SHORT REPORT

Contractile properties of lower leg muscles are normal in Parkinson’s disease

A Hufschmidt, K Stark, C H Lücking

Abstract

Contraction parameters (time-to-peak and half relaxation time), responses to short tetani and resistance to stretch were studied in the lower leg muscles of Parkinsonian patients and in age-matched controls. It was possible to distinguish between muscle groups of different fibre type composition in normal subjects on the basis of their contraction and relaxation velocities. These parameters, however, failed to show any abnormalities in the patient group. The only abnormal finding in Parkinsonian subjects was an increased resistance to passive stretch under static conditions, presumably elastic in origin. The results are evidence against a contribution of altered contractile properties to muscular rigidity in Parkinsonism.

The pathophysiology of rigidity is still controversial. Among the mechanisms thought to contribute to the tone increase in Parkinson’s disease (PD) are: enhanced flexor reflexes with responses appearing also in antagonist muscles,12 disinhibited tonic stretch reflexes3 as well as abnormal long-loop reflexes.4,6 Current theory agrees that the lower motor neuron is the final common pathway over which rigidity is expressed. Electromyographic recordings of lower leg muscles in Parkinsonian and spastic patients during walking have, however, failed to show an increase in activity in the gastrocnemius muscle while it was stretched during the swing phase.7 In the upper extremity of Parkinsonian subjects, the neutral angle of the elbow joint has been shown to be smaller than in normal controls, and stiffness is enhanced in the absence of EMG activity.8 As a result, it has been suggested that both rigidity and spasticity arise partly from altered mechanical properties of the muscles. Lower leg muscles of patients with long-standing spasticity have indeed been shown to have some increase in plastic stiffness.9 The actual contribution, however, of this change to the overall increase in tone, as well as to the disability experienced by the patient, remains open.

The influence of neuronal discharge patterns on the differentiation of skeletal muscle is well established and has been demonstrated after spinal transections10 cross-reinnervation of fast and slow units11-14 and long-term stimulation.15-17 Thus a relationship between altered central innervation patterns and changes in muscular properties is conceivable.

Muscles of spastic patients have delayed contraction times.16-20 As yet, there are no comparable data for PD. Morphological and histochemical studies of this condition have revealed an atrophy of fast (Type II) and hypertrophy of slow (Type I) fibres.21 In addition, ultrastructural abnormalities have been shown, in particular, defects of the outer mitochondrial membrane and disarrangement of the contractile filaments.22 It is not, however, known whether these changes affect muscle tone.

Direct measurements of muscle stiffness using imposed movements are difficult in Parkinsonism, due to shortening reactions, which are more frequent in extrapyramidal disorders,23 and which may lead to gross under-estimation of the passive resistance to stretch. In this study, responses to single twitches and to brief tetanic trains were recorded under resting conditions and during passive movements of the ankle. We have failed to demonstrate any abnormality in the contractile properties of lower leg muscles in Parkinsonian patients.

Methods

Subjects

The experiments were performed on 16 Parkinsonian patients (mean age 67.8 years, male/female = 11/5) and 18 controls (mean age 63.7 years, m/f = 9/9), who had no history or clinical signs of motor abnormalities. For statistical comparison of the two groups, the two youngest normal subjects were eliminated from the control group (thus n = 16, mean age 65.4 years, m/f = 9/7). The subjects gave their informed consent to the procedures involved in the study. The patients were Grade 3 and 4 according to the Hoehn and Yahr scale. All were receiving anti-Parkinsonian treatment, including levodopa.

Experimental procedure

The subjects were seated comfortably in a chair and were instructed to relax their legs as completely as possible. One foot was strapped to a pedal which could either be locked into place or moved by a torque motor. The axis of the pedal was aligned approximately with the subject’s ankle. The twitch force (later converted to ankle torque) and the resisting torque during passive movement were measured by strain-gauges registering a minimal deformation of the pedal along a breaking-line. Tension will therefore be given
as ankle torque in this paper. Assuming a relationship between force output and cross-sectional area, all tension values were normalised to a standard calf with a cross-sectional area corresponding to a maximum circumference of 35 cm. The torque and position signals were digitised and stored by a Nicolet 1070 averager and then passed on to an Apple II+ computer for further evaluation. The subject's lower leg was positioned so that the tibial edge was perpendicular and at a right angle to the sole of the foot. Muscles were stimulated with brief (0.1–1 ms), current-stabilised square-wave pulses of up to 25 mA at the motor point, with the second electrode positioned over the distal tendon. For the triceps surae muscle, stimuli were applied to one head of the gastrocnemius muscle. There was often also a twitch of the soleus muscle elicited by the indifferent electrode.

