Reticular reflex myoclonus: a physiological type of human post-hypoxic myoclonus

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SUMMARY A patient with post-hypoxic myoclonus, sensitive to therapy with 5-hydroxytryptophan and clonazepam, was subjected to detailed electrophysiological investigation. Brief generalised jerks followed the critical stimulus of muscle stretch. The electroencephalogram showed generalised spikes that were associated with, but not time locked to, the myoclonus. The cranial nerve nuclei were activated upward. Analysis of the findings suggests that the mechanism of the myoclonus is hyperactivity of a reflex mediated in the reticular formation of the medulla oblongata.

Myoclonus is a complicated and ill-understood phenomenon associated with a variety of electrophysiological abnormalities (Halliday, 1967) and caused by many pathological conditions (Marsden and Parkes, 1973). No one pathophysiological mechanism can account for all the types of myoclonus seen in clinical practice. In some patients the myoclonic jerks occur spontaneously while the subject is at rest. In others they appear only on action and in such patients myoclonic jerks may be triggered by sensory stimuli. Such reflex myoclonus is seen in response to visual or auditory stimuli. Occasionally a tendon tap will elicit a myoclonic response, in which case the critical stimulus may be muscle stretch, a joint movement, or cutaneous stimulation. In one particularly well-studied case under the care of Dr A. Carmichael at the National Hospital, Queen Square, careful clinical observation by Dr Alan Norton established the critical stimulus to be muscle stretch. This patient was found to have giant cortical potentials evoked by electrical stimulation of peripheral nerves (Dawson, 1947), stimuli which also caused myoclonic jerks.

Recently, Sutton and Mayer (1974) have described a patient with focal reflex myoclonus due to cerebral vascular disease with infarction in the left cerebral hemisphere. She suffered from intermittent jerking of the right fingers and hand, and to a lesser extent the right face, jaw, tongue, and foot. The onset was acute at the age of 66 years and was associated initially with a mild right hemiparesis and hemisensory loss, as well as speech disturbance and hemianopia. When studied seven years after the onset, it was noted that the myoclonic jerks of the right limbs increased on voluntary action and were precipitated by eliciting tendon jerks. Electrophysiological investigation revealed that stimulation of the median nerve at the wrist or the posterior tibial nerve at the ankle evoked a late myoclonic jerk (termed C reflex in the original paper). The latency recorded from the thenar muscles from stimulating the median nerve at the wrist was some 51 ms. When the response was recorded in plantar muscles after stimulating the posterior tibial nerve at the ankle, the latency was some 103 ms. The stimulus at either site evoked abnormal large somatosensory cortical potentials recorded from the opposite hemisphere. The authors concluded that this reflexly evoked myoclonic jerk occurred after an interval sufficient to allow passage of afferent impulses to the cerebral cortex and efferent impulses from cortex down the corticospinal tract to motoneurones. In a subsequent article, Sutton (1975) analysed the critical stimulus required to elicit reflex myoclonus in this patient. Touch or pressure, in the absence of movement, was an adequate stimulus, whereas muscle stretch by itself was not, in keeping with the observation that reflex myoclonus could be induced as easily by stimulation of the digital nerves as by stimulation of the median nerve at the wrist.

The case described by Carmichael and Dawson is
similar to that described by Sutton and Mayer in
that both showed reflex myoclonus in response to a
tendon tap or to mixed nerve stimulation, and both
had giant cortical potentials evoked by the latter
stimulus. The two cases differ in that the critical
stimulus in the first case was stretch of muscle
while in the latter it was touch or pressure.

In the present study, we have investigated a patient
with post-anoxic reflex myoclonus in whom the
critical stimulus would appear to be stretch or
movement, rather than touch or pressure, and in
whom cortical evoked potentials were not abnormally
large.

Case history

A 55 year old man, resident in Bermuda for 20 years,
but originating from Great Britain, had been
asthmatic for 15 years. He had sustained a period of
respiratory and cardiac arrest during an attack of
asthma six months previously. After this he was
unconscious for two weeks before his level of
consciousness gradually improved. However, as he
improved, it was noticed that all four limbs were
rigid and akinetic, and that there were continuous
involuntary movements affecting all four limbs, head,
and trunk. His involuntary movements had persisted
despite treatment with phenobarbitone and diphenyl-
hydantoin and during the six months between the
initial period of anoxia and investigation he had had
two generalised convulsions.

On examination he was alert but disoriented in
time and space. Spontaneous speech was simple in
form and he showed a moderate mixed dysphasia
with perseveration, as well as a marked slurring
dysarthria. He was markedly bradykinetic and
muscle tone was increased in all limbs, having the
characteristics both of extrapyramidal rigidity and
spasticity. His facial expression was rather fixed, and
vertical gaze restricted. His overall posture was one
of flexion, particularly of the trunk, neck, elbows,
and hips, and there were marked bilateral grasp and
rooting reflexes. His tendon jerks were brisk through-
out and plantar responses extensor. Myoclonic jerks
occurred spontaneously and in response to stimuli,
and will be described in detail below.

