CASE REPORT
The Change of Grip Strength in a Patient with Congenital Myotonic Dystrophy Over a 4-year Period

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ABSTRACT. Myotonic dystrophy (MyD) is a neuromuscular disease that is autosomal dominant and the most common form of muscular dystrophy affecting adults. The clinical features of MyD include a multisystemic disorder characterized by myotonia, progressive muscle weakness and wasting, cataracts, premature balding and mental retardation. The most severe type of MyD is classified as congenital MyD (CMyD). The muscle weakness in CMyD is very severe, but muscle development can be observed in the period of growth. However, no clinical case of this type has been reported yet. Therefore, we report on a girl with CMyD who had an increase in muscle strength over a four-year period. The girl with CMyD participated in this study from the age of 9 to the age of 12. The measurement of muscle strength was recorded as the maximum score of grip strength with the use of dynamometers. Grip strength was assessed once a year by the same two physical therapists. Grip strength of CMyD for each year was markedly weak when compared with the normal controls, but muscle strength changed within some specific growth areas. The muscle weakness in CMyD was remarkable, but the result showed that specific muscle strength of CMyD in childhood was actually increased.

Key words: congenital myotonic dystrophy, development, grip strength

Myotonic dystrophy (MyD) is a neuromuscular disease caused by amplification of an unstable CTG repeat in the 3' untranslated region on human chromosome 19q13.31. MyD has an incidence of 5.5 in 100,000 and is the most common form of adult-type muscular dystrophy2). The clinical features of MyD are markedly variable, characterized by myotonia, progressive muscle weakness and atrophy, cataracts, premature balding and mental retardation3). The clinical severity shows extreme variation even among family members. The symptoms and severity of MyD are related to age at the time of onset. The most mild and late-onset form of MyD is classified as adult type myotonic dystrophy (AMyD). On the other hand, the type that is most severe, with onset from birth is congenital myotonic dystrophy (CMyD)4). Progressive muscle weakness is one of the cardinal symptoms in MyD but, in CMyD, it has been suggested that those who survive CMyD for a year after birth have an improvement in their muscle strength5). However, there is no detailed report in the literature about muscle strength improvement in CMyD.

The main symptoms of MyD are progressive muscle weakness and atrophy. The degree of muscle weakness and the severity of other clinical symptoms of MyD are correlated with the length of the CTG repeat6-10). Gharehbaghi-Schnell et al. indicated a correlation between clinical severity and the CTG repeat based on the results of comparing the CTG repeat and the Muscular Disability Rating Scale (MDRS) scores of 18 patients with MyD8). Eguchi et al. reported a correlation between the number of CTG repeats and three clinical features (age at the time of onset, degree of mental retardation and severity of muscle

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We encountered one patient with CMyD whose grip strength increased over four years. The aims of this report, through this one case, were to show the change specificity of grip muscle strength in CMyD as a basic factor for prognosis and to design an effective physical therapy program for CMyD.

**Case**

**Progress before evaluation of muscle strength**

A female child, diagnosed with CMyD, participated in this study. Her CTG repeat size was about 2,200 repeats \(^{10}\). The subject’s growth was obtained from her clinical chart and the data collection was continued until she was 8 years old.

History of subject from birth to 2 months: She was born at 35 weeks of gestation after a cesarean section was performed due to polyhydramnios and fetal asphyxia; her birth weight was 2,238 grams. Her Apgar scores were 1 and 3 at 1 and 5 minutes respectively. She had general muscle hypotonia (frog leg position) and required ventilation for one month after birth. At 2 months, she was discharged from the hospital because she could sufficiently swallow and suck.

Milestones assessment: Tracking, 4 months; Moro reflex, 4 months; smile, 5 months; handkerchief on face, 6 months; rolling over, 10 months; crawl, 12 months; standing and walking with support, 16 months; walking alone (about 10 steps), 22 months; running, 3 years; speech (one word), 3 years 9 months; climbing up and down the stairs, 3 years 9 months; pedaling a bicycle with support, 4 years 1 month; express the need to urinate, 4 years 1 month; speech (over 3 words), 7 years; jumping, 7 years; speech (making a sentence), 8 years.

