with “KAFO” (knee ankle foot orthoses) may prolong standing and walking for 2 years. There is level III-2 evidence that the use of corticosteroids delays the onset of scoliosis by extending independent ambulation. Steroid therapy has been shown to reduce the need for surgical treatment to reduce scoliosis in DMD from 90% to 15%.

Long-term corticosteroid treatment, however, carries important side effects including worsened bone fragility and the risk of vertebral fractures.

When the scoliosis curvature exceeds the 40° of Cobb angle, scoliosis rapidly progresses and may impair breathing by reducing lung capacity on both the convex and the concave sides. Since spinal orthoses seem to have minimal effects on preventing, delaying or minimally correcting the curve progression, surgical correction becomes the indication. Because of the cardiopulmonary impairment of DMD, spinal surgery is better when performed during early adolescence in order to avoid the high risk of developing postoperative pneumonia, prolonging ventilatory dependence, cardiac failure, and eventually death. In the literature, there are conflicting reports about the effect of surgical spine stabilization on pulmonary function. They all consider the evolution of FVC. There are studies reporting positive effects of spinal fusion in terms of stabilization of FVC or slightly slowing FVC deterioration with respect to a control non-operative group or to pre-operative patients. It should be noted that in these papers there was a continuous reduction in FVC after surgery, although to a lesser extent. Other studies have found no ameliorative effects on the decline of pulmonary function.

The decision to have surgical fixation of the spine therefore has to be considered mainly to improve the general well-being of the patient (maintenance of comfortable sitting balance, release of pain due to muscle contractures, and improvement of quality of life) rather than to improve his respiratory function which will continue to decline anyway. Nevertheless, in DMD patients the combination of spinal fusion with nocturnal ventilation prolongs median survival to 30 years.

**Cough assisted devices**

The main causes that require intubation for DMD patients are episodes of pneumonia and acute respiratory failure secondary to benign upper-respiratory infection. This is a consequence of retained secretion due to inability to cough effectively. In DMD, cough is considered adequate when CPF >270 L/min and ineffective if CPF <160 L/min and therefore cough augmentation is required. When CPF is
between 160 and 270 L/min cough is considered effective when the patient is in good and stable condition, but it becomes inadequate in case of viral illness when secretions increase.¹¹,¹³,¹⁴ Once assisted coughing is recommended, it should be used not only when the patient is ill, but once or twice-per-day as maintenance therapy to improve airway clearance and reduce atelectasis. Since CPF strongly depends on the volume of air inspired pre-coughing,⁵⁶ the first goal of cough assisted techniques is to maximally inflate the lungs. This can be achieved autonomously by GPB, manually through the use of an inflating Ambu bag or mechanically assisted. The expulsive cough phase can be spontaneous, when the patient is still able to generate sufficient intrathoracic pressures, manually assisted by abdominal compression or mechanically forced by a cough assisted device. The combination of inspiratory volume support with manual assisted expiratory cough phase is more efficacious than the single intervention per se.¹⁴⁵,¹⁴⁶ The use of mechanical in-exsufflator is effective in preventing pulmonary complications like pneumonia and respiratory failure and in decreasing the number of hospitalizations due to respiratory illness.⁴ The addition of intrapulmonary percussive ventilation to assist mucus clearance techniques is a safe method to augment the efficacy of these methods to keep the airways of tracheotomized DMD patients clear.¹⁴⁷ Care must be taken to adequately pre-set the timing and the levels of pressure in order to be well tolerated by the patient, to maximize cough effectiveness, and to avoid possible complications like pneumothorax, gastroesophageal reflux, and pulmonary hemorrhage.

CPF correlates with MIP and MEP⁹,¹⁴⁶ and all the three parameters improve after long-term steroid therapy. Therefore the steroid-induced preservation of both inspiratory and expiratory muscle strength translates to better cough efficiency.⁹⁷

The combination of cough assisted devices with NIV before surgery helps in easy extubation after surgery without postoperative respiratory complications.¹⁴⁸

NIV
The onset or chronic respiratory insufficiency is a fatal inevitable complication of DMD. Patients not treated with mechanical ventilation die by the end of the second decade of life.⁷,³⁷,⁷³,⁷⁷ The use of NIV combined with adequate airway clearance prolongs survival up to the third decade of life in DMD patients.⁷,⁹,¹⁰,⁷⁷ In these patients, nocturnal NIV helps to reduce the fatigue of respiratory muscles by lowering TTI. As a result dyspnea reduces, as indicated by lower Borg scale values. After nocturnal NIV, vital capacity slightly increases while MIP does not change. Two additional hours of diurnal NIV improves the respiratory condition of severe DMD patients who present with nocturnal hypoventilation symptoms like breathlessness in the evening. In these patients, the benefits of nocturnal NIV last 6 hours. The same levels of lower TTI and reduced dyspnea are restored and maintained after the 2 additional hours of diurnal NIV that represents a good tool to reduce the load on the respiratory muscles rising from morning to evening.²⁷ This is an important therapeutic intervention for DMD patients who are dyspneic during diurnal unassisted breathing.

