Health-related quality of life and functional changes in DMD: A 12-month longitudinal cohort study

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Abstract

In Duchenne muscular dystrophy (DMD) little has been reported on the association between clinical outcome measures and patient health-related quality of life (HRQOL) tools. Our study evaluated the relationship between 12 month changes on the Generic Core Scales (GCS), the Multidimensional Fatigue Scale and the Neuromuscular Module of the PedsQLTM with several outcome measures (6 minute walk test, North Star Ambulatory Assessment and timed items) in ambulatory DMD. Ninety-eight ambulatory DMD in a multicentric setting were included in the study. At baseline, the PedsQLTM inventories correlated with almost all the functional measures On the Child Self-Report there was a significant decrease between baseline and 12 months on the PedsQL TM GCS and its first domain, in parallel with the decrement in the functional outcome measures. Correlation between the 12 month changes on the PedsQLTM inventories and functional measures were almost all negligible. Similar results were obtained on the Parent Proxy-Report.

In conclusion, PedsQLTM correlates with the level of impairment at baseline, but this does not hold true when 12 month changes are considered. Further studies comparing different tools are needed to better elucidate the complexity of the relationship between HRQOL and functional performances.

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Keywords: Duchenne muscular dystrophy; Quality of life; Outcome measures; PedsQLTM

1. Introduction

In the last few years promising new therapies for Duchenne muscular dystrophy (DMD), including nonsense mutation suppression via stop codon read through and exon skipping via antisense oligonucleotides, have been introduced into registration-directed international multicentre investigational drug clinical trials. Safety and efficacy trials have involved both ambulatory and
nonambulatory subgroups of DMD populations [1–6], even though most of the recent and ongoing studies focus on the ambulatory phase [7–9]. The 6 minute walk test (6MWT) has been recently chosen as the primary outcome measure in both international multicentre investigational drug clinical trials and longitudinal natural history studies in DMD ambulant patients [7–13]. It provides an integrated global assessment of ambulatory function that is influenced by decreased lower extremity strength, biomechanical inefficiencies during gait, diminished endurance, and compromised cardio-respiratory status [14]. Functional scales, such as the North Star Ambulatory Assessment (NSAA), represent an ideal additional tool to the 6MWT, as they provide information on a wider spectrum of functions that reflect everyday life activities [15]. We have recently reported data from multicentric longitudinal studies in ambulant DMD boys, describing the changes on both 6MWT and NSAA and providing cut-off values to predict the risk of losing ambulation within 12, 24 and 36 months [10–12].

There has been an increasing pressure from regulatory authorities to include health-related quality of life (HRQOL) assessments in order to test the impact of possible functional changes on activities of daily living and more generally, on patient’s quality of life. The Paediatric Quality of Life Inventory with its Generic Core Scales (PedsQL™ GCS) is the most used tool to assess quality of life in clinical studies and a specific module for neuromuscular disorders, the PedsQL™ Neuromuscular Model (PedsQL™ NMM), has been validated in DMD and spinal muscular atrophy (SMA) [16,17].

The PedsQL™ Multidimensional Fatigue Scale (PedsQL™ MFS) was designed as a child self-report and parent proxy-report generic symptom-specific instrument to measure fatigue in pediatric patients [18] and has never been used in neuromuscular disorders.

A few studies have recently reported the use of these modules in DMD cohorts [19–21], but little has been reported on if and how the PedsQL™ scores change in relation to longitudinal changes on the functional measures commonly used in therapeutic trials and clinical practice, such as the 6MWT and NSAA [21].

The aim of our study was to assess if 12 month changes in function, assessed by 6MWT, NSAA, 10 meter timed walk/run and Gowers test, were associated to changes on HRQOL tools in a large cohort of ambulant DMD boys. In order to assess possible differences among the most commonly used HRQOL tools we used the PedsQL™ GCS, the PedsQL™ MFS and the PedsQL™ NMM, both in the Child Self-Report and Parent Proxy-Report.

2. Subjects and methods

The study is a prospective longitudinal multicentric cohort study involving patients recruited from 10 tertiary neuromuscular centers in Italy and followed for at least one year.

Patient inclusion criteria were:

(i) genetically proven DMD diagnosis, aged between 5 and 13 years. The age of 5 as lower limit was chosen as both 6MWT and NSAA have been validated from this age and the upper limit of 13 years as we wished to select a population similar to that included in most of the current clinical trials in DMD.

(ii) still ambulant and able to walk independently for at least 75 meters, and

(iii) without severe or moderate learning difficulties or behavioral problems, that could affect compliance or the level of performance.

All consecutive patients attending the 10 participating centers who fulfilled the inclusion criteria were enrolled in the study. As part of this study, all centers performed the NSAA followed by the 6MWT at each visit. As we wished to obtain the best compliance in the functional assessments also in younger patients, the PedsQL™ modules were filled in after the functional tasks at variance with the administration guideline suggestions. A research assistant assisted the younger children (5–7 years) in completing the questionnaires and was available to assist the older group if needed. Data were collected at baseline and at 12-month follow-up assessment.

This study is part of a longer longitudinal study on 6MWT and NSAA in DMD involving the same centers [10–12,22,23]. Details of the training for the participants and of the interobserver reliability for measures among the centers have already been reported in our previous studies [22,23].

2.1. Standard protocol approvals, registrations and patient consents

The study was approved by the ethical committee of each center. Written informed consent was obtained from participants.

2.1.1. PedsQL™

In order to assess the suitability of the different PedsQL™ modules in DMD, we used the Italian version not only of the general module (PedsQL™ GCS), but also of the dedicated module for neuromuscular disorders (PedsQL™ NMM) and of the PedsQL™ Multidimensional Fatigue Scale (MFS). We also separately analyzed the results of the first domain of each module (respectively: Physical Function Score, General Fatigue Scale, About My Neuromuscular Disease), as they are supposed to be more related to patient functional status [20].

The PedsQL™ is a modular instrument designed to measure HRQOL in children and adolescents aged 2–18 years. The 23−item PedsQL™ 4.0 GCS encompasses: (1) Physical Functioning (8 items), (2) Emotional Functioning (5 items), (3) Social Functioning (5 items) and (4) School Functioning (5 items). The 18-item PedsQL™ MFS was designed to measure fatigue in pediatric patients and comprises the General Fatigue Scale (6 items), Sleep/Rest Fatigue Scale (6 items), and Cognitive Fatigue Scale (6 items). The 25-item PedsQL™ 3.0 NMM encompasses 3 Scales: (1) About My/My Child’s Neuromuscular Disease (17 items), (2) Communication (3 items) and 3) About Our Family Resources (5 items). The PedsQL™ NMM young child form (5–7 years) does not contain the Communication and About Our Family Resources Scales.

All three inventories include parallel Child Self-Report and Parent Proxy-Report formats for children aged 5–18 years. The versions for patients between 5 and 7 years have 3-choice
answers, the other versions have 5 choices answers. The
instructions, Likert response scale, and scoring method for the
PedsQL™ modules are identical. For each item the score can
vary between 0 and 4. Items are linearly transformed to a 0 to
100 scale (0 = 100, 1 = 75, 2 = 50, 3 = 25 and 4 = 0), with
higher scores indicating better HRQOL.

2.1.2. 6MWT

6MWT was performed in all DMD ambulant boys older than
5 years using a modified version of the American Thoracic
Society guidelines for the test [24]. Modifications include the
addition of continuous encouragement from the testing staff,
and a “safety chaser” to walk along behind the subject during
testing. The test is generally completed within 15–20 minutes.
Suitability and interrater and intrarater reliability in DMD for
the 6MWT have already been reported [14,23,25].

2.1.3. NSAA

The scale consists of 17 items, ranging from standing (item
1) to running (item 17) and includes several items assessing
abilities that are necessary to remain functionally ambulant.
Each item can be scored on a 3 point scale using simple criteria:
2 – Normal, achieves goal without any assistance; 1 – Modified
method, but achieves goal independent of physical assistance
from another person; 0 – Unable to achieve independently.
The score can range from 0, if all the activities are failed, to 34, if all
the activities are achieved. The scale is generally completed in
a maximum of 15 minutes.

2.1.4. Timed items

The NSAA also includes the possibility to record timed
items (10 meter timed walk/run test and time to rise from
the floor or Gowers test) [22]. The time taken to complete these
tasks is not part of the score, but provides an additional measure
that can be monitored over time. In children who were or
became unable to perform these tasks, conventionally a time
equal to the worst performance in the group was subjectively
given to indicate poor performance [10]. Reliability of the
NSAA and timed items in the same multicentric setting has
already been tested with positive results [22].

2.1.5. Statistical analysis

Continuous variables are presented as mean ± standard
deviation (SD) and categorical variables as frequencies and
percentages. Paired t-tests were used for comparisons of the
measures across time (baseline and 12 months). Delta of the
measure evaluations were calculated as the difference between 12
month and baseline data. The relation between the PedsQL™
and functional data was assessed by bivariate correlations (Pearson’s
correlations). All hypothesis tests conducted were 2-tailed. A p
value < 0.05 was considered statistically significant. Statistical
analyses were performed using SAS (SAS version [9.2] of the
Cary, NC, USA).

3. Results

One hundred and seven patients fulfilled the inclusion
criteria and participated in the study, but 9 patients were lost at
follow-up. Ninety-eight patients (mean age = 8.4 years, SD
2.29) had both baseline and follow-up data and were included in
the analysis. Forty-three patients were aged 7 years or below.
Within the year when the study was performed, 9 of the 98
patients lost the ability to complete the 6MWT and timed items
and were included in the analysis with a result equal to the worst
performance in the group. All the tests were performed safely
without any major fall during the assessments.

3.1. Baseline data

Descriptive statistics for PedsQL™ Child Self-Report,
Parent Proxy-Report and functional data at baseline are shown in
Tables 1 and 2.

| Table 1
| Descriptive statistics for PedsQL™ Child Self-Report and Parent Proxy-Report data at baseline and at 12 month follow-up. Mean (±SD). |

<table>
<thead>
<tr>
<th>Children</th>
<th>Parents</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Baseline</td>
</tr>
<tr>
<td>PedsQL™ Generic Core Scales (GSC)</td>
<td>74.5 (±15.8)</td>
</tr>
<tr>
<td>PedsQL™ GSC – 1st domain (Physical Function Score)</td>
<td>68.9 (±21.9)</td>
</tr>
<tr>
<td>PedsQL™ Multidimensional Fatigue Scale (PedsQL MFS)</td>
<td>76.3 (±17.4)</td>
</tr>
<tr>
<td>PedsQL™ MFS – 1st domain (General Fatigue Scale)</td>
<td>75.9 (±21.1)</td>
</tr>
<tr>
<td>PedsQL™ Neuro muscular Module (NMM)</td>
<td>81.4 (±12.8)</td>
</tr>
<tr>
<td>PedsQL™ NMM – 1st domain (About My Neuromuscular Disease)</td>
<td>82.3 (±10.7)</td>
</tr>
</tbody>
</table>

p significant values in bold. *p < 0.05, **p < 0.01, ***p < 0.001 or above.

| Table 2
| 6 minute walk test (6MWT), North Star Ambulatory Assessment (NSAA), 10 meter timed walk/run test (10 meter) and Gowers test data at baseline, at 12 month follow-up and changes over 12 months (Delta). Mean (±SD). |

<table>
<thead>
<tr>
<th></th>
<th>Baseline</th>
<th>12 Months</th>
<th>DELTA</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>6MWT (m)</td>
<td>383.8 (±88.9, 119.5–546.2)</td>
<td>345.5 (±139.8, 0–563)</td>
<td>−38.3 (±82.6, −285–184.5)</td>
<td>***&lt;0.0001</td>
</tr>
<tr>
<td>North Star Ambulatory Assessment Score (34/34)</td>
<td>24.2 (±6.8, 4–34)</td>
<td>21.6 (±9.1, 0–34)</td>
<td>−2.6 (±4.7, −14.7–12)</td>
<td>***&lt;0.0001</td>
</tr>
<tr>
<td>10 meter timed walk/run test (s)</td>
<td>7.4 (±2.6, 3.6–17.4)</td>
<td>8.9 (±5.4, 2–24)</td>
<td>1.5 (±3.8, −5.4–14.9)</td>
<td>**0.0009</td>
</tr>
<tr>
<td>Gowers test (s)</td>
<td>8.0 (±6.8, 1.41–27)</td>
<td>11.8 (±10.7, 2–31)</td>
<td>3.8 (±6.2, −3.6–29.6)</td>
<td>***&lt;0.0001</td>
</tr>
</tbody>
</table>

