Natural History of Facioscapulohumeral Dystrophy in Children
A 2-Year Follow-up

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Abstract

Background and Objectives
Data on the natural history of facioscapulohumeral dystrophy (FSHD) in childhood are limited and critical for improved patient care and clinical trial readiness. Our objective was to describe the disease course of FSHD in children.

Methods
We performed a nationwide, single-center, prospective cohort study of FSHD in childhood assessing muscle functioning, imaging, and quality of life over 2 years of follow-up.

Results
We included 20 children with genetically confirmed FSHD who were 2 to 17 years of age. Overall, symptoms were slowly progressive, and the mean FSHD clinical score increased from 2.1 to 2.8 \((p = 0.003)\). The rate of progression was highly variable. At baseline, 16 of 20 symptomatic children had facial weakness; after 2 years, facial weakness was observed in 19 of 20 children. Muscle strength did not change between baseline and follow-up. The most frequently and most severely affected muscles were the trapezius and deltoid. The functional exercise capacity, measured with the 6-minute walk test, improved. Systemic features were infrequent and nonprogressive. Weakness-associated complications such as lumbar hyperlordosis and dysarthria were common, and their prevalence increased during follow-up. Pain and fatigue were frequent complaints in children, and their prevalence also increased during follow-up. Muscle ultrasonography revealed a progressive increase in echogenicity.

Discussion
FSHD in childhood has a slowly progressive but variable course over 2 years of follow-up. The most promising outcome measures to detect progression were the FSHD clinical score and muscle ultrasonography. Despite this disease progression, an improvement on functional capacity may still occur as the child grows up. Pain, fatigue, and a decreased quality of life were common symptoms and need to be addressed in the management of childhood FSHD. Our data can be used to counsel patients and as baseline measures for treatment trials in childhood FSHD.

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Facioscapulohumeral muscular dystrophy (FSHD) is a slowly progressive muscular dystrophy and one of the most common hereditary muscle diseases, with an estimated prevalence of 12 per 100,000 in the Netherlands. FSHD typically causes progressive, asymmetric weakness of the facial, scapulohumeral, tibial, and axial muscles, with a highly heterogeneous disease severity.

Type 1 FSHD, responsible for >95% of the FSHD cases, is associated with a contraction of the D4Z4-repeat on chromosome 4q35. In the normal population, this array contains 11 to 100 D4Z4 repeats, whereas in FSHD, there is a contraction of the D4Z4 region to 1 to 10 repeat units, leading to D4Z4 hypomethylation. In type 2 FSHD, hypomethylation of the D4Z4 repeats is caused by mutations in SMCHD1 or DNMT3B gene. Both types lead to disease in the setting of a permissive allele, with aberrant DUX4 expression in skeletal muscle.

The age at symptom onset in FSHD is variable; in approximately one-fifth of all patients, FSHD starts in childhood. Approximately half of the children with FSHD fulfill the criteria of early-onset FSHD, a subgroup defined by facial weakness before the age of 5 years and scapular weakness before the age of 10 years. This subgroup is associated with a fewer number of D4Z4 repeats, more extensive muscle weakness, and more systemic features (including hearing loss, retinal abnormalities, epilepsy, intellectual disability, and cardiac arrhythmias) compared to those of similar age and disease duration with classic-onset FSHD.

Childhood FSHD is characterized by significant variability in symptoms and clinical disease course. However, so far, only cross-sectional observations are available, and little is known about the natural course in this age group. The rate of disease progression and the presence of systemic features within this population are natural course in this age group. The rate of disease progression and the presence of systemic features within this population are also unknown. Identification of prognostic factors that can be used in patient care and counseling is needed. Knowledge of the natural history is also required for the design of clinical trials.

In this study, we present the 2-year follow-up of a nationwide prospective cohort study of 28 children diagnosed with FSHD in the Netherlands. Baseline data were reported as the iFocus FSHD study. The primary aim of the current work was to assess the clinical features and natural course of FSHD in childhood in a 2-year follow-up period, to identify prognostic factors of disease severity and progression rate, and to find promising outcome measures that can serve as a base for further clinical trials.

**Methods**

**Patients and Design**

This is a prospective cohort study of FSHD in childhood. The iFocus FSHD study cohort includes 28 children with genetically confirmed type 1 FSHD who were 0 to 17 years of age and living in the Netherlands, of whom 1 patient was asymptomatic at baseline. A detailed protocol and description of the baseline characteristics can be found elsewhere.

**Clinical Assessments**

Clinical assessment of the patients was performed by the same examiners at study initiation and after 2 years of follow-up. The follow-up measurements took place from March 2018 until March 2020. All assessments were performed as a part of this study. The study visit took ≈4 hours, including a lunch break. When patients were unable to visit the study site, information was obtained by a phone interview with the patient or a parent, supplemented with information from the electronic health record, including a recent evaluation by the participant’s pediatric neurologist or rehabilitation physician. A full description of the study procedures can be found in the published protocol.

In brief, the clinical phenotype of each participant was assessed by the Motor Function Measure (MFM); the shoulder dimension of the Performance of the Upper Limb module; manual muscle force testing with the Medical Research Council (MRC) sum score from the trapezius, deltoid, biceps brachii, finger flexors, quadriceps femoris, tibialis anterior, and gastrocnemius muscle on both sides; and testing motor performance with the 6-minute walk test. The MFM assesses proximal, distal, and axial motor functions and has been developed for the broad spectrum of neuromuscular diseases. A short form is validated for the use in children 2 to 7 years of age. The Performance of the Upper Limb scale was developed to measure upper limb function in patients with Duchenne muscular dystrophy. Weakness is tested through 3 domains: shoulder, mid, and distal, of which the shoulder domain is relevant for the FSHD population. Results of the 6-minute walk test were expressed as a z score (the number of SDs from the mean for sex and age in healthy individuals).

In addition, the FSHD clinical score (an evaluation scale that assesses the strength and functionality of 6 different groups of muscles typically involved in FSHD; total score ranges from 0 when no symptoms are present to a maximum of 15) and (age-corrected) FSHD clinical severity scale (a score ranging from 0–10 that evaluates the extent of weakness in FSHD, considering the descending spread of symptoms) were scored. The burden of disease on quality of life was assessed by asking about the presence and location of pain, evaluating fatigability (NeuroQol fatigue domain), and scoring the Kidscreen questionnaire. These questionnaires were completed by children ≥8 years of age; if participants were younger, their caregivers were asked to complete these forms.
Information about (para)medical treatments was obtained from an interview with the patient and/or parent and concerned the doctors and therapists involved in the treatment and the frequency of these visits. Muscle imaging was performed with muscle ultrasonography, with visual grading according to the Heckmatt scale (a 4-point visual grading scale to classify muscle intensity, ranging from 1 for normal ultrasound appearance to 4 for very strongly increased echogenicity) and quantitative gray scale analysis to determine echogenicity of the bilateral trapezius, biceps brachii, rectus abdominis, rectus femoris, and tibialis anterior muscles. Gray scale results were expressed as a z score, that is, the number of SDs from the mean, after comparing the echogenicity to a muscle-specific reference value. Patients were screened for systemic disease features with ophthalmologic screening, including a fundoscopy and visual acuity testing by an ophthalmologist, tone- and speech audiometry by an audiometrist, an ECG, and observation of possible spinal deformities.

Statistical Analyses
Statistical analyses were performed with SPSS version 25 (IBM SPSS Inc, Chicago IL). Continuous parametric variables were expressed as mean and SDs, whereas continuous nonparametric variables were expressed as median and interquartile ranges. Categorical data were given as a percentage. The Kolmogorov-Smirnov test was used for testing normality assumptions in the distribution of the data. For continuous variables with a normal distribution, a paired t test was performed to test a mean difference between baseline and the 2-year follow-up. The Wilcoxon signed-rank test was performed to compare nonparametric continuous variables, and the McNemar test was used for comparing paired categorical data. For comparison between the early-onset and classic-onset groups at the 2-year follow-up, the Mann-Whitney U test was used for nonparametric continuous variables, the independent t test was used for continuous variables with a normal distribution, and the Fisher exact test was used for categorical variables. Linear mixed models were applied to analyze differences in disease progression. The continuous variable of the FSHD clinical score was used as the primary outcome, whereas the repeat length (divided into categories of 1–3, 4–6, and 7–10 D4Z4 repeat units), age at baseline, and sex were used as the fixed-effect predictors. The level of statistical significance was set at ≤ 0.05.

Data Availability
The data that support the findings of this study are available from the corresponding author on reasonable request.

Results
Demographics and Genetic Characteristics
Of the 28 participants in the baseline study, a total of 20 children from 17 families, including 3 sibling pairs, could be included for this follow-up study. Thirteen of them were clinically examined at the study location. An electronic health record review combined with a telephone interview was performed in 7 participants; 2 of them could not be examined at the study location because of the coronavirus disease 2019 (COVID-19) pandemic restrictions; the other 5 were unable to visit for other reasons, mainly because of the added burden of a hospital visit (Figure 1). Participants were very cooperative; all of them gave permission to approach them again for future study visits.

The demographic and genetic characteristics of the cohort of 20 children who completed the 2-year follow-up are shown in Table 1. We observed a significant correlation between mean number of units in the pathogenic D4Z4 repeat and the age at onset, with a lower number of D4Z4 units associated with a younger age at onset (r = 0.498, p = 0.015). No significant correlation between the number of D4Z4 repeats and disease severity as measured by the FSHD clinical score and clinical severity scale was found at either baseline or follow-up. In addition, no correlation between the age at onset and disease severity (FSHD clinical score and clinical severity scale) was found.

Table 1. Flow Diagram of Patient Inclusion in Previously Published Baseline Study (Gray) and Current 2-Year Follow-up Study (Blue)

| Baseline | iFocus baseline study (n = 28) |
|----------|--------------------------------|
| Genetic classification and demographics (N = 32) |
| Excluded (n = 4); Declined to participate (1) |
| Other reasons: could not be traced or identified (3) |
| iFocus baseline study (n = 28) |
| Clinical evaluation (19) |
| Phone interview and medical file review (1) |
| Excluded (n = 8); Medical file review (8) |
| Two year follow-up |
| iFocus two year follow-up study (n = 20) |
| Clinical evaluation (13) |
| Phone interview and medical file review (7) |

The 8 children for whom only an electronic health record review had been performed at baseline were not invited for follow-up. However, their baseline characteristics, including the mean age at time of baseline examination, age at onset of symptoms, age at time of diagnosis, sex, and mean number of D4Z4 repeats, were similar to those of the group of children who were included in this follow-up study.
Table 1: Demographic and Genetic Characteristics in Current 2-Year Follow-up Study

| Demographics                              | Value | Mean | Range  | SD  |
|-------------------------------------------|-------|------|--------|-----|
| Age at baseline examination, y            | 20    | 10.8 | 2 to 17 | 4.6 |
| Male sex, n (%)                           | 8 (40)|      |        |     |
| D4Z4a repeat units, n                    | 20    | 5.1  | 2 to 10 | 2.2 |
| Delta 1 methylation, %b                   | 13    | −7.2 | −13 to 5 | 5.3 |
| Hereditary pattern, n/total (%)           |       |      |        |     |
| Maternal AD                               | 10/20 (50) |     |        |     |
| Paternal AD                               | 7/20 (35)  |     |        |     |
| Sporadic (de novo mutations)              | 3/20 (15) |    |        |     |
| Mosaic Inheritance, n/total (%)          | 0/16 (0)   |     |        |     |
| Onset type, n/total (%)                   |       |      |        |     |
| Early onset                               | 8/20 (40) |     |        |     |
| Classic onset                             | 10/20 (50) |    |        |     |
| Too young for classification              | 1/20 (5)   |     |        |     |
| Asymptomatic                              | 1/20 (5)    |     |        |     |
| Age at symptom onset, y                   | 19c   | 6.9  | 1–16   | 5.1 |
| Age at diagnosis, y                       | 20    | 9.5  | 0–17   | 5.1 |

Abbreviation: AD = autosomal dominant.

a Mean number of units within the pathogenic D4Z4 repeat.
b The observed methylation minus the predicted methylation based on the D4Z4 repeat size.
c One child was from a family with facioscapulohumeral dystrophy and had been tested and diagnosed asymptotically.

Clinical Characteristics and Changes Over 2 Years

Muscle Function
At baseline, 15 of 20 patients (75%) had facial weakness; 2 years later, this had increased to 19 of 20 (95%). One child developed new scapular weakness during follow-up; a total of 6 of 15 patients had functional impairment of the shoulder or arm after 2 years. The mean MRC sum score of the group did not change between baseline and follow-up and varied between 60 and 70 (median 69, maximum score 70). The most frequently and most severely affected muscles were the trapezius and deltoid, but we also observed weakness of the biceps brachii, quadriceps, tibialis anterior, and gastrocnemius muscles. The mean clinical severity scale was stable over the 2-year period, despite variability between individuals. The mean FSHD clinical score increased from 2.1 to 2.8 (range 1–8, $p = 0.003$). The mean result of the 6-minute walk test and MFM showed a tendency toward an increase, but this did not reach statistical significance. Despite better test scores during follow-up, the 6-minute walk distance was on average still below the mean for sex and age (mean −1.3 SDs, range −2.7 to 0.2, SD 1.0), and a large variability between patients was observed. During follow-up, 11 of 20 patients (55%) were given physical therapy compared to 8 of 20 (40%) at baseline.

All muscle measurements are summarized in Table 2. Figure 2 displays the change in FSHD clinical score, MFM, the 6-minute walk test, and the quantitative muscle ultrasonography results of individual patients over the course of 2 years.

Muscle Imaging
Visual muscle ultrasound grading from 14 children at baseline and 9 children at follow-up showed a mean Heckmatt score of 1.4 and 1.6, respectively. Quantitative muscle ultrasound showed an increase in the mean z scores from 1.0 to 2.1 after 2 years (Table 2).

The visual grading correlated moderately to strongly with the quantitative score (baseline $r = 0.55 p = 0.02$, 2-year follow-up $r = 0.88, p = 0.001$). Figure 3A shows mean z score per muscle at the 2-year follow-up. All clinically affected muscles had an increased echogenicity except for the rectus abdominis. Three patients with an increased lumbar lordosis, considered to be a sign of weakness of the abdominal muscles in FSHD,3 did not have an increased echo intensity of the bilateral rectus abdominis. In addition, spinal and hip girdle muscle weakness contributes to postural instability in FSHD.33 These muscles were not evaluated by muscle ultrasound. Not all muscles with an increased echogenicity were clinically affected. We found a negative correlation between the mean quantitative gray scale analysis z score and the MRC sum score ($r = −0.74, p = 0.01$). Disease severity as measured by the FSHD clinical score demonstrated a trend toward a positive correlation with the mean z scores, but this did not reach statistical significance ($r = 0.51, p = 0.07$). One child showed a deterioration on muscle ultrasound with a z score increase of 1.06 but no clinical changes as measured by the FSHD clinical score. Figure 2D shows the changes in mean echogenicity in individual patients over the course of 2 years and demonstrates the variability in progression. Figure 3, B and C shows the distribution of the measured muscles at baseline and at follow-up by visual (Heckmatt score) and quantitative (z score) analysis.

Systemic Features
The presence of systemic features is shown in Table 3.

Vision
None of the children or parents reported vision loss. Despite normal visual acuity, retinal abnormalities were frequently observed on fundoscopy in 6 of 8 patients tested. Retinal changes consisted of mild tortuosity of the retinal arteries. The degree of retinal abnormalities did not change over 2 years. Coats disease with retinal detachment was not observed.

Hearing
One child reported hearing problems, and in 2 children (12.5%), a hearing loss was detected by audiometry. One
child had a conductive hearing deficit at baseline and a normal audiometry during follow-up. There were no patients with a newly diagnosed hearing deficit on follow-up.

**Cardiac Function**

No changes were observed in the frequency of abnormalities seen on the ECG. We found a right-axis deviation without clinical consequences in 1 child.

**Skeletal Features**

The percentage of children with lumbar hyperlordosis showed an increase from 35% to 53% over 2 years, but this change was not found to be statistically significant (Table 3). At baseline, a mild scoliotic posture was seen in 2 participants; however, this was no longer observed during the neurologic examination at the 2-year follow-up visit.

**Speaking and Swallowing**

Bulbar dysarthria was found in 25% of children on baseline, and its occurrence increased to 50% after the 2-year follow-up. All children with severe facial weakness had dysarthric speech. No changes were observed in the number of children with swallowing difficulties over time.

**Central Nervous System**

None of the participants had an intellectual disability or developmental impairment. One child was diagnosed with dyslexia, and 1 child had psychological support to deal with anxiety. Seventeen of 20 children (85%) attended regular education. None of the participants had or developed epilepsy.

**Pain, Fatigue, and Quality of Life**

Pain and fatigue were frequently reported, and their occurrence increased during the 2-year follow-up (75% and 70%, respectively). The location of pain was mainly the lower legs (53%) but also in the scapular region (27%) and lower back (20%). The pain was typically induced by muscle exercise and exertion. The NeuroQoL fatigue questionnaire showed that the children experienced more fatigue compared to healthy children, both at baseline and after the 2-year follow-up. The mean z score did not increase during follow-up (Table 3).

Quality of life was lower in children with FSHD compared to their healthy peers. Low scores were found especially in the domains of physical well-being and social acceptance. Overall, there was no difference in quality of life at baseline and after 2 years. On follow-up there was a significant improvement of the scores in the domains of parent relation and home life and of school environment. Details are shown in Table 3.
Differences in Early-Onset and Classic-Onset Type

The differences between children with an early-onset and classic-onset type are listed in Table 4. Two of the 20 children were too young to determine their specific phenotype and were excluded from this analysis. Patients with early-onset FSHD had a shorter number of units within the pathogenic D4Z4 repeat compared to classic-onset patients, as expected. Patients with an early-onset type were significantly younger than patients in the classic-onset group. Motor functioning after 2 years as measured by MRC sum score, FSHD score, clinical severity scale, and 6-minute walking test did not differ significantly between these subgroups. The age-adjusted clinical severity scale score was higher in the early-onset subgroup ($p = 0.02$). No significant difference in disease progression was found between the 2 onset types, with a mean change in FSHD clinical score of 0.63 points in the early-onset subgroup and 0.80 in the classic-onset subgroup ($p = 0.65$).

The visual and quantified muscle echogenicity tended to be higher in early-onset patients, but these changes did not reach statistical significance ($p = 0.11$ and $p = 0.07$). During follow-up, early-onset patients showed a bigger mean echogenicity change compared to classic-onset patients: in early-onset patients, an increase in mean z score of $2.0 \pm 1.4$ was seen compared to an increase of $0.1 \pm 0.5$ in the classic-onset subgroup ($p = 0.02$). The presence of pain, fatigue, and systemic features did not vary significantly between the 2 groups, although hearing abnormalities were observed only in the children with early-onset FSHD who had a D4Z4 repeat size of 2 units.

Determinants of Disease Progression

The rate of disease progression in 2 years as measured by the change of FSHD clinical score was not influenced by sex, current age, the number of D4Z4 repeats, or the mean echogenicity z score at baseline.

Discussion

This 2-year follow-up study in children with FSHD clinically showed mild progression of facial weakness, pain and fatigue, and lumbar hyperlordosis. This was reflected by an increase of disease severity as measured by the FSHD clinical score. As in adults, the disease course was variable. The muscle ultrasound abnormalities increased over 2 years and correlated with disease severity. The early-onset subtype had shorter D4Z4 repeats and a more severe phenotype as indicated by a higher age-corrected clinical severity score, more ultrasound abnormalities, and a higher prevalence of hearing and retinal abnormalities.
While the clinical scores indicated disease progression, the actual changes were limited. The FSHD clinical score, a sum score of 6 independent sections that describes strength and functionality of several muscle regions, was the only motor function outcome measurement that showed a significant deterioration. The children’s performance on functional exercise tests improved over 2 years, although it was still below the average performance for healthy peers. This most likely indicates the delayed but still ongoing motor development in children with FSHD compared to healthy children, which is in line with observations in patients with Duchenne muscular dystrophy that a delay in gross motor development is seen in one-third of children.34 Physical therapy and functional training might also improve functional capacities,35 but we did not find a relationship between attending physical therapy and improvement. This result might also point to limitations of the 6-minute walk test: its result is affected by the effort of the child. It gives us an estimate of the child’s functional status but does not assess the mechanism of exercise intolerance.36 The improvement found emphasizes the importance of natural history studies to ensure correct implementation of these outcome measurements in future clinical trials in children.

Pain, fatigue, and a decreased quality of life appear to be major problems in childhood, with a prevalence similar to that in the adult population.44-46 The current study reports an even higher proportion of children experiencing pain compared to another recent study on FSHD in childhood (83% vs 61%), despite the fewer patients with systemic features or loss of ambulation in our cohort.12 This difference could be explained by the different approach to exploring the presence of pain in the current study. We consciously asked for the presence of pain without the use of a pain scale, thus including all levels and patient perceptions of pain. Another possible
explanation is that the children in our cohort experience more pain secondary to exercise because of their maintained ambulation. The high prevalence of pain and fatigue in children with FSHD is a reason for concern, especially when we take into account the correlation between age and fatigue severity, with increasing age leading to a further deterioration of the quality of life. For children, this could have a substantial effect on participation in school and sports and their professional development. We suggest placing more focus on the management of pain and fatigue to ameliorate physical impairment and to optimize the child’s abilities.

A strength of this nationwide study is that it provided information about the full spectrum of FSHD in childhood. However, our study also has some limitations. First, the small number of participants makes it hard to determine correlations. This could be addressed by an even more encompassing international study or prospective data collection initiative for patients with FSHD. Second, FSHD in childhood appears to be a slowly progressive disease, so a 2-year follow-up is a short period to evaluate clinically relevant changes. A longer follow-up period is expected to provide further information about the natural history and support longer running treatment trials. Last, missing data were caused by

### Table 3: Systemic Features, Pain, Fatigue, and Quality of Life at Baseline and 2-Year Follow-up

| Systemic features, n/total (%) | No. of participants (at baseline—2-y follow-up) | Baseline scoring | 2-y Follow-up scoring | p Value |
|-------------------------------|-----------------------------------------------|------------------|----------------------|---------|
| **Hearing lossa**             | 18–16                                        | 3/18 (17)        | 2/16 (12.5)          | 1.00e   |
| **Vision loss**               | 20–12                                        | 0/18 (0)         | 0/12 (0)             | 1.00e   |
| **Retinal abnormalities**b    | 11–8                                         | 8/11 (73)        | 6/8 (75)             | 1.00e   |
| **Cardiac abnormalitiesc**   | 15–13                                        | 3/15 (20)        | 1/13 (8)             | 1.00e   |
| **Assisted ventilation**      | 20–20                                        | 0/20 (0)         | 0/20 (0)             | 1.00e   |
| **Lumbar hyperlordosis**     | 20–19                                        | 7/20 (35)        | 10/19 (53)           | 0.38e   |
| **Dysarthria**               | 20–20                                        | 5/20 (25)        | 10/20 (50)           | 0.13e   |
| **Swallowing difficultiesd** | 20–20                                        | 5/20 (25)        | 7/20 (35)            | 0.69e   |
| **Intellectual disability**  | 20–20                                        | 0/20 (0)         | 0/20 (0)             | 1.00e   |
| **Epilepsy**                 | 20–20                                        | 0/20 (0)         | 0/20 (0)             | 1.00e   |