The subject had never done any specific muscle strengthening training or physical therapeutic exercise until the time of this study. The measurements were done in August every year from 2001 to 2004, from the age of 9 to 12 years. She did not undertake any specific physical therapy or muscle training. Her lifestyle showed no difference until she reached 12 years of age. Her mother was diagnosed with AMyD with no symptoms, except for grip myotonia. The mother had no problem with ADL.

Before any measurement of muscle strength was done, the normal range of motion was verified and no deformity was found. Hyper- or hypomobility was not observed in the screening test for ROM in the subject. At the ages of 9 to 12, she was 121.0, 125.1, 128.2 and 133.4 cm tall and weighed 21.2, 22.0, 26.0 and 26.2 kg, respectively.

The subject and her mother were informed about the parameters of this study and consented to a series of investigations. In addition, this study obtained approval from the Committee for Graduate School of Health Sciences, Sapporo Medical University (approval number: 2001MP-02-12).

**Methods**

**Measurement protocol**

Grip strength tends to decrease in inverse relation to the size of the CTG repeats. The grip strength was measured by the same handgrip dynamometer (TK-1201, TAKEI KK) each year. The grip strength test was performed by the same two physical therapists and was measured in accordance with a standard protocol. Grip strength was measured 3 times for each hand (total 6 times in all), each separated by a 30 sec rest period. The highest result of the six trials was used as the grip strength. The results were compared with the normal strength that was matched by age (9, 10, 11 and 12 years old) and gender from the investigation of physical strength and activity in the 2002 fiscal year \(^{20}\).

Intraclass correlation coefficients (ICC) were used to analyze the reliability of each measurement using SPSS Ver13.0J for Windows. Analysis of reliability for grip strength data was conducted for the 3 measurement values for each hand (right and left hand) for each year (4 years).

**Validity of the instrument**

The validity of the handgrip dynamometer was measured with Fess’s method each year for the entire four-
## Table 1. Summary of grip strength measured investigation in subjects with myotonic dystrophy

<table>
<thead>
<tr>
<th>Source</th>
<th>Number of subjects</th>
<th>Gender</th>
<th>Mean age (range)</th>
<th>Number of CTG repeats (range)</th>
<th>Materials of measurement</th>
<th>Mean grip strength ± S.D.³ (range)§§§</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moxley et al. 2007¹¹</td>
<td>29</td>
<td>M*= 18 F**= 11</td>
<td>45.9 (18–80) years</td>
<td>Mean= 563 (169–1731) Jamar Dynamometer</td>
<td>10.4 (NR) kg</td>
<td></td>
</tr>
<tr>
<td>Guimaraes et al. 2007¹²</td>
<td>37</td>
<td>M= 20 F= 17</td>
<td>M= NR*** (8–64) years, F= NR (15–69)</td>
<td>NR (NR) Electronic dynamometer (Kratos Equipamentos Industriais Ltd.)</td>
<td>M= 5.00 (0.20–40.07) kg, F= 3.98 (0.51–28.96) kg</td>
<td></td>
</tr>
<tr>
<td>Whittaker et al. 2006¹³</td>
<td>158</td>
<td>M= 79 F= 79</td>
<td>M= 43.6 (12–75) years, F= 47.0 (19–75)</td>
<td>NR (NR) Handgrip dynamometer (Takei)</td>
<td>M= 16.1 (NR) kg, F= 11.1 (NR) kg</td>
<td></td>
</tr>
<tr>
<td>Aldehag et al. 2005¹⁴</td>
<td>5</td>
<td>M= 2 F= 3</td>
<td>M= 47.5 (28 and 67) years, F= 55.7 (54, 56 and 57)</td>
<td>28 years M= 500–800, 67 years M= 600–900, 54 years M= 600–700, 56 years F= 200–300 and 57 years F= 40–145 Grippit (Goteborg)</td>
<td>Rt</td>
<td>¹⁵= 10.94±2.92 (NR) kg, Lt</td>
</tr>
<tr>
<td>Logigian et al. 2005¹⁵</td>
<td>25</td>
<td>M= 16 F= 9</td>
<td>NR (18–80) years</td>
<td>Mean= 550 (169-1469) Grip ergometer</td>
<td>10.12 (2.42–27.06) kg</td>
<td></td>
</tr>
<tr>
<td>Mathieu et al. 2003¹⁶</td>
<td>50</td>
<td>M= 27 F= 23</td>
<td>41.1 (16–67) years</td>
<td>16 %&lt; 200, 2 %= 201–400, 30 %= 401–850, 16 %= 851–1100, 18 %= 1101–1500 and 18 %&gt;1500 Jamar dynamometer</td>
<td>M= 16.33±13.48 (NR) kg, F= 12.39±7.59 (NR) kg</td>
<td></td>
</tr>
<tr>
<td>Nitz et al. 1999¹⁷</td>
<td>36</td>
<td>M= 18 F= 18</td>
<td>36.9 (NR) years</td>
<td>NR (NR) Jamar dynamometer (Asimow engineering)</td>
<td>Rt= 10.51±6.77 (NR) kg, Lt= 9.89±6.23 (NR) kg</td>
<td></td>
</tr>
<tr>
<td>Grant et al. 1987¹⁸</td>
<td>10</td>
<td>M= 6 F= 4</td>
<td>40.4 (28–64) years</td>
<td>NR (NR) Grip strength ergometer</td>
<td>M= 10.25±12.3 (NR) kg, F= 10.78±9.01 (NR) kg</td>
<td></td>
</tr>
<tr>
<td>Omdahl et al. 1986¹⁹</td>
<td>5</td>
<td>M= 3 F= 2</td>
<td>M= 39.9 (33, 39 and 45) years, F= 42.5 (39 and 46)</td>
<td>NR (NR) Vigotimeter</td>
<td>Rt= 0.18 (0–0.6) kg, Lt= 0.12 (0–0.4) kg after treatment; Rt= 0.28 (0.1–0.6) kg, Lt= 0.40 (0.1–0.8) kg</td>
<td></td>
</tr>
</tbody>
</table>