p significant values in bold. *p < 0.05, **p < 0.01, ***p < 0.001 or above.
3.2. Correlations between PedsQL™ and functional data at baseline

3.2.1. PedsQL™ child self-report

At baseline the PedsQL™ GCS and the GCS-first domain, The GSC-Physical Function Score, correlated with age and all the functional measures with the strongest correlation found with Gowers test. The PedsQL™ MFS correlated with Gowers test and 6MWT and its first domain, the MFS-General Fatigue Scale, with 6MWT only. The PedsQL™ NMM correlated with all the functional measures with the exception of NSAA and its first domain, PedsQL™ NMM-About My Neuromuscular Disorder, did not show any significant correlation. All the other correlations were not significant. Details in Table 3.

3.2.2. PedsQL™ parent proxy-report

At baseline the PedsQL™ GCS and the GCS-first domain correlated with age and all the functional measures with the strongest correlations found between GCS-Physical Function Score and NSAA, Gowers test and 6MWT.

Both the PedsQL™ MFS and the MFS-first domain correlated with 6MWT.

The PedsQL™ NMM correlated with 10 meter and 6MWT and its first domain, the PedsQL™ NMM-About My Neuromuscular Disease, correlated with all outcome measures, with the strongest correlation with 6MWT. All the other correlations were not significant. Details in Table 3.

3.3. Longitudinal data

On the Child Self-Report scores there was a significant decrease between baseline and 12 months in the PedsQL™ GCS and its first domain (GSC-Physical Function Score). All the other inventories did not significantly differ between baseline and 12 months (Table 1).

On the Parent Proxy-Report there was a significant decrease between baseline and 12 months on all tools with the exception of the GCS – Physical Function Score (Table 1).

All the selected functional outcome measures detected a significant decline in function over the 12-month period (Table 2). The decline was more obvious in the subgroup of boys above the age of 7 years (p < 0.001), as the younger ones did not show a significant difference over the 12-month period, with several patients showing an improvement in all the selected outcome measures. Significant correlations were found between baseline values and the 12-month changes of functional scales [10].

3.4. Correlation of 12 month changes

3.4.1. PedsQL™ child self-report

The correlation between the 12 month changes on the PedsQL™ GCS and the other PedsQL™ tools were all significant with the strongest correlation with its first domain (PedsQL™ GCS-Physical Function Score) r = 0.64, p < 0.0001.

Table 3

Correlations between PedsQL™ Child Self- and Parent Proxy-Report and age and functional measures at baseline.