Moreover, intermittent positive-pressure breathing (IPPB) shows positive effects on regional lung ventilation. A session of 30 consecutive breaths up to 30 cm H₂O performed in different positions (supine, left and right lateral position) increases the regional ventilation of the lungs. Such increment is maintained also 3 hours after the end of IPPB if the patient wears an abdominal belt. The supine position is characterized by the greatest improvement in lung ventilation particularly in the anterior regions. In order to reduce risks of hyperinflation in only one part of the lung, it is recommended to use several positions for the IPPB protocol.¹⁴⁹

With the progression of the clinical course of the disease, DMD patients develop respiratory failure characterized by daytime hypercapnia, hypoxia, and impairment in swallowing. The traditional approach to deal with this last stage of the disease would be to introduce tracheostomy ventilation.¹⁵⁰ Twenty-four hours’ NIV represents a safe and non-invasive alternative to tracheostomy.⁹,¹⁴³,¹⁵¹ The advantage of 24-hour NIV compared to invasive ventilation is the possibility to engage normal verbal communication, to eat, to breath-stack as desired, and to avoid the clinical complications of tracheotomized patients. Dysphagia also benefits from 24 hours of NIV maybe because oropharyngeal muscles improve their strength and endurance associated with the reduced work of breathing. The swallowing mechanism may improve and therefore it can support better nutrition.¹⁵²

Tracheostomy ventilation becomes necessary in case of severe aspiration risk, severe respiratory infection, and after onset of severe bulbar weakness that is rarely developed in patients with DMD. The combined use of NIV and mechanically assisted cough allows the decannulation and extubation even for unweanable DMD patients.⁹,¹⁵³

Animal studies
DMD animal models, both mice and canine, are important reliable models for a physiological study of the disease and/or to test new pharmacological treatment efficacy, particularly
for the diaphragm, the most important respiratory muscle. Exposure to episodic hypoxia mocks the sleep disorder breathing of DMD patients and it results in a 30% reduction in diaphragmatic strength in mdx mice.\textsuperscript{154} Fourteen weeks’ treatment with phosphodiesterase 5 inhibitors reduced the dysfunction of the diaphragm in mdx mice by enhancing the contractility of the diaphragm, by slowing muscle fibrosis, and by normalizing the expression of pro-inflammatory cytokine.\textsuperscript{155} Improvement in diaphragmatic health, in terms of muscle mass normalization and reduced fiber necrosis, can be obtained in neonatal mdx mice with the combination of antisense therapy with dystrophin restoration and myostatin inhibition.\textsuperscript{156} Reduction of the oxidative damage and of the inflammatory response in the mdx mouse diaphragm can be achieved using deferoxamine, a potent iron chelating agent.\textsuperscript{157} It is very important to restore diaphragmatic function not only from a respiratory point of view but also since the rescue of the diaphragm in mdx mice, by increasing its UTRN or dystrophin expression, prevented heart dysfunction. The decline in lung function secondary to respiratory muscles’ impairment increases pulmonary vascular pressure therefore leading to pulmonary hypertension.\textsuperscript{158} In DMD patients, pulmonary hypertension increases right atrial pressure with possible right ventricular failure. An early component of the hypercapnic ventilatory reflex has been revealed in mdx mice, demonstrating a malfunction in the peripheral chemosensory drive. This is supposed to be a further contributor to the respiratory failure in mdx mice. Therefore the monitoring of peripheral chemosensory drive function may be useful to better manage DMD patients with respiratory failure.\textsuperscript{159} The DMD mdx mouse model with haploinsufficiency of the UTRN gene is characterized by more severe diaphragmatic fibrosis and may improve when testing antifibrotic therapies, especially in the most severe stage.\textsuperscript{160}

Golden Retriever muscular dystrophy (GRMD) is a canine model of DMD. GRMD dogs guarantee an adequate level of ventilation at rest by increasing their respiratory rate in order to compensate for a lower $V_{\text{A}}$ compared to normal dogs. GRMD does not show abdominal paradox. When the respiratory drive is stimulated by doxapram, abdominal expiratory muscles are recruited early to compensate for the loss of diaphragmatic function. The diaphragm in GRMD is weaker, shorter, thicker, and severely fibrotic compared to the other respiratory and locomotor muscles.\textsuperscript{161}

**Conclusion**

Until now clinical trials have not considered respiratory function variables as primary outcomes. This, however, would be highly recommended, as respiratory problems are among the main causes of death in DMD patients. The non-invasive and non-volitional methods and techniques for respiratory functional assessment which are nowadays available should be considered as a tool not only for experimental trials, but also for clinical monitoring starting during the very early stages of the disease.

**Disclosure**

The authors have no conflict of interest to declare.

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