*: M= Male, **: F= Female, ***: NR= not reported, ³: S.D.= Standard deviation, ⁴: The unit of force was converted to kg, ⁵: Rt= right, ⁶: Lt= left.
year period\textsuperscript{21}). The handgrip dynamometer was in error by less than 0.75 kg.

### Results

**Grip strength**

The ICC score was 0.83, indicating that the results were very reliable.

As shown in Table 2, the grip strength results of the subject by year were 6.5 kg (Left [L]: 5.5–6.0 kg, Right [R]: 6–6.5 kg) (minimum-maximum), 5.5 kg (L: 5–5.5, R: 4.5–5.0 kg), 7.5 kg (L: 6.0–7.0 kg, R: 6.5–7.5 kg) and 7.5 kg (L: 6–7.5 kg, R: 6–7.5 kg). The grip strength percentages were 45.8, 32.7, 37.5 and 33.6\% for the ages of 9, 10, 11 and 12, respectively, compared with normal control matched subjects by age and gender. The grip strength showed a moderate increase with aging, but not linearly. The difference between grip strength in this case and the normal control became larger year by year during the measurement over the four-year period.

**The changes of weight and height**

The height and weight values increased year by year (Table 2). However, the percentiles of the subject’s height were below the normal average and decreased year by year (90.4, 89.0, 87.0 and 87.6\%), but not to under 85\%. Percentiles of her weight were not linearly plotted (70.5, 63.3, 65.1 and 58.7\%), and the change pattern of weight was similar to that for grip-strength changes.

### Discussion

Mathieu \textit{et al.} reported that muscle strength tended to decrease with age in MyD\textsuperscript{16}. Eguchi \textit{et al.}, however, showed that the degree of muscle weakness varied in each case of AMyD, and there was little agreement on the muscle weakness found in MyD\textsuperscript{9}. Until now it has been considered that high-resistance strength training causes overwork weakness in MyD. However, Tollback \textit{et al.} demonstrated that patients with MyD had improved muscle strength without any observed negative side effect after high-resistance training\textsuperscript{7}. Recently, Lideman \textit{et al.} substantiated that strength training improved muscle function of patients with slowly progressive hereditary neuromuscular disorders\textsuperscript{22} such as CMyD. However, most of the subjects of these studies did not actually have CMyD, but rather AMyD, and there are actually very few studies on the changes of muscle strength with aging in CMyD. Therefore, we studied muscle strength changes in a patient with CMyD from the age of 9 to 12 years old.

The results of this study showed that the grip strength at 12 years of age was increased in comparison with that at 9 years of age. This result might show the possibility of increased muscle strength with development in CMyD. The development rate of muscle strength in CMyD was very slow in comparison with the normal control. The grip strength in this case was 50\% when compared with the normal control, and the gap tended to increase year by year. In addition, the muscle strength did not linearly increase. The pattern of percentile change showed the similarity between grip strength and weight. In particular, the measured values of grip strength and weight decreased from 9 years old to 10 years old. Weight values are affected by many elements and a decrease of weight in childhood usually suggests an abnormal condition or a nutritional problem. The cause of the weight decrease was not known in this case, but the decrease of grip strength from ages 9 to 10 might have been connected with not only pathological

\begin{table}[h]
\centering
\begin{tabular}{lccccc}
\hline
 & \textbf{9 y}\textsuperscript{*} & \textbf{10 y} & \textbf{11 y} & \textbf{12 y} \\
\hline
\textbf{Height (cm)} & & & & & \\
subject & 121.0 & 125.1 & 128.2 & 133.4 \\
N.C.\textsuperscript{**} (mean ± S.D.)\textsuperscript{***} & 133.8 ± 6.3 & 140.6 ± 6.9 & 147.4 ± 6.6 & 152.3 ± 5.6 \\
% & 90.4 & 89.0 & 87.0 & 87.6 \\
\hline
\textbf{Weight (kg)} & & & & & \\
subject & 21.2 & 22.0 & 26.0 & 26.2 \\
N.C. (mean ± S.D.) & 30.1 ± 5.4 & 34.7 ± 6.9 & 39.9 ± 7.6 & 44.7 ± 7.5 \\
% & 70.5 & 63.3 & 65.1 & 58.7 \\
\hline
\textbf{Grip strength (kg)} & & & & & \\
subject (maximum) & 6.5 & 5.5 & 7.5 & 7.5 \\
N.C. (mean ± S.D.) & 14.2 ± 3.3 & 16.8 ± 3.9 & 20.0 ± 4.5 & 22.3 ± 4.4 \\
% & 45.8 & 32.7 & 37.5 & 33.6 \\
\hline
\end{tabular}
\caption{Height, weight and grip strength and percentiles of normal controls in each year}
\end{table}

\textsuperscript{*}: y=years old, \textsuperscript{**}: N.C.=normal control, \textsuperscript{***}: the values for normal control height, weight and grip strength are from an investigation of physical strength and activity in the 2002 fiscal year\textsuperscript{20).}
effects, but also the total body condition. Therefore, the
decrease of grip strength from 9 to 10 in this particular case
is not necessary by the negative evidence for an increase of
muscle strength in patients with myotonic dystrophy.

In the skeletal muscle immature fibers are observed
predominantly in childhood in patients with CMyD23), but
in adults with CMyD pathological observations are similar
to those of late onset MyD24). Kikuchi and colleagues
reported that the muscle pathological observation of CMyD
with growth to 10 years old was similar to that of adult
onset MyD and type IIC (immature) fibers were not
detected25). These reports suggest that the maturation of
the skeletal muscle fiber type in CMyD progresses with aging,
but that for muscle the pathological observations are
the same as for AMyD. Therefore, in this case, muscle strength
increased but the muscle expanded in a different manner
from the normal control.

This case, suggests that CMyD patients have the
possibility to increase muscle strength from 9 to 12 years of
age, and Tollback et al. reported that muscle strength was
improved without any negative side effect after high-
resistance training7). For physical therapy in CMyD, our
results and the previous report suggest a need for physical
exercise including a high-resistance training program to
limit the growing gap between children with CMyD and
normal children, after first confirming the total body
development and condition.

**Conclusion**

The importance of this study is that the change of
muscle strength in CMyD showed an unstable increase and the
difference of muscle strength between the patient with
CMyD and the normal control became greater within the
period of 9 to 12 years of age. It is difficult to arrive at a
general conclusion as to the change of muscle strength in
CMyD, because there was only one participant in this study.
Therefore, we think it is worthwhile to expand this research
with a greater number of similar subjects.

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