Experiments 1 and 2, involving isometric twitches and tetani, were performed on the tibialis anterior and the triceps surae muscle:

1 **Isometric twitches** at near-threshold intensity ("weak twitch"); near the tolerance threshold intensity ("strong twitch"), and at an intermediate intensity ("intermediate twitch"); stimulating current mid-way between the previous two. The tolerance threshold, as a rule, was reached at two to three times motor threshold intensity. The intermediate stimulus intensity was maintained for the subsequent tetanic stimulation experiments. Recorded parameters were: peak tension (PT), time-to-peak (TTP), and half-relaxation time (HRT).

2 **Isometric tetani** were applied as brief trains (5 pulses) at 10, 20, 40, 60 and 100 Hz. Peak tension (PT) was recorded. The twitch/tetanus ratio was computed as the quotient between peak tensions for the "intermediate" twitch and the 100 Hz tetanus, the stimulus intensity being identical in the two conditions.

Experiments 3 and 4 involved controlled stretches:

3 **Elastic stiffness**. The foot was slowly (6–6 deg/s) dorsiflexed by the pedal and then maintained in a 10 degree dorsiflexed position for 4.7 s. Elastic stiffness was calculated from the difference in ankle torque at the starting position and at the dorsiflexed position immediately before the return of the pedal, that is, after termination of phasic stretch responses.

4 **Stiffness of stimulated muscle**. The influence of low-level activation on the resistance to stretch was initiated by 10 Hz tetanic stimulation (5 pulses) of the triceps surae muscle while it was being slowly stretched at a rate of 16 deg/s of ankle dorsiflexion. The first impulse of the train was delivered 100 ms after onset of the passive movement. As a control condition, the same stretch was imposed without stimulation. Tension increase was measured as the difference between the peak tension during stretch under tetanic stimulation and (in the control runs without stimulation) the resting tension at the point where peak tension had occurred in the previous condition. In the tetanically stimulated muscle, the resistance to stretch (stiffness) was compared with the active tension by the stiffness ratio

\[ R_{\text{stiff}} = \frac{\text{tensile tension of stretched muscle}}{\text{tensile tension of resting muscle}} \]

which is invariably > 1, reflecting the fact that the tetanically stimulated muscle resists stretch with a force far greater than its contractile force.

**Trials with superimposed shortening reactions could be identified by irregular deformations of the length-tension curve and were discarded.**

**Results**

**Normal subjects**

The absolute twitch force, in our experiments, was a poorly controlled parameter and varied considerably between subjects. We must begin therefore by showing that the kinetic parameters are not dependant on twitch amplitude. This has been done by comparing TTP and HRT at the three different levels of stimulus intensity. Table 1 shows that there are no systematic changes of contraction or relaxation velocity with changes in stimulation intensity.

Both parameters of twitch duration (TTP and HRT) were significantly shorter in the tibialis anterior compared with the triceps surae (table 1). This reflects the difference in fibre type composition: the ratio of Type 1: Type II fibres is 46:54 for tibialis anterior, but between 82:18 (gastrocnemius, lateral head) and 89:11 (soleus) for the triceps surae.24 The difference was not, however, reflected in the tetanic PT at increasing frequencies.

There was a significant co-variation (r = 0.75; p < 0.001) of TTP and HRT for strong twitches in the tibialis anterior, suggesting that contraction velocities are scattered over a certain range within the control group. Correlations for weak and intermediate twitches as well as for the triceps surae (all twitches) were non-significant.

**Table 1** Contraction parameters of lower leg muscles in elderly normal subjects (n = 18). PT: peak tension, TTP: time-to-peak, HRT: half-relaxation time.

<table>
<thead>
<tr>
<th></th>
<th>M tibialis ant. (mean ± SD)</th>
<th>M triceps surae (mean ± SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weak twitch</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PT (Nm)</td>
<td>0.169 (0.115)</td>
<td>0.159 (0.134)</td>
</tr>
<tr>
<td>TTP (Nm)</td>
<td>98.7 (12.6)</td>
<td>113.7 (26.5)</td>
</tr>
<tr>
<td>HRT (ms)</td>
<td>103.2 (25.2)</td>
<td>135.0 (54.1)</td>
</tr>
<tr>
<td>Intermediate twitch</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PT (Nm)</td>
<td>0.335 (0.163)</td>
<td>0.378 (0.266)</td>
</tr>
<tr>
<td>TTP (Nm)</td>
<td>90.2 (10.7)</td>
<td>118.3 (21.4)</td>
</tr>
<tr>
<td>HRT (ms)</td>
<td>98.1 (23.0)</td>
<td>146.6 (32.6)</td>
</tr>
<tr>
<td>Strong twitch</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PT (Nm)</td>
<td>0.504 (0.280)</td>
<td>0.595 (0.476)</td>
</tr>
<tr>
<td>TTP (Nm)</td>
<td>90.0 (12.5)</td>
<td>114.7 (20.1)</td>
</tr>
<tr>
<td>HRT (ms)</td>
<td>92.7 (25.3)</td>
<td>139.1 (31.7)</td>
</tr>
</tbody>
</table>