| Pain and quality of life      |                                              |                  |                      |         |
|-------------------------------|-----------------------------------------------|------------------|----------------------|---------|
| **Pain, n/total (%)**         | 20–20                                        | 12/20 (60)       | 15/20 (75)           | 0.51e   |
| **Fatigue, n/total (%)**      | 20–20                                        | 12/20 (60)       | 14/20 (70)           | 0.69e   |
| **NeuroQoL 8-item fatigue bank score, mean SD** | 10–11 | 1.1 | 1.0 | 0.37e |
| **Kidscreen total score, mean SD** | 9–11 | −0.9 | −0.7 | 0.13e |
| **Kidscreen subdomain scores, mean SD** | 9–11 | −1.5 | −2.0 | 0.14e |
| **Physical well-being**       | 9–11                                        | −1.0             | −0.4                 | 0.20e   |
| **Psychological well-being**  | 9–11                                        | −0.7             | −0.4                 | 0.40f   |
| **Autonomy**                  | 9–11                                        | −0.8             | 0.2                  | 0.04e   |
| **Parent relation and home life** | 9–11 | −0.8 | 0.2 | 0.21f |
| **Financial resources**       | 9–11                                        | −0.7             | −0.3                 | 0.27e   |
| **Social support and peers**  | 9–11                                        | −0.8             | −0.9                 | 0.03e   |
| **School environment**        | 9–11                                        | −0.7             | −0.1                 |         |
| **Social acceptance**         | 9–11                                        | Insufficient numbers or replies | −1.6 |         |

a Based on audiometry abnormalities.

b Defined as mild or severe tortuosity of retinal arteries in at least 1 eye measured by fundoscopy.

c Based on ECG abnormalities.

Based on a score of ≤7 on the Neuromuscular Disease Swallowing Status Scale.

McNemar test.

Wilcoxon signed-rank test.

Paired t test.

Insufficient numbers or replies.

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COVID-19 and by the substantial proportion of patients or their parents who preferred not to visit the study center and participated by a phone interview and chart review only.

The slowly progressive course of childhood FSHD offers the opportunity for a future drug trial to delay or thwart this progression rate. According to our results, the most promising outcome measures for analyzing the effects of future therapeutics are the FSHD clinical score and muscle ultrasound. We expect to further define the natural history and course in a subsequent 5-year follow-up study of this cohort.

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**Table 4** Comparison of Motor Functioning and Systemic Complications Between Early-Onset and Classic Onset FSHD After 2-Year Follow-up

|                                      | Early onset (n = 8) | Classic onset (n = 10) | p Value     |
|--------------------------------------|--------------------|------------------------|-------------|
| Demographics                         |                    |                        |             |
| Age at baseline examination, y       | 8.3 ± 2.5          | 14.2 ± 2.9             | <0.001c     |
| Age at 2-y follow-up, y              | 10.3 ± 2.5         | 16.4 ± 3.0             | <0.001c     |
| Age at onset of symptoms, y          | 2.8 ± 2.2          | 10.5 ± 4.2             | 0.01b       |
| D4Z4, mean ± SD                      | 3.8 ± 2.2          | 6.4 ± 1.7              | 0.02b       |
| Motor functioning                    |                    |                        |             |
| MRC sum score (0–70), mean ± SD     | 66.7 ± 3.9         | 69.0 ± 1.4             | 0.17b       |
| CSS score (0–10), mean ± SD         | 3.1 ± 2.0          | 2.7 ± 1.8              | 0.70b       |
| Age-corrected CSS score (0–2000), mean ± SD | 288 ± 131        | 168 ± 112              | 0.02b       |
| FSHD clinical score (0–15), mean ± SD | 3.0 ± 1.9         | 2.5 ± 1.1              | 0.27b       |
| 6-min walk test score, No. of SDs, mean ± SD | −1.5 ± 1.1       | −1.0 ± 0.9             | 0.40c       |
| Motor Function Measure score, mean ± SD | 97.2 ± 3.9      | 100 ± 0                | 0.06b       |
| Muscle imaging                       |                    |                        |             |
| MUS score (z score), mean ± SDa     | 3.4 ± 2.5 (n = 4)  | 1.1 ± 0.4 (n = 5)      | 0.07c       |
| Heckmatt score (1–4), mean ± SD     | 1.9 ± 0.6 (n = 4)  | 1.4 ± 0.1 (n = 5)      | 0.11d       |
| ΔMUSf score (z score), mean ± SDa   | 2.0 ± 1.4 (n = 4)  | 0.1 ± 0.5 (n = 5)      | 0.02c       |
| Pain, fatigue, and systemic features, n/total (%) |          |                        |             |
| Pain                                 | 7/8 (87.5)         | 8/10 (80)              | 0.59a       |
| Fatigue                              | 7/8 (87.5)         | 7/10 (70)              | 0.38a       |
| Hearing abnormalities                | 2/7 (28.6)         | 0/9 (0)                | 0.18b       |
| Retinal abnormalities                | 3/3 (100)          | 3/5 (60)               | 0.36c       |
| Lumbar hyperlordosis                 | 4/8 (50)           | 5/10 (50)              | 0.68c       |
| Swallowing difficulties              | 2/8 (25)           | 4/10 (40)              | 0.43c       |
| Dysarthria                           | 5/8 (62.5)         | 4/10 (40)              | 0.32c       |

Abbreviations: FSHD = facioscapulohumeral dystrophy; CSS = clinical severity scale; MRC = Medical Research Council; MUS = muscle ultrasound.

a Fisher exact test, 1-sided p value.

b Mann-Whitney U test.

c Independent t test.

d Mean number of units within the pathogenic D4Z4 repeat.

* Mean z score of the quantified echogenicity per muscle measured by MUS.

Change in mean z score of the quantified echogenicity during 2 years of follow-up.
Disclosure

J.N. Dijkstra and R.J.M. Goselink report no disclosures relevant to the manuscript. N. van Alfen consults for Dynacure and provides editorial duties for Wiley Inc; all fees are paid to her employer. I.J.M. De Groot, M. Pelsma, N. Van der Stoop, and T. Theelen report no disclosures relevant to the manuscript. B.G.M. Van Engelen received grants from Global FSH, Stichting Spier for Spieren, Princes Beatrix Spierfonds, and Dutch FSHD foundation. N.C. Voermans and C.E. Erasmus report no disclosures relevant to the manuscript. Go to Neurology.org/N for full disclosures.

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Appendix

| Name               | Location                                                                 | Contribution                                                                 |
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| MD, PhD             | University Medical Centre, Nijmegen, the Netherlands                      | for content; major role in the acquisition of data; analysis or interpretation of data |
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|                    | Department of Rehabilitation, Donders Centre for Biomedical and Clinical | Study concept or design                                                      |
|                    | Sciences, Linköping University, Linköping, Sweden                        |                                                                             |
| Imelda J.M. de     | Department of Rehabilitation, Donders Centre for Biomedical and Clinical |                                                                             |
| Groot, MD, PhD     | Sciences, Linköping University, Linköping, Sweden                        |                                                                             |
|                    | Department of Neurology, Donders Centre of Neurosciences, Radboud         | Major role in the acquisition of data                                        |
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| Nienke van der     | Department of Clinical Genetics, Leiden University Medical Centre, the    | Major role in the acquisition of data; analysis or interpretation of data    |
| Stoop, PhD          | Netherlands                                                               |                                                                             |
| Thomas Theelen,     | Department of Ophthalmology, Radboud University Medical Centre, Nijmegen, | Major role in the acquisition of data                                       |
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| Baziel G.M. van     | Department of Neurology, Donders Centre of Neurosciences, Radboud         | Drafting/revision of the manuscript for content, including medical writing   |
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Appendix (continued)

| Name               | Location                                                                 | Contribution                                                                 |
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| Corrie E. Erasmus  | Department of Pediatric Neurology, Amalia Children’s Hospital, Radboud University Medical Centre, Nijmegen, the Netherlands | Drafting/revision of the manuscript for content, including medical writing for content; study concept or design; analysis or interpretation of data |
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