Investigation revealed a normal haemoglobin and
white blood cell count, with an ESR of 42 mm/h.
Serum urea and electrolytes, calcium, inorganic
phosphate, and liver function were all normal. Skull
and chest radiographs and ECG were within normal
limits. CSF obtained by routine lumbar puncture
showed no cells and a normal protein content. CSF
5-hydroxyindolacetic acid was 23.7 ng/ml (control:
32.6 ± 9.6 ng/ml (mean ± SD)) and CSF homo-

vanillic acid was 26.7 ng/ml (control: 52.3 ± 30.4

ng/ml).

PROCEDURE

The electromyogram and electroencephalogram
were recorded from various sites using surface
electrodes. Signals were preamplified by Devices 3160
amplifiers and fed into a PDP 12 computer. Sponta-
naneous myoclonic jerks were recorded using the
programme PASTIME triggering the computer
from the EMG of one of the muscles in order to
observe events both before and after the trigger.
Elicited jerks were recorded using the programme AV
triggering the computer at the time of delivery of the
stimulus. Data were collected in single trials or in
averages of up to 128 trials.

Averaged data gave inconsistent results on
occasions; this turned out to be due to jitter in the
timing of EMG bursts in different muscles with
respect to each other, and to jitter of the bursts in
different muscles with respect to the stimulus. Thus
most of the results presented were taken from series
of single trials.

The differences in the time of onset of the EMG
burst in one muscle with respect to another muscle
(or the stimulus) was taken as the average difference
in those trials where both muscles were active and the
time of onset could be determined clearly. The
standard deviation of the individual differences was
taken as a measure of the jitter.

Results

NERVE CONDUCTION STUDIES

In order to derive motor and sensory conduction
times, peripheral nerve conduction velocities, mono-
synaptic tendon jerks responses, distal latencies,
and the latencies of F waves and H reflexes were
obtained in the standard manner.

The minimum time needed for a neural signal to
travel down a limb from one muscle to the next was
calculated from motor nerve stimulation (Table 1).
That these times were appropriate in certain physio-
logical circumstances was confirmed using the
monosynaptic tendon jerk data. For example, the
tendon jerk latency in biceps to a tap on the biceps
tendon was 15 ms and in finger flexors to a finger tap
was 21 ms. The difference, 6 ms, represented the
aferent and efferent time from finger flexors to
biceps. Assuming that motor and afferent nerve
conduction velocities were approximately the same,
3 ms was the one-way time from finger flexors to
biceps, a figure confirming the result obtained with
direct motor nerve stimulation. A similar calculation
using the latencies of the quadriceps jerk (25 ms)
and soleus jerk (43 ms) gave 9 ms for the time between
upper leg and calf, slightly longer than the fastest motor conduction time of 6 ms.

SPONTANEOUS JERKS

Clinical observations

Myoclonic jerks were present at rest, were intensified by attempted voluntary and passive movement, were diminished during drowsiness or light sleep, and were absent in deep sleep. The jerks occurred irregularly, on average five to 10 times per minute and usually involved the whole body, although occasionally the jerk was limited to a single limb or part of a limb. Muscles of the four limbs, head, neck, trunk, and face were involved in the myoclonus. Depending on the amount of activity in particular muscles, the jerks would vary in clinical form, but flexors tended to be more active than extensors. Commonly there would be nodding of the head, bending of the trunk, flexion of the arms, shrugging of the shoulders, and flexion withdrawal of the legs.

EMG studies

Electrophysiological surveys were made of the cranial nerve musculature and the four limbs. Large, relatively synchronous bursts of activity in the muscles corresponded to the clinical jerks (Fig. 1). In most spontaneous jerks all muscles surveyed participated regardless of the clinical pattern.

<table>
<thead>
<tr>
<th>Biceps to finger flexors</th>
<th>Motor nerve stimulation</th>
<th>Spontaneous</th>
<th>Toe taps</th>
<th>Finger taps</th>
<th>Median nerve stimulation</th>
</tr>
</thead>
<tbody>
<tr>
<td>6</td>
<td>10 ± 3 (12)</td>
<td>10 ± 6 (16)</td>
<td>13 ± 4 (8)</td>
<td>9 ± 3 (13)</td>
<td></td>
</tr>
<tr>
<td>Biceps femoris to soleus</td>
<td>11</td>
<td>14 ± 3 (14)</td>
<td>7 ± 8 (14)</td>
<td>10 ± 6 (7)</td>
<td></td>
</tr>
<tr>
<td>‘Biceps to biceps femoris’</td>
<td>3</td>
<td>4 ± 4 (12)*</td>
<td>5 ± 6 (14)</td>
<td>0 ± 6 (12)</td>
<td></td>
</tr>
</tbody>
</table>

*Mean latency (± 1 SD) is given in ms and is based on a number of observations (in parentheses) of single records in which both muscles were active, except in the case of finger taps where only averaged records were available.

The mean difference between a pair of muscles shown in Table 1 may differ from that shown in Table 3, in which latencies from stimuli were measured in all records in which the particular muscle was active, irrespective of whether the other muscle fired. These minor differences reflect the jitter of latency for a given muscle with respect to the stimulus and to other muscles.

Fig. 1 Electromyographic recording from right limbs of two spontaneous myoclonic jerks. Muscles recorded from are biceps, finger flexors (F. flex.), biceps femoris (Bic. fem.), quadriceps (Quad.), tibialis anterior (Tib. ant.), and soleus. In part A, the jerk in the arm precedes that in the leg, while, in part B, the earliest activity in leg precedes that in the arm.
Usually both components of an agonist–antagonist pair were active, but, corresponding to the clinical impression of flexor predominance, the amplitude of the flexor component was usually greater than the amplitude of the extensor and at times the extensor component was apparently absent. The EMG of a single jerk was simple in form, having only a few phases and lasting 10–30 ms. The time of onset of activity in one muscle with respect to another of a flexor–extensor pair varied by about 3 ms. The latency from proximal to distal muscles in one limb in individual records varied by 4–5 ms. Despite this jitter, the average time of one muscle’s onset with respect to another was approximately proportional to the distance of the muscles from the neuraxis. By using the average difference in time of the onset of the EMG in different muscles during the spontaneous jerks, it was possible to calculate ‘motor conduction times’ from one muscle to the next; these times were somewhat slower than the maximal motor conduction times obtained by direct nerve stimulation (Table 1).

The same muscles in different limbs had a 6–8 ms jitter with respect to each other; this applied to the two arms, the two legs, or one arm and the ipsilateral leg. On average, one side of the body did not become active earlier than the other. The arms tended to precede the legs, but only by 5–6 ms (biceps with respect to biceps femoris), and in some jerks the legs preceded the arms. The cranial nerve musculature, at least that supplied by the lower cranial nerves, tended to precede the limbs (Fig. 2 and Table 2). The cranial nerve nuclei seemed to be activated in ascending order.

**EEG studies**

The EEG showed predominant small amplitude intermediate slow and fast activity over both hemispheres without any definite alpha rhythm. There were very frequent spikes, often doublets or triplets, followed by slow waves. The spikes were usually triphasic, with successive positive, negative and positive waves (Figs. 2–5); they were generalised in distribution with slightly higher amplitude at the vertex (Fig. 3). The spikes were usually, but not always, associated with the myoclonic jerks; there was a marked variation in the timing of the spike (measured at the beginning of the initial positive component) and the onset of the EMG activity. Activity of the lower cranial nerve musculature (sternocleidomastoid and trapezius), but not of the upper cranial nerve musculature (orbicularis oris and masseter), usually preceded the cortical spike (Table 2). Even the activity of the leg muscles preceded the cortical spikes at times. The failure of the EEG spikes to be tightly time-locked to the EMG discharges was illustrated by the effect of averaging the EEG by triggering the averaging sweep from EMG bursts in an active muscle (Fig. 4). No EEG activity averaged with respect to the myoclonic muscle jerks.

Cortical evoked responses in this patient after median nerve stimulation at the wrist were not abnormally large. Only contralateral potentials could be identified; the initial positive phase was 4 μV in amplitude after left median nerve stimulation and

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**Fig. 2** Electrical record of spontaneous myoclonic jerk including activity in cranial nerve muscles. The electroencephalographic recording (EEG) is from a point 1 cm to the left and 2 cm behind the vertex referred to a mid-frontal electrode (a positive deflection is downward). Other records are from the right masseter, left orbicularis oris (Orb. oris), left sternocleidomastoid (SCM), right trapezius, and right biceps. Note the activation of the cranial nerves up the brain stem, and the onset of activity in trapezius before that in the EEG. The upper voltage calibration refers to the EEG record and the lower voltage calibration to all of the EMG records.

**Table 2** Spontaneous jerks in cranial nerve musculature

<table>
<thead>
<tr>
<th>Muscle</th>
<th>Activity (ms ± 1 SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sternocleidomastoid</td>
<td>3 ± 3 (26)</td>
</tr>
<tr>
<td>Orbicularis oris</td>
<td>10 ± 6 (24)</td>
</tr>
<tr>
<td>Masseter</td>
<td>19 ± 4 (16)</td>
</tr>
<tr>
<td>EEG spike (initial positive)</td>
<td>4 ± 5 (25)</td>
</tr>
<tr>
<td>Biceps</td>
<td>11 ± 4 (24)</td>
</tr>
</tbody>
</table>

The onset of activity in a given muscle or the time of onset of the initial positive wave in the EEG, was measured from the time of onset of activity in trapezius. Mean latency in ms (± 1 SD) is shown for a number of observations (in parentheses).
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In each part of the diagram the EEG is recorded from the vertex referred to the left ear (a positive deflection is downward), and the EMG is from the left sternocleidomastoid muscle. Part A shows the events in a single spontaneous jerk. Part B is the average of 84 spontaneous jerks with the averaging being triggered on the occurrence of the EMG event. Since the EEG record in part B shows essentially no activity, the timing of the EEG spike seen in part A must vary considerably with respect to the EMG jerk.

