Neurophysiological mechanisms in abnormal reflex activities in cerebral palsy and spinal spasticity

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Summary

Dorsal root stimulation, carried out during operation in 80 spastic cerebral palsy patients and in one spastic quadriplegic patient, allowed a study of abnormal monosynaptic and polysynaptic reflexes. Frequency-related depression of the monosynaptic reflex was not present; increased activity through non-suppressed polysynaptic pathways was shown.

The exact pathophysiological mechanisms underlying spasticity which determine its clinical features have not been fully elucidated. Even a concise and universal definition of what entities the term spasticity includes is not presently available.

Much of the investigative efforts have been focused on the gamma loop and muscle spindles, on the assumption that the main pathophysiology was related to gamma hyperactivity. In recent years, however, evidence has accumulated that simple gamma hyperactivity cannot account for all the features of spasticity and even the classical view of the gamma loop organisation has undergone modifications.

New evidence has shown that other mechanisms, such as lack of presynaptic inhibition and other abnormalities in interneuronal activity, may be involved in spasticity. These deductions resulted from the use of the tonic vibration reflex (TVR) and studies of the ratios of H reflex (vibration) to H reflex (control) and of Achilles tendon reflex to M response. Preliminary results of dorsal root stimulations (DRS) in cerebral palsy patients and its usefulness in the surgical management of spasticity have been reported previously by one of the authors. This paper deals with a detailed neurophysiological analysis of the abnormal reflexes in patients with spinal cerebral palsy and adds the findings in a neurophysiological study of a case of spinal spasticity.

Materials and methods

A Cerebral spasticity

Eighty subjects with cerebral palsy, ages between 5 and 25, underwent functional posterior rhizotomy for the relief of spasticity. Dorsal root stimulation (DRS) was used to show which roots or rootlets were involved in abnormally excitable circuits; we will refer to these as pathological roots or rootlets. All of the subjects presented with spasticity, that is, increased stretch reflexes, hyperactive tendon reflexes and abnormal postural patterns. In a few of them, dystonic or athetoid attitudes or both were associated with spasticity.

With the patient in the prone position, a D12-L1 laminectomy exposed the lumbar roots; the roots and rootlets were electrically stimulated in sequence. An average of about 105 rootlets were stimulated in each patient using bipolar hook electrodes which were separated by 1 cm. Care in handling the roots was paramount, as even a minimal stretch or compression could modify their electrical excitability. The stimulation and recording equipment used was a Modular Electrophysiological System: Display and Recording Unit DFO6 MKII-N6, AVM6 Signal Averager and Biological Amplifier AA6 MK II-PA 467/15 (Medelec Limited, Woking, Surrey, England). Pulse duration was 0.5 msec. Ketamine hydrochloride was used for the anaesthesia. Reflex muscle responses were recorded using surface or needle electrodes. The EMG response following DRS is representative of synchronous multimotoneuronal activity.

Stimulation sequence for each root or rootlet was as follows. At 1 Hz, the pulse amplitude was adjusted...
to obtain a reflex muscle contraction just above threshold. The muscle groups involved were then identified. With this pulse amplitude adjusted, the stimulation frequency was set at 50 Hz. The reflex contraction patterns were observed, active muscle groups and patterns of contraction were identified. Occasionally, intermediate frequencies were used and adjustments to the DRS strength were made.

According to the various patterns observed, the criteria as to which roots or rootlets giving pathological responses should be sectioned were established. Thirty-six of the 80 cerebral palsy cases were analysed regarding the number and patterns of abnormal reflexes, and their clinical correlations.

**B SPINAL SPASTICITY**

Seven patients with chronic spinal cord injury underwent Bischof's myelotomy for the relief of intractable spasticity. Essentially, the lower thoracic and lumbar cord was split laterally into anterior and posterior halves, resulting in a disruption of spinal reflexes. Technically, the extent of the myelotomy was determined by stimulating the dorsal roots and observing muscle contractions in the abdomen and lower extremities.

Prior to myelotomy, in one patient records were made from the T11 ventral root while the corresponding dorsal root was stimulated, providing a study of spinal pathophysiology in a very spastic subject. This patient was a 21 year old female who, four years previously, had a C5 fracture followed by immediate complete tetraplegia. The patient presented with severe spