Short Report

Single fibre EMG studies in chronic fatigue syndrome: a reappraisal

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Abstract
Single fibre EMG studies were carried out on the right extensor digitorum communis muscle in 30 subjects with chronic fatigue syndrome and in 30 age and sex matched controls. Abnormal jitter was seen in five patients with chronic fatigue syndrome. Slight but significant differences between the mean consecutive differences in the remainder of the chronic fatigue subjects and the control subjects were recorded. Overall the differences were so minor that it seems unlikely that a disturbance of neuromuscular function as reflected by jitter measurement has a pathogenetic role. It is suggested that the increased jitter seen may be explained by the effects of the variability of motor unit firing rates on the myogenic component of the jitter.

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Chronic fatigue, often with muscle pain, is a prominent complaint in western countries and often a definable physical or psychiatric basis cannot be established. Idiopathic chronic fatigue syndrome has been defined by a Centre for Disease Control (CDC) committee1 as disabling chronic fatigue of over six months’ duration in the absence of a definable physical or psychiatric basis and a number of minor criteria are also required to establish the diagnosis. A great deal has been published about chronic fatigue syndrome in the immunological,2 virological,3 and psychiatric fields4 but its aetiology is still poorly understood. The suggestion that abnormal muscle metabolism may be at fault5 is not supported by other studies6,7 and muscle exercise performance8-10 is normal. A finding of abnormal single fibre jitter in 75% of cases in one study is perhaps the only remaining suggestion of a peripheral neuromuscular abnormality in this syndrome.11 The purpose of the present study was to re-examine this question in a cohort of patients with chronic fatigue syndrome with a carefully selected control group.

Methods
Thirty patients were admitted to the study, all of whom fitted CDC criteria for diagnosis. All patients had moderate or severe fatigue and 92% had moderate or severe myalgia. All had symptoms for over 6 months and 54% had symptoms for over 5 years. Although a history of antecedent infection was given in 40% of patients, serological confirmation was usually not available.

After informed consent was obtained, single fibre jitter studies were performed in the right extensor digitorum communis muscle. A standard Medelec SFEMG needle was used with a recording surface of 25 µm diameter. Recordings were made with a Medelec MS6 electromyograph. The traces were digitised and transferred on line by a Medelec CI6 interface to an Epson PC (IBM compatible) computer for subsequent analysis. Eighteen to 20 recordings of each pair were acquired and stored in this way. At least 10 potential pairs were analysed. The recording was made with voluntary activation of the muscle, with the subject maintaining a slight, steady contraction.

The interpotential interval (IPI) and mean consecutive differences (MCD) were calculated. Thirty age and sex matched control subjects were studied in parallel. The mean age of the patient group was 39 (range 21 to 58) years and the mean age of the control group was 34 (range 21 to 55) years.

Results
The mean MCD of the controls was 25.7 (SD 4.3) µs. The jitter was considered abnormal if it was greater than 3 SD above the control mean, or if more than two pairs had jitter in excess of 55 µs or if blocking was seen. Using these criteria five subjects with chronic fatigue syndrome had abnormal jitter. Blocking was seen in only two patients. One of these had 5% blocking in only 1/10 pairs studied and the mean jitter was increased (50-3 µs). The other had 5% blocking in 2/10 pairs and the mean jitter was 49.3 µs. Three other subjects with chronic fatigue syndrome had abnormal jitter (each with 2/10 pairs greater than 55 µs, the mean jitter being 33.3, 39.2, and 43.0 µs). The cases with abnormal results had no clinical distinguishing features from other cases. The mean MCD of these five patients was 43.0 (SD 7.1) µs. The mean MCD of the remaining 25 subjects with chronic fatigue syndrome was 27.8 (SD
therefore be increased than intervals recorded interpotential jitter; distribution is less than axon transmission is not.

Discussion
This study has shown a small subgroup of patients with chronic fatigue syndrome who have abnormal single fibre jitter without any other evidence of motor unit abnormality. All of our patients with abnormal jitter had variable innervation rates associated with abnormal recruitment of motor units, which is a common finding in chronic fatigue syndromes. This finding was also common among the other patients. There was a slightly higher jitter in the patient with the control group. The amount of this difference is very small and not clinically important.

Fatigue due to neuromuscular transmission defects depends upon the presence of blocking and this was not present in most patients.

The reasons for the discrepancy between our findings and those of Jamal and Hansen, who found 75% of their patients had abnormal studies, is not clear. Comparison of data of this type with an age and sex matched control group, as in this study, rather than with standard laboratory reference ranges, is clearly essential, however.

Variability in jitter is usually due largely to neuromuscular transmission at the motor end plates. Variability of conduction in the terminal axon branches or variability of muscle fibre propagation may also contribute significantly to jitter.

Trontelj et al10 have shown that the contribution to jitter from muscle fibre propagation (myogenic jitter) is normally small and with more or less constant rates of stimulation of axons, is less than 5 µs. Increased myogenic jitter can potentially be produced, however, by long mean interpotential intervals and with erratic discharge rates. In this study all interpotential intervals recorded were less than 2.5 ms and therefore variability in the interpotential intervals is not likely to be an important factor.

Erratic rates of stimulation of a muscle fibre will under normal conditions significantly change the velocity of propagation and this variation does occur at physiological rates of stimulation. Abnormalities of recruitment may therefore be sufficient to contribute to increased myogenic jitter; this may explain the abnormal studies reported here and also the slightly higher jitter seen in the chronic fatigue syndrome group. Also, in this study, calculations were made on a small number of motor unit discharges. This may have resulted in artefactually increasing the effects of erratic recruitment on the myogenic component of the jitter.

First, stimulation of muscle fibres and stimulated SFEMG studies would eliminate the contribution of the effects of variation in rates of transmission in the axons and across the neuromuscular synapses and may help to localise the site of origin of the abnormal jitter. The cause of the blocking seen in two patients is not clear. This was, however, only a minor finding present in very few pairs and it is probably of no clinical importance.

Conclusion
In this study only five out of 30 patients with chronic fatigue syndrome had an abnormal jitter and the mean of the jitter in the remaining patients was only slightly higher than in the controls. These abnormalities are considerably less than in a previously reported study. It is suggested that the increased jitter may be explained by variable rates of innervation producing an unusually large component of myogenic jitter rather than being related to primary effects of the disorder at the neuromuscular junction, terminal axon, or muscle membrane. The low incidence of abnormal jitter and the minor abnormalities seen suggest that the increased jitter is not due to an important pathogenetic factor in chronic fatigue syndrome. In our hands, SFEMG is not a useful test in chronic fatigue syndrome, other than to exclude other conditions such as myasthenia gravis.

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