Fig. 3 Electroencephalographic correlate of a spontaneous myoclonic jerk. The electroencephalogram is sampled from five widely separated sites in the left hemisphere as indicated in the diagram and referred to a mid-frontal electrode (R) (a positive deflection is downward). Activity in the right biceps is taken as a marker of the myoclonic jerk. It is apparent that the electrocortical spike activity is widespread. The upper voltage calibration marker refers to all of the EEG records and the lower voltage calibration marker refers to the EMG record.

5 μV after right median nerve stimulation. The latency to onset of these potentials was 29 ms and 22 ms respectively. The flash evoked visual response and the auditory evoked potential were similarly not exceptional in size or latency.

ELICITED JERKS

Clinical observations

Myoclonic jerks could be precipitated by a variety of sensory stimuli, including startle, an abrupt noise, or a tendon tap. In the latter case, light taps might evoke jerks confined to the stimulated limb, but usually involved the whole body. With regard to the sensitivity to a tendon tap, the arms were much more sensitive than the legs, and the hands and feet were much more sensitive than proximal structures. The most reactive site was the fingers. A light tap delivered to the tips of the fingers or thumb to elicit a finger jerk would regularly cause a myoclonic jerk. Taps to the tendons of biceps or triceps rarely caused jerks. Taps to the pads of the toes often caused jerks, but taps to the Achilles tendon and patellar tendon were ineffective. Careful analysis of the adequate stimulus required to elicit a myoclonic jerk in response to stimulation of the fingers or thumb revealed that pinprick, light touch, or deep touch to the tips of the fingers were ineffective. Even a brisk tap with a tendon hammer on the pads of the fingers did not necessarily cause jerks when they were firmly supported on a flat surface to prevent movement at joints. However, a tap to the...
finger tips such as to elicit a tendon jerk regularly caused a myoclonic response.

**Muscle stretch**

A tendon hammer, incorporating a triggering device, was used in the usual clinical manner to stretch the finger flexors, biceps, triceps, and toe (and foot) plantar flexors. The resulting EMG responses correlated with the visible myoclonic jerks evoked and were similar in appearance to those seen with the spontaneous jerks.

Single responses to taps to the toes were studied extensively (Fig. 5). The EMG activity elicited with this stimulus was most prominent in the ipsilateral lower leg, but was usually seen in the upper leg as well. A larger amplitude of EMG activity in flexor muscles (similar to spontaneous jerks) correlated with the clinical appearance of flexion at hip and knee and dorsiflexion at the ankle. Activity was frequently seen in the opposite leg and sometimes in the ipsilateral arm, but this latter response was so erratic that it was uncertain whether it was part of the myoclonic jerk being studied. The latency of response from the stimulus to the individual muscles was markedly variable with a jitter of 9 to 13 ms (Table 3). The jitter of a muscle to its antagonist was about 3 ms, the jitter to another muscle in the same limb was about 5 ms, and the jitter to the same muscle in the opposite limb about 9 ms. All these values were similar to those obtained with the spontaneous jerks. Within the same limb the average time of onset of the EMG varied with the distance from the neuraxis; ‘conduction times’ from one muscle to another could be derived and were generally similar to those found for spontaneous jerks (Table 1). Only averaged data were available for the myoclonic jerks produced by taps to the fingers (Fig. 6a). Responses were seen only in the biceps and finger flexors of the limb being stimulated. Derived conduction time between these two muscles was similar to that obtained by other methods (Table 1) and latencies of response were similar to those obtained by mixed nerve stimulation (see below and Table 3).

**Mixed nerve stimulation**

Direct electrical stimulation of the median nerve at the wrist or elbow was a very effective stimulus for myoclonic jerks, whereas stimulation of the posterior tibial nerve at the ankle gave inconsistent responses. As with spontaneous jerks and those produced by muscle stretch, the electromyographic correlate of the myoclonus was a large, relatively synchronous burst of activity in the muscles. The results of stimulation of the median nerve at the wrist were studied in detail (Fig. 7). In the figure, there are three responses in abductor pollicis brevis. The first is the direct muscle response (M response) to the nerve stimulation, which has overloaded the amplifier at the gain employed. This is followed by the F wave. Last, corresponding in time with activity in the other muscles, is the myoclonic burst of activity. The myoclonus was widespread with a regular response in all the muscles of the same arm and usually in the ipsilateral leg. The latency of response (Table 3) in the various muscles of the arm showed a variation of 4–8 ms and in the leg of 11–15 ms. The jitter of various muscle pairs with respect to each other and the ‘motor conduction times’ found by considering onset of activity for muscles within the same limb (Table 1) were generally similar to those calculated from records of spontaneous jerks or those elicited by muscle stretch. The arms (biceps) preceded the legs (biceps femoris) by $8 \pm 13$ ms. For median nerve stimulation at the
wrist the motor threshold for EMG response in abductor pollicis brevis was about 80–90 V with a 100 µs pulse. Myoclonic activity was obtained with voltages of 75% of threshold (Fig. 8a).