<table>
<thead>
<tr>
<th></th>
<th>Age</th>
<th>NSAA</th>
<th>10 meters</th>
<th>Gowers test</th>
<th>6MWT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline child self-report</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PedsQL™ Generic Core Scales (GSC)</td>
<td>r = −0.33</td>
<td>r = 0.26</td>
<td>r = −0.26</td>
<td>r = −0.43</td>
<td>r = 0.27</td>
</tr>
<tr>
<td></td>
<td>*p &lt; 0.001</td>
<td>*p &lt; 0.01</td>
<td>*p &lt; 0.009</td>
<td>***p &lt; 0.0001</td>
<td>*p &lt; 0.008</td>
</tr>
<tr>
<td>PedsQL™ GSC – 1° domain (PFS)</td>
<td>r = −0.36</td>
<td>r = 0.32</td>
<td>r = −0.34</td>
<td>r = −0.41</td>
<td>r = 0.25</td>
</tr>
<tr>
<td></td>
<td>***p = 0.0004</td>
<td>***p = 0.002</td>
<td>***p &lt; 0.0007</td>
<td>***p &lt; 0.0001</td>
<td>*p &lt; 0.02</td>
</tr>
<tr>
<td>PedsQL™ Multidimensional Fatigue Scale (PedsQL™ MFS)</td>
<td>r = −0.04</td>
<td>r = 0.20</td>
<td>r = −0.18</td>
<td>r = −0.21</td>
<td>r = 0.23</td>
</tr>
<tr>
<td></td>
<td>p = 0.68</td>
<td>p = 0.06</td>
<td>p = 0.09</td>
<td>***p = 0.04</td>
<td>*p &lt; 0.02</td>
</tr>
<tr>
<td>PedsQL™ MFS – 1° domain (GFS)</td>
<td>r = 0.11</td>
<td>r = 0.14</td>
<td>r = −0.16</td>
<td>r = −0.17</td>
<td>r = 0.26</td>
</tr>
<tr>
<td></td>
<td>p = 0.31</td>
<td>p = 0.17</td>
<td>p = 0.13</td>
<td>*p = 0.1</td>
<td>*p &lt; 0.01</td>
</tr>
<tr>
<td>PedsQL™ NM</td>
<td>r = −0.10</td>
<td>r = 0.18</td>
<td>r = −0.24</td>
<td>r = −0.26</td>
<td>r = 0.20</td>
</tr>
<tr>
<td></td>
<td>p = 0.30</td>
<td>p = 0.07</td>
<td>*p = 0.02</td>
<td>*p &lt; 0.01</td>
<td>*p &lt; 0.04</td>
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<tr>
<td>PedsQL™ NM – 1° domain</td>
<td>r = −0.02</td>
<td>r = 0.15</td>
<td>r = −0.16</td>
<td>r = −0.18</td>
<td>r = 0.13</td>
</tr>
<tr>
<td></td>
<td>p = 0.88</td>
<td>p = 0.14</td>
<td>p = 0.12</td>
<td>p = 0.08</td>
<td>p = 0.22</td>
</tr>
<tr>
<td>Baseline parent proxy-report</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PedsQL Generic Core Scales (GSC)</td>
<td>r = −0.34</td>
<td>r = 0.36</td>
<td>r = −0.31</td>
<td>r = −0.36</td>
<td>r = 0.36</td>
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<tr>
<td></td>
<td>***p = 0.0007</td>
<td>***p = 0.0004</td>
<td>***p = 0.002</td>
<td>***p = 0.0002</td>
<td>***p = 0.0003</td>
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<tr>
<td>PedsQL GSC – 1° domain (PFS)</td>
<td>r = −0.30</td>
<td>r = 0.43</td>
<td>r = −0.39</td>
<td>r = −0.45</td>
<td>r = 0.49</td>
</tr>
<tr>
<td></td>
<td>***p = 0.003</td>
<td>***p &lt; 0.0001</td>
<td>***p &lt; 0.0001</td>
<td>***p &lt; 0.0001</td>
<td>***p &lt; 0.0001</td>
</tr>
<tr>
<td>PedsQL Multidimensional Fatigue Scale (PedsQL MFS)</td>
<td>r = −0.03</td>
<td>r = 0.18</td>
<td>r = −0.16</td>
<td>r = −0.12</td>
<td>r = 0.27</td>
</tr>
<tr>
<td></td>
<td>p = 0.77</td>
<td>p = 0.08</td>
<td>p = 0.13</td>
<td>p = 0.25</td>
<td>*p &lt; 0.006</td>
</tr>
<tr>
<td>PedsQL MFS – 1° domain (GFS)</td>
<td>r = −0.02</td>
<td>r = 0.16</td>
<td>r = −0.16</td>
<td>r = −0.14</td>
<td>r = 0.26</td>
</tr>
<tr>
<td></td>
<td>p = 0.9</td>
<td>p = 0.12</td>
<td>p = 0.13</td>
<td>p = 0.18</td>
<td>***p &lt; 0.009</td>
</tr>
<tr>
<td>PedsQL NM</td>
<td>r = −0.05</td>
<td>r = 0.15</td>
<td>r = −0.25</td>
<td>r = −0.15</td>
<td>r = 0.21</td>
</tr>
<tr>
<td></td>
<td>p = 0.6</td>
<td>p = 0.13</td>
<td>*p = 0.01</td>
<td>p = 0.15</td>
<td>*p = 0.04</td>
</tr>
<tr>
<td>PedsQL NM – 1° domain</td>
<td>r = −0.17</td>
<td>r = 0.38</td>
<td>r = −0.35</td>
<td>r = −0.29</td>
<td>r = 0.46</td>
</tr>
<tr>
<td></td>
<td>p = 0.1</td>
<td>***p = 0.0001</td>
<td>***p = 0.0004</td>
<td>***p = 0.0001</td>
<td>***p &lt; 0.0001</td>
</tr>
</tbody>
</table>

*p significant values in bold. *p < 0.05, **p < 0.01, ***p < 0.001 or above.
The correlation between the 12 month changes on the PedsQL™ inventories and the functional measures were all negligible with the exceptions of the PedsQL™ MFS with 10 meter timed walk/run test and PedsQL™ NMM-About My Neuromuscular Disease and Gowers test.

A correlation with age was shown for some of the inventories (Table 4). Fig. 1 shows as an example the distribution of individual changes on PedsQL™ GCS and functional scales. When we considered the subgroup of patients above the age of 7, the correlation between the 12 month changes on the PedsQL™ inventories and the functional measures were also all negligible with the exceptions of the 12 month changes on PedsQL™ GCS-Physical Function Score and Gowers test (r = 0.23, p = 0.01) and between PedsQL™ MFS and 6MWT (r = 0.25, p = 0.01).

A correlation with age was shown for some of the inventories (PedsQL™ MFS-General Fatigue Scale, r = 0.23, p = 0.01, PedsQL™ NMM, r = 0.28, p = 0.003; PedsQL™ NMM-About My Neuromuscular Disease, r = 0.38, p = 0.0001).

4. Discussion
The use of relatively new functional outcome measures in clinical trials in boys with DMD has highlighted the need to better understand how changes in these measures relate to quality of life. In the last few years there has therefore been increasing interest to identify HRQOL tools to be included as a secondary outcome measures in clinical trials in children with DMD. Although HRQOL measurements have been increasingly acknowledged as essential health outcome measures in clinical trials, the available pediatric tools are very scanty and the choice is even poorer if we look for disease-specific measures. Generic pediatric HRQOL instruments have been widely used in several disorders. They have the advantage that they enable comparisons across pediatric populations and facilitate benchmarking with healthy population norms. Disease-specific measures on the other hand have the advantage of enhancing measurement sensitivity for health domains germane to a particular chronic health condition. The PedsQL™ is a versatile instrument widely used in children with chronic diseases [26]. Several disease-specific modules have also
been developed, such as those for asthma, arthritis, cancer, cardiac disease, diabetes. The recent PedsQL™ NMM, originally developed for children with SMA [17], was subsequently also applied in DMD boys, also in comparison with an other DMD version [16,19]. In this study we used both the generic PedsQL™ GCS and the dedicated neuromuscular module. Furthermore, as fatigability is an important issue in DMD boys and is partly recorded by the 6MWT, we also decided to use the fatigability scale, the PedsQL™ MFS. We focused on their global scores and on the first domains, supposed to better relate to functional aspects, this resulting in a choice of 6 measures of HRQOL.

Our results showed that the total scores of PedsQL™ GCS Child Self-Report were between 25.78 and 100 at baseline and between 38.81 and 100 at 12 months with an average of 74.5 and 71 respectively. The average scores in our cohort are lower than those reported for healthy controls in two previous studies (between 87 and 88.8) [16,19], but higher than in their DMD cohorts, as these ranged between 60.4 and 62.7 [16,19]. The results however are not easily comparable because of different inclusion criteria as we included only ambulant boys from the age of 5 and in the first of the two previous studies [16] nearly 90% of the DMD boys were non ambulant and the second focused on the 8–12 years age range [19].

Another difference with previous studies is that by also including younger boys, we were able to compare parents’ and children’s responses in all ambulatory age range. With one exception parents’ responses were always lower than children’s self report both at baseline and at 12 months. These findings confirm and expand previous evidence of poor agreement between parents’ and children’s responses [16,19].

The aim of our study however was not to duplicate previous reports on quality of life in DMD or their correlation with different demographic data or intervention such as steroids, but to establish the value of the questionnaires in assessing changes in HRQOL. At variance with previous studies [19–21], we also used the PedsQL™ NMM, a module specifically devised for patients with NMD and validated in DMD, with selected items thought

![Fig. 1. Distribution of individual 12 month changes of PedsQL™ Child Self-Report Generic Core Scales scores versus 6MWT (A), NSAA (B), 10 meter timed walk/run test (10 meter) (C) and Gowers test (D) according to age (≤ or >7 years).](image-url)
to be more relevant for the disease [16]. Surprisingly, its use did not appear to increase the level of significance compared to the general module with the exception of the first domain, that better correlated with all functional measures in the parents’ questionnaire.

It is of interest that the 6MWT appeared to be the measure that at baseline better correlated with all the inventories.

When we assessed the possible concordance of 12 month changes between PedsQL™ GCS versus its first domain PedsQL™ GCS – PFS and the other PedsQL™ measures, we found significant correlations with the strongest correlation with its first domain in both parents’ and children’s versions.

In contrast, the correlation was often not significant between the 12 month changes on the different PedsQL™ tools and the functional assessments. While the functional scores significantly decreased over 12 months, the results of the inventories were more variable. On the Child Self-Report scores, only the GCS and its first domain PedsQL™ GCS – PFS had a significant decrement over 12 months. At variance, on the parents’ version the scores decreased with age and disease progression as previously reported by other studies [19–21] with the exception of the PedsQL™ GCS – PFS, although this tool is targeted to functional aspects. Even when the changes observed on the functional tests and on the inventories were concordant, trending in the same direction, the changes in the two groups of tools did not always happen in the same direction in individual patients (e.g. in Fig. 1). As a result, the overall correlation between changes in function and PedsQL™ was, with few exceptions, not significant both in the whole cohort and in the subset of patients above the age of 7 years. Interestingly, in the subset of patients above the age of 7 years we found a correlation between changes in PedsQL™ MFS and 6MWT both on children’s and parents’ versions. This is not surprising considering that the 6MWT is a measure of endurance and may better reflect fatigability, as also shown in the correlation between PedsQL™ MFS parents’ version and 6MWT at baseline.

The only previous longitudinal study reporting both PedsQL™ parental self-report and functional measures changes in 24 DMD boys with age range similar to our cohort showed a weak correlation at baseline between PedsQL™ GCS and 6MWT and timed items. Thirteen of the 24 also had follow-up assessments and the correlation of the 12 month changes was negligible [21]. Our study, performed in a larger cohort and using additional measures, confirm that 12 month changes in functional performances, particularly at this disease stage and over 12 months.

Previous studies using other tools, such as the Pediatric Outcomes Data Collection Instrument (PODCI), have shown higher sensitivity to functional changes over 12 months in ambulant DMD compared to PedsQL™ [21]. This difference may be explained by the fact that the PODCI response options address the perceived physical difficulty of activities: e.g. “during the last week has it been easy or hard for your child to walk one block?”, response options include: “easy, a little hard, very hard, can’t do at all”. In contrast, the PedsQL™ response in based on self-selected frequency and/or importance: e.g. “In the past one month how much of a problem has your child had with . . .”, response choices include “never, almost never, sometimes, often and almost always”.

Our findings highlight the complexity of the relationship between HRQOL tools and functional measures. Functional changes on the 6MWT and on the NSAA reflect changes on activities of daily living, that are not always addressed by PedsQL™ and other available HRQOL tools. Furthermore, PedsQL™ scores reflect also social and emotional aspects that may not parallel to functional changes.

Further studies on longer time frame or using other tools, such as the PODCI and/or specific questionnaires addressing changes in activity of daily living, may help to elucidate to which extend the available pediatric HRQOL tools are capable of mirroring the functional changes and/or eventually to identify new valuable tools.

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References


