Fatigue and daytime sleepiness rating scales in myotonic dystrophy: a study of reliability

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OBJECTIVES: To assess the reliability of the Epworth Sleepiness Scale (ESS), Daytime Sleepiness Scale (DSS), Chalder Fatigue Scale (CFS), and Krupp's Fatigue Severity Scale (KFSS) in patients with myotonic dystrophy type 1 (DM1).

METHODS: In total, 27 patients with DM1 were administered the questionnaires on two occasions, with a 2 week interval. Internal consistency and test-retest reliability were measured using intraclass correlation coefficients (ICCs), and Cronbach's α, Cohen's κ, and Goodman-Kruskal's γ coefficients.

RESULTS: Internal consistency of the CFS and KFSS were adequate (α>0.70) but that of the ESS was weak (α = 0.24). Both daytime sleepiness and fatigue rating scales showed significant test-retest reliability. Test retest reliability for individual items revealed inconsistencies for some ESS and CFS items.

CONCLUSIONS: Reliability of the CFS, DSS, and KFSS was high, allowing their use for individual patients with DM1, but that of the ESS was lower, rendering its current usage in DM1 questionable. Fatigue rating scales such as the KFSS, which are based on the behavioral consequences of fatigue, may constitute a more accurate and comprehensive measure of fatigue severity in the DM1 population.

Myotonic dystrophy type 1 (DM1), an autosomal dominant disorder, is the most common adult form of muscular dystrophy. DM1 results from an unstable CTG repeat expansion in the 3' untranslated region of the myotonin kinase gene at 19q13.3.2 DM1 is not only a muscle disease but a multisystemic disorder, including impairment of the central nervous system (CNS).3 Daytime sleepiness has been referred as the most frequent non-muscular symptom of the disorder5 and has repeatedly been reported as one of its earliest symptoms.5–8 Available evidence suggests that daytime sleepiness is primarily the result of a CNS dysfunction.9 Fatigue is yet another prominent complaint of patients with DM1.1 A recent study reported no difference in fatigue severity between DM1 patients with and without daytime sleepiness. Nevertheless, both these groups exhibited abnormal fatigue levels, thus advocating simultaneous use of daytime sleepiness and fatigue outcome measures in studies of patients with DM1.

In view of the deleterious impact of these latter symptoms on health and social function,6 their accurate documentation has relevant implications for the quality of life of patients with DM1. There are tests available, such as the multiple sleep latency test,10 which quantify objective daytime sleepiness, but these techniques require sophisticated equipment, and are expensive and time consuming to perform. As for fatigue, there is no such “gold standard” measuring tool; however, clinicians frequently rely on rating scales to guide treatment decisions and measure progress regarding these symptoms. Even though it is well recognised that the soundness of clinical evidence substantially depends upon the applicability of an instrument to the population of interest,12 no daytime sleepiness or fatigue rating scale has, to our knowledge, been thoroughly assessed for reliability in patients with DM1. The present study was thus conducted in order to document the reliability of the Epworth Sleepiness Scale (ESS),13 the Daytime Sleepiness Scale (DSS) specifically devised for DM1,14 the Chalder Fatigue Scale (CFS),15 and the Krupp's Fatigue Severity Scale (KFSS)16 in patients with DM1.

METHODS

Sample
A list of 30 individuals was randomly selected from a registry of adult patients with classic DM1 attending the Saguenay Neuromuscular Clinic (Québec, Canada), from which 27 patients with DM1 (11 men, 16 women) aged 37–66 years (mean 49.6) agreed to participate in the study (participation rate 90%). Each patient met the diagnostic criteria for definite DM1 or for an obligate carrier.17 Molecular confirmation of the diagnosis was available for 24 patients (88.9%), but a founder effect had already been demonstrated in this population.18–19 CTG repeat classes were: <200 (n = 3), 200–400 (n = 2), 401–850 (n = 5), 851–1100 (n = 4), 1101–1500 (n = 6), and >1500 (n = 4). All individuals were examined by a neurologist (JM) and had their muscular impairment categorised as mild (grades 1 and 2, no or minimal signs of muscular impairment; n = 4), moderate (grade 3, distal weakness; n = 7) or severe (grades 4 and 5, mild to severe proximal weakness; n = 16), based on our previously published muscular impairment rating scale.20 All patients completed the four questionnaires on two occasions, with a 2 week interval between them. This study was approved by the ethics committee Complexe hospitalier de la Sagamie.

Measures

Daytime sleepiness rating scales
The ESS was developed to measure the general level of sleepiness, conceptually defined as sleep propensity. It consists of eight questions asking the subject to rate their chance of falling asleep in situations commonly encountered in daily life. The unidimensional aspect of this scale has been confirmed by factor analysis.21 An evaluation in normal subjects suggests that the measure is stable over time.22
The DSS consists of five items derived from the Stanford University Sleep Questionnaire and Assessment of Wakefulness. Principal component analysis in 157 patients with DM1 revealed that the DSS measured a single factor. Cronbach’s α reliability coefficients for the five items was 0.72, reflecting a level of internal consistency suitable for application in both research and practice.

Fatigue rating scales

The CFS was originally used in a hospital based case-control study and further refined by Chalder et al on a sample of 374 primary care patients. This 11 item scale measures physical and mental fatigue. The validity and reliability of the CFS has been examined in general practice attendees and in patients with chronic fatigue syndrome. Principal components analyses performed by Chalder et al supported the notion of a two factor scale for fatigue (physical and mental).

The KFSS assesses the effect of fatigue on daily activities. Its nine items were selected to identify common features of fatigue in both multiple sclerosis (MS) and systemic lupus erythematosus (SLE). In the initial validation study, individuals with MS and SLE were compared with healthy adults. Internal consistency of the KFSS was high for both illness groups. The KFSS demonstrated adequate concurrent validity by clearly distinguishing between patients and controls, and criterion related validity by a moderate correlation with Centre for Epidemiologic Studies Depression Scale scores in the MS, SLE, and control groups.

Statistical analysis

Internal consistency (the degree of item interrelatedness), was assessed with Cronbach’s α reliability coefficients from the first questionnaire completion. The test retest reliability (the estimate of the instrument’s reproducibility over time assuming that no change in condition has taken place) was assessed by intraclass correlation coefficients (ICCs). Finally, test retest reliability for individual items (that is, the measure of association for a given item computed from both completions) was assessed with Cohen’s κ for nominal measures (for the CFS, of which scoring is bimodal) and with Goodman-Kruskal’s γ coefficient for ordinal measures (ESS, DSS, KFSS). Significance testing was two sided, with α set at 0.05.

RESULTS

Fatigue and daytime sleepiness levels

The mean (SD) scores were 8.6 (3.5) for ESS, 5.7 (3.5) for DSS, 4.9 (2.4) for CFS, and 5.1 (1.6) for KFSS.

Reliability

Cronbach’s α for the ESS was 0.24, and ICC 0.68 (p<0.001). Goodman-Kruskal’s γ was significant for ESS items 1, 2, 4, 5, and 7 (range 0.54–0.90, p<0.05). However, it was not significant for ESS item 8 (In a car, while stopped for a few minutes in traffic). In particular, all patients with DM1 except one indicated that they would “never fall asleep” in this situation on the second ESS completion. In addition, Goodman-Kruskal’s γ could not be computed for ESS items 3 and 6 because all patients with DM1 answered that they would “never fall asleep” on the first completion of ESS item 3, and on both completions of ESS item 6.

The ICC of the DSS was 0.82 (p<0.001), and Goodman-Kruskal’s γ was significant for all DSS items (range 0.60–0.86, p<0.05).

Cronbach’s α reliability coefficient of the CFS was 0.70, and ICC 0.81 (p<0.001). Cohen’s γ could not be computed for CFS item 11 (How is your memory?) because all patients with DM1 answered either “better than usual” or “no worse than usual” on the second CFS completion. Cohen’s κ was significant for all other CFS items (range 0.40–0.76, p<0.05) except item 3 (Do you feel sleepy or drowsy?). Cronbach’s α reliability coefficient of the KFSS was 0.86, and ICC 0.88 (p<0.001). Goodman-Kruskal’s γ was significant for all KFSS items (range 0.55–0.88, p<0.05).

DISCUSSION

The most widely used instrument to assess daytime sleepiness is probably the ESS, but its measurement properties are questionable, particularly its reliability in clinical samples. In DM1, the ESS was used as the primary efficacy variable or as one primary outcome measure in the most recent clinical trials of modafinil for patients with daytime sleepiness. In DM1, the ESS was used as the primary efficacy variable or as one primary outcome measure in the most recent clinical trials of modafinil for patients with daytime sleepiness. 

In contrast with previous reports that documented an acceptable internal consistency for the ESS in both controls and patients with various sleep disorders, the present results show a weak internal consistency for the ESS in patients with DM1, indicating poor correlation between ESS items. Some insights regarding this discrepancy are provided by the test retest reliability analyses that were performed on individual ESS items. Indeed, Goodman-Kruskal’s γ coefficients could not be computed or were not significant for three of the eight "real life" situations depicted by the ESS. Particularly, patients with DM1 estimated that they would "never doze" when "sitting and talking to someone", "in a car, while stopped for a few minutes in the traffic", or "sitting inactive in public place (for example, a theatre or a meeting)". The estimation of sleep propensity by patients with DM1 may have been biased by the fact that severely affected patients actually experience the latter two situations infrequently, if ever. It must be noted that the dependence of the DM1 population upon social welfare for their income may also restrain such activities. More importantly, falling asleep in such situations is uncharacteristic of patients with DM1.

Indeed, it has long been established that the tendency to sleep in DM1 does not occur during activity. It is probable that reporting no chance of dozing on some ESS items while reporting slight to high chances of dozing on other items diminished the consistency of intrasubject responses and intersubject variability, and produced low homogeneity of variances among items, consequently yielding a low Cronbach’s α. On the other hand, the present results indicate significant test retest reliability of the ESS in DM1, consistent with that previously observed in 87 medical students who filled the ESS twice with a 5 month interval.

The higher test retest reliability observed for the DSS probably ensues from the fact that it was compiled from answers to questions relating to sleep and sleepiness given by patients with DM1. The DSS was found to correlate with the extent of muscular impairment, and its items are consistent with the clinical features most commonly noted in association with DM1 related daytime sleepiness, such as daytime napping and sleepiness when attention is not being held. However, its appropriateness as an evaluation instrument that measures change over time must be assessed.

The KFSS was conceived as a fatigue function measure that integrates dimension of fatigue intensity and functional outcomes associated with fatigue, in contrast to the CFS, which measures fatigue intensity alone. It was accordingly suggested that the KFSS is a more effective measure of disability related fatigue in diseases that specify a state of marked functional disability. Hence, patients with DM1, who exhibit significant limitations in daily living activities, may more readily comprehend rating scales that specifically assess the effect of fatigue on daily activities, partially

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explaining the better reliability of the KFSS over the CFS. A second reason may pertain to the difference in samples used in the development of these scales. The KFSS was evaluated using patients with MS, a chronic disease for which modafinil has been shown to significantly decrease fatigue levels, as has also been shown in DM1.9 Regarding the CFS, the poor reproducibility of CFS item 3 (Do you feel sleepy or drowsy?) may relate to our observation that patients with DM1 may not present complaints of daytime sleepiness on a given day, while on a different day assessment, it may represent a troublesome symptom. Consequently, a more reliable clinical assessment of daytime sleepiness is likely to be provided by inquiring about the effect of daytime sleepiness on the daily activities or behaviours of patients with DM1. With regards to CFS item 11, it is a matter of debate as to whether patients with DM1 show normal22 or impaired memory function.82 Their answers could either reflect a normal memory functioning or an incapacity to report a putative memory deficit. This latter CFS item, as well as those relating to sleepiness or apathy, may address constructs other than fatigue per se, perhaps further influencing the slightly higher reliability of the KFSS with respect to the CFS.

CONCLUSION
This first study to assess the reliability of daytime sleepiness and fatigue rating scales in DM1 revealed acceptable internal consistency and reproducibility for the CFS, DSS, and KFSS, suggesting that they are suitable for application in both research and practice in this patient population. Our results, however, indicate a lesser reliability for the ESS, probably due to the poor correlation between some of its items and the habitual activities and/or clinical realities of patients with DM1, rendering its current usage in DM1 questionable. In addition, other studies must determine the construct validity of daytime sleepiness and fatigue rating scales in this disease. Although daytime sleepiness and fatigue rating scales obviously provide invaluable information to clinicians and researchers, there should also be a concern about obtaining objective measurement of sleep tendency17 and exploring potential physiological correlates of fatigue.41 Given that virtually all systems may be affected in DM1, clarifying the mechanisms underlying daytime sleepiness and fatigue entails considering such potential aetiological factors as the cerebral, genetic, metabolic, muscular, psychological, and respiratory abnormalities that are characteristic of the disorder.

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