Conduction velocity in the peripheral part of the afferent path responsible for the myoclonus can be deduced by comparing the latency of the EMG activity after a distal and a proximal stimulus. Using averages of 32 single trials, the latency of the myoclonic activity in biceps was compared after median nerve stimulation at the wrist and elbow. The average latency to the onset of biceps activity was 33 ms in 25 averages of 32 trials after median nerve stimulation at the wrist and was 29.5 ms in five averages of 32 trials after stimulation at the elbow. The distance between the sites of stimulation was 28 cm; thus the afferent conduction velocity was about 80 m/s.

Sensory nerve stimulation
Using ring electrodes the sensory nerves to the fingers were stimulated. The thumb, index, and middle fingers were stimulated separately with 100 µs pulses of up to 90 V, parameters which were effective when stimulating the median nerve at the wrist (Fig. 8B). (The patient's mental state made it impossible to determine a sensory threshold.) With this form of stimulation myoclonic jerks were not seen clinically or in EMG records of single trials or averages of 32.

DRUG EFFECTS
Clonazepam
An intravenous bolus of 1 mg clonazepam was given. Within one minute the spontaneous myoclonus had stopped. Myoclonic jerks could no longer be averaged by tendon taps to the fingers or median nerve stimulation at the wrist (100 µs, 90 V), conditions which were previously effective (Fig. 6b and 9b).

5-Hydroxytryptophan (5-HTP)
The patient was given an intravenous infusion of 150 mg 5-HTP in 500 ml of normal saline over two hours. At the end of the infusion the myoclonus had disappeared and was still absent five hours later at the time of physiological testing. The patient was nauseated, depressed, and rather drowsy, but was easily

Table 3  Latencies for elicited jerks

<table>
<thead>
<tr>
<th></th>
<th>Toe taps</th>
<th>Finger taps</th>
<th>Median nerve shocks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Biceps</td>
<td>35</td>
<td>39 ± 4 (12)</td>
<td></td>
</tr>
<tr>
<td>Triceps</td>
<td>No response</td>
<td>43 ± 8 (16)</td>
<td></td>
</tr>
<tr>
<td>Finger flexors</td>
<td>37</td>
<td>39 ± 5 (19)</td>
<td></td>
</tr>
<tr>
<td>Finger extensors</td>
<td>No response</td>
<td>43 ± 5 (19)</td>
<td></td>
</tr>
<tr>
<td>Abductor pollicis brevis</td>
<td>No response</td>
<td>49 ± 4 (13)</td>
<td></td>
</tr>
<tr>
<td>Biceps femoris</td>
<td>74 ± 9 (16)</td>
<td>44 ± 13 (12)</td>
<td></td>
</tr>
<tr>
<td>Quadriceps</td>
<td>75 ± 9 (15)</td>
<td>Not studied</td>
<td></td>
</tr>
<tr>
<td>Tibialis anterior</td>
<td>87 ± 12 (17)</td>
<td>58 ± 15 (8)</td>
<td></td>
</tr>
<tr>
<td>Soleus</td>
<td>89 ± 13 (17)</td>
<td>Not studied</td>
<td></td>
</tr>
<tr>
<td>Flexor hallucis brevis</td>
<td>96 ± 11 (14)</td>
<td>70 ± 11 (11)</td>
<td></td>
</tr>
</tbody>
</table>

Mean latency (± 1 SD) is given in ms and is based on a number of observations (in parentheses), except in the case of finger taps where only averaged records were available. Latencies were measured in ms from the stimulus trigger produced by the tap or shock to the initial EMG deflection in the response evoked in the muscles stated.

Fig. 6 Myoclonic responses to finger tap (sudden extension of the fingers). The records, which are averages of 64 trials, are taken from biceps (BIC), finger flexors (FF), and abductor pollicis brevis (APB). Part A is the control, part B is after the administration of clonazepam, and part C is after the administration of 5-HTP (see text for details). Because APB showed no averaged myoclonic activity even on the control run, it is not illustrated in parts B and C. Note in the control record the initial burst of activity at tendon jerk latency in finger flexors, which is followed by the myoclonic burst. After clonazepam, the tendon jerk is unaffected, but the myoclonic burst is absent or reduced. After 5-HTP, the tendon jerk is enhanced, but the myoclonic burst is reduced.
aroused. Clinically the tendon jerks were enhanced. Averaged responses were obtained to tendon taps of the fingers (Fig. 6c) and to stimulation of the median nerve at the wrist (Fig. 9c) and were compared with the responses the previous day (with similar EMG electrode placement) before the drug was given. After the administration of 5-HTP myoclonic response to finger taps was abolished in finger flexor muscles, but interpretation of the EMG response from biceps was difficult because of a large monosynaptic potential not previously present. The myoclonic response to median nerve stimulation in abductor pollicis brevis was abolished and the amplitudes of responses in finger flexors and biceps reduced by approximately 60%.

**Discussion**

In this paper we have assumed that the spontaneous myoclonic jerks are equivalent to the elicited jerks and that their site of origin in the central nervous system is the same. The same muscles are involved in the same temporal sequence in both and the electromyographic activity in individual muscles and the electroencephalographic discharges are similar. It is reasonable to suppose that the spontaneous jerks are elicited by stimuli which have not been identified, but which are always present, although we cannot exclude the possibility of spontaneous discharges.

The jerks have been studied in averages or in series of single sweeps. As has been pointed out, averaged data can be confusing, for a component can become smaller because of an actual decline in size, less frequent appearance, increased jitter, or any combination. It was jitter of the muscle bursts that first drew our attention to the difficulty in interpreting averaged data. There was jitter of any particular muscle with respect to an eliciting stimulus, and there

**Fig. 7** Electromyographic recording of a myoclonic jerk in response to right median nerve stimulation at the wrist. Muscles recorded from, all on the right side, are biceps, triceps, finger flexors (F. flex.), finger extensors (F. ext.), abductor pollicis brevis (APB), biceps femoris (Bic. fem.), tibialis anterior (Tib. ant.), and flexor hallucis brevis (FHB). In APB, three responses can be seen: the direct M responses (which overload the amplifiers), an F wave, and then the myoclonic jerk.

**Fig. 8** A comparison of the response to mixed digital nerve stimulation. Electromyographic records, which are averages of 128 trials, are taken from biceps, finger flexors (F. flex.) and abductor pollicis brevis (APB). In part A, the median nerve was stimulated at the wrist with 67 V for 100 μs, which was below threshold for a direct motor response in APB. In part B the middle finger was stimulated with 90 V for 100 μs utilising ring electrodes.
was jitter of one muscle with respect to another. The further distal a muscle was from another, the greater was the jitter.

The phenomenon probably reflects the nature of the supraspinal organisation of the myoclonus—for example, variable spread in a complex polysynaptic network—since the signal sent to the spinal cord is powerful, producing absolutely stereotyped and almost synchronous firing of each muscle.

The times required for the neural signals to traverse peripheral motor nerves from point to point have already been noted (Table 1). The data from the myoclonic jerks gave values similar but slightly slower than those obtained with direct electrical stimulation. This and other data can also be used to make some deductions for approximate conduction times in central pathways mediating the myoclonus (Table 4).

The time from biceps to the cervical spinal cord can be estimated from the time for the monosynaptic tendon jerk (15 ms). Half of this time, about 8 ms, was needed for the signal to go one way. The same time can be deduced and confirmed from other observations including the finger flexor monosynaptic stretch and the F wave in abductor pollicis brevis, knowing the motor conduction delay from biceps to those muscles.

In similar fashion, the time from biceps femoris to the lumbar spinal cord can be estimated from the latency of the quadriceps tendon jerk, assuming quadriceps and biceps femoris are about equally distant from the cord (as seems to be the case; see, for example, Table 3). The tendon jerk latency was 25 ms, which implies that the time from biceps femoris to the cord was about 13 ms. Similar values can be calculated from the timing of the H reflex or the ankle jerk.

At least for the purpose of computing latencies, we can define a site in the central nervous system, M,

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**Table 4  Derived 'central conduction times'**

<table>
<thead>
<tr>
<th>Site</th>
<th>Time (ms)</th>
<th>Data used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cervical spinal cord to biceps</td>
<td>8</td>
<td>Biceps tendon jerk, finger flexor jerk, F wave</td>
</tr>
<tr>
<td>Lumbar spinal cord to biceps femoris</td>
<td>13</td>
<td>Quadriceps tendon jerk, ankle jerk, H reflex</td>
</tr>
<tr>
<td>Cervical spinal cord to M and back</td>
<td>18</td>
<td>Latencies of myoclonic jerks in arm muscles after median nerve stimulation and finger taps</td>
</tr>
<tr>
<td>Cervical spinal cord to lumbar spinal cord</td>
<td>2</td>
<td>Latencies of myoclonic jerks in leg muscles with respect to arm muscles in spontaneous and median nerve elicited jerks</td>
</tr>
<tr>
<td>Lumbar spinal cord to cervical spinal cord</td>
<td>14</td>
<td>Comparison of latencies of myoclonic jerks in leg muscles after median nerve stimulation and toe taps</td>
</tr>
<tr>
<td>11th nucleus to 7th nucleus</td>
<td>10</td>
<td>Latency in spontaneous jerks</td>
</tr>
<tr>
<td>7th nucleus to 5th nucleus</td>
<td>9</td>
<td>Latency in spontaneous jerks</td>
</tr>
</tbody>
</table>