Peak tetanic tension

| 100 Hz (%)     | 100                           | 100                           |
| 60 Hz (%)      | 100-7 (26.1)                 | 96.2 (14.8)                  |
| 40 Hz (%)      | 93.4 (18.3)                  | 92.9 (20.3)                  |
| 20 Hz (%)      | 93.0 (15.1)                  | 98.1 (20.2)                  |
| 10 Hz (%)      | 70.3 (21.2)                  | 82.2 (18.3)                  |
| Twitch/tetanus | 0.22-2 (0.01)                | 0.24-2 (0.13)                |

*p < 0.01 (Wilcoxon test for paired samples), ns, not significant.
Patients with Parkinson's disease

The patient group did not show any slowing of TTP or HRT compared with the control group (table 2). The curves relating the frequency of tetanic stimulation to tetanic PT lay as close together as in normal subjects and did not saturate earlier, as might have been expected in slower muscles. The only abnormality found in the patient group was a twofold increase in the passive resistance to stretch under static conditions (table 3).

Discussion

Two mechanisms are conceivable by which altered muscle properties may contribute to rigidity. Firstly, the elastic stiffness of muscle and joint capsule may be enhanced by fibrosis. Secondly, a transformation of the contractile apparatus may result in a disproportional increase in stiffness at low rates of stimulation. It would be expected that the latter abnormality would be seen to some degree when the fibre type composition of a muscle is shifted towards slow (Type 1) fibres, with a lower tetanic fusion frequency. This should manifest itself as a slowing of contraction and relaxation of the whole muscle and as a large increase in stiffness when the muscle is stretched during low rate tetanic stimulation.

This study failed to produce any evidence for a slowing of intrinsic muscular contraction in the lower leg muscles of Parkinsonian patients. It is unlikely that this failure is attributable to a lack of sensitivity of our recording equipment, as the method was able to show differences in contraction kinetics between the tibialis anterior and triceps surae muscles in normal subjects. Furthermore, we found an inter-individual co-variation between TTP and HRT for the tibialis anterior which obviously reflects individual differences in contraction velocity within the control group. If the method is sensitive enough to detect these differences, it should also be able to record changes related to rigidity. The quantification of peak tension during short tetani was found to be of little use in this context and did not even reveal noticeable differences between muscle groups of different fibre type composition in normal subjects. To spare our subjects from extremely painful sustained tetanic stimulation, we confined ourselves to very brief trains of five stimuli. This is probably insufficient to reach a steady state of tetanic contraction and therefore inadequate as a means of detecting differences in tetanic fusion frequency.

Our only abnormal finding in Parkinsonian patients was an increase in the resistance to passive dorsiflexion. A similar finding has been made in the upper extremities of Parkinsonian subjects, who exhibited a shift of the resting position of the elbow towards flexion and an increased resistance to stretch in the absence of EMG activity. In the latter case, the authors were uncertain whether to attribute these changes to the elastic structures or the contractile apparatus. In this study, we did not record EMG activity and therefore cannot exclude a neural contribution to the increased stiffness observed. Even so, our experiments did not produce any evidence for associated changes of active contraction kinetics and would therefore indirectly support the first alternative. This increase in passive elastic stiffness may be a consequence of a failure to use the whole range of joint excursions during daily life of Parkinsonian patients.

This work was supported by the Deutsche Forschungsgemeinschaft (SFB 325). The authors are grateful to Drs G Deuschl, S Fellows and J Noth for reviewing the manuscript.

---


17 Pette D, Müller W, Leisner E, Urbova G. Time dependant effects on contractile properties, fibre population, myosin light chains and enzymes of energy metabolism in intermittenly and continuously stimulated fast twitch muscles of the rabbit. Pflügers Arch 1976;364:103–12.


Contractile properties of lower leg muscles are normal in Parkinson's disease.

A Hufschmidt, K Stark and C H Lücking

J Neural Neurosurg Psychiatry 1991 54: 457-460
doi: 10.1136/jnnp.54.5.457

Updated information and services can be found at:
http://jnnp.bmj.com/content/54/5/457

These include:
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/