M refers to the hypothetical source of the myoclonus in the central nervous system.
where the myoclonus is 'generated'. As we know the
time required for neural signals to travel back and
forth from the spinal cord, the time from the cord to
M can be determined. The peripheral part of the
efferent limb of the myoclonus consists of rapidly
conducting fibres, just slightly slower than the most
rapid motor fibres (Table 1). The peripheral part of
the afferent limb also consists of rapidly conducting
fibres as deduced from a comparison of median nerve
stimulation at the wrist and elbow (see p. 259). The
latency of the evoked myoclonic jerk in biceps after
median nerve stimulation at the wrist was 39 ms.
Basing our calculations on motor conduction vel-
cocities, it would have taken about 6 ms for the afferent
volley to reach the level of biceps, and, as already
indicated, it would have taken 15 ms for the signal to
traverse the segment biceps to cervical spinal cord and
back. This leaves 18 ms for the loop from the spinal
cord to M and back. A similar value can be calculated
from the latency of the myoclonic jerk in biceps
evoked by finger taps (35 ms). Subtracting 3 ms for the
time taken for the afferent volley from the finger
taps to reach the level of biceps, and 15 ms for the
segment biceps to spinal cord and back, leaves 17 ms
for the spinal cord to M loop. Another method for
calculating the time around this loop comes from a
direct comparison of the latencies in abductor
pollicis brevis of the F wave (32 ms) and the myo-
clonic activity after median nerve stimulation at the
wrist (49 ms). Since the F wave latency reflects the
time to the spinal cord and back, the excess latency of
the myoclonic activity (17 ms) is the central loop time.

The time required for the efferent signal to traverse
the spinal cord can be deduced from the difference in
time of a myoclonic jerk appearing in the legs com-
pared with one in the arms. In spontaneous jerks,
biceps preceded biceps femoris by 5 ms and in jerks
evoked by median nerve stimulation biceps preceded
biceps femoris by 8 ms (average of these two values is
about 7 ms). Basing calculations on tendon jerk
latencies (and other measures, cf. Table 4), it was 5 ms
longer from the lumbar spinal cord to biceps femoris
than from cervical spinal cord to biceps (13 ms–8 ms).
So it must take about 2 ms (7 ms–5 ms) for the myo-
clonus signal to go from the cervical cord to the lum-
bar cord. Thus, the efferent conduction velocity for
the myoclonic signal in the spinal cord is rapid, sug-
gest that the signal travels in a strongly influential,
rapidly conducting oligosynaptic pathway.

The time that the afferent signal takes to traverse
the spinal cord can be deduced from the difference in
timing between jerks in the legs produced by stimula-
tion of the arms and that produced by stimulation of
the legs. The biceps femoris latency was 74 ms after
toe taps and 44 ms after median nerve stimulation at
the wrist (Table 3), a 30 ms difference. Some of this
time is accounted for by a longer afferent path in the
periphery. From values previously calculated, the
time for median nerve at the wrist to cervical spinal
cord was 14 ms and the time from toe flexors to lum-
bar cord was 30 ms, a difference of 16 ms. This leaves
14 ms as the afferent time from lumbar to cervical
cord. This suggests that the afferent path in the spinal
cord is much slower than the efferent path. If we
assume that the long toe flexors and foot plantar
flexors are also stimulated by toe taps, as they
probably are, then the peripheral time in the leg is
reduced and the apparent afferent conduction time
further prolonged. This apparently long afferent
signal time could be explained, at least in part, if
muscle stretch takes longer to produce myoclonus
than does mixed nerve stimulation; but this does not
seem to be the case, as the latency of arm muscle
response after finger taps was similar to the latency
after median nerve stimulation (Table 3). Another
explanation would be that it takes longer for afferent
signals to be processed in M if they are from the legs
than if they are from the arms.

Considering the cranial nerve musculature, the 11th
nerve nucleus (trapezius and sternocleidomastoid) was
activated first, followed by the 7th nerve nucleus
(orbicularis oris) 10 ms later, and the 5th nerve nu-
cleus (masseter) 19 ms later. Thus the signal producing
the myoclonus seemed to travel (rather slowly) up
the brain stem.

The appearance of the EMG burst in trapezius and
sternocleidomastoid was the earliest electrophysio-
logical event detectable in a myoclonic jerk. Taking
half of the 8 ms required for the monosynaptic stretch
reflex in sternocleidomastoid gives 4 ms for the time
from the 11th cranial nerve nucleus to its muscles. Re-
calling that biceps was 8 ms removed from the cervical
cord (Table 4) and that the 11th nerve musculature
preceded biceps by 8 ms (Table 2), we see that the 11th
nerve nucleus was activated 4 ms before the cervical
spinal cord. In regard to cortical spikes, as the EMG
burst in sternocleidomastoid preceded these by 1 ms,
the 11th nerve nucleus was activated 5 ms before the
generation of the spike. Thus, activation of the 11th
erve nucleus was the earliest phenomenon on the
output side of the myoclonic jerk that we have
identified. The myoclonus generating centre, M, must
be closer to the 11th nerve nucleus than to any other
CNS structure examined.

The myoclonic jerks elicited in this case by mixed
nerve electrical stimulation or tendon taps occurred
after a delay too long for spinal cord mediated mono-
synaptic and other 'short latency' reflexes, yet too
short for voluntary responses. They correspond to
so-called 'long latency' reflexes. Some long latency
reflexes can be mediated entirely in the spinal cord,
while others, called long loop reflexes, require the
participation of supraspinal structures. Several different long latency reflexes have been discovered in animals and man, and derangement of one (or more) of these may be responsible for the myoclonus.

The association of a type of myoclonus with such a reflex system would provide a rational system for classification of some of the different types of myoclonus.

The characteristics of the present case that seem important are:

1. The myoclonic jerk had a simple form and when it occurred it tended to involve all of the muscles in the body, including proximal muscles.

2. The EEG showed generalised spikes that were associated, but not time locked, to the jerk.

3. Afferent cord conduction time was long, while efferent cord conduction time was short.

4. Activation of cranial nerves was upward and the earliest event in the jerk could be localised in the medulla.

5. The critical stimulation for producing the myoclonus was muscle stretch.

6. The myoclonus was responsive to 5-HTP and clonazepam.

The first four characteristics point to a type of myoclonus mediated in the lower brain stem and probably in the reticular formation. Halliday (1975) has recently summarised the literature suggesting that the reticular formation may play an important role in myoclonus. The spinobulbospinal reflex, best studied in the cat, has tentatively been identified in man and is the only known long latency reflex mediated in the reticular formation (Shimamura et al., 1964; Meier-Ewert et al., 1972). The myoclonus in our patient, however, cannot be identified with that spinobulbospinal reflex, as the latter is characterised by fast afferent cord conduction and slow efferent cord conduction velocity. Rapid oligosynaptic pathways are known to originate in reticular formation and these must be involved in the kind of myoclonus under discussion here.

Other patients, with similar characteristics, have now been identified and we refer to this group as the 'reticular reflex myoclonus' type. They are contrasted with another group of patients who share the hypersynchronous EMG appearance of the myoclonus. In these patients the jerks are restricted to the area of stimulus, they have large sensory evoked potentials and the EEG correlate of the myoclonic jerk is a time-locked fragment of the sensory evoked potential. We refer to this group as the 'cortical loop reflex' type, as this seems to be the underlying mechanism. Details of the other patients will be reported elsewhere.

The fifth point in the above list characterising the present patient concerns the critical stimulus for the reflex myoclonus. As in the patient of Dawson (1947), stretch was the critical stimulus and simple touch was ineffective. On the other hand, for the patient of Sutton and Mayer (1974), touch was the critical stimulus and pure stretch was ineffective. We must conclude that both phenomena are possible, but then ask how this feature helps in the separation of different types of myoclonus. Theoretically, identification of the critical stimulus should help in identifying the reflex which has become deranged in producing the myoclonus. Practically, this does not seem useful yet, partly because the critical stimuli for the known long latency reflexes have not been well established. Additionally, even though both the present patient and Dawson's patient require stretch to produce the myoclonus, the present patient falls into the category of reticular reflex myoclonus while Dawson's patient would most probably be in the category of cortical loop reflex myoclonus. The patient of Sutton and Mayer would also most probably be in the cortical loop reflex category, yet the critical stimulus differs from that of Dawson's patient. These facts, together with the analysis of our other patients, suggest that critical stimulus is not a feature that separates the two identified categories. It may be, however, necessary eventually to subdivide these categories.

The excellent response of this patient's myoclonus to 5-HTP and clonazepam is shown by other patients with the physiological characteristics of the reticular reflex type of myoclonus (Chadwick et al., 1976). In such patients there is biochemical evidence that the myoclonus is related to a relative deficiency of brain serotonin (5-HT) and that the therapeutic effects of 5-HTP and clonazepam may be mediated by their action on 5-HT metabolism. In man the highest concentrations of 5-HT are found in the midbrain and medulla (Gottfries et al., 1974). Thus, the physiological and biochemical data are complementary in implicating brain-stem structures in the myoclonus of these patients. It is likely that the unknown reflex mediating the myoclonus is usually inhibited by the action of 5-HT systems of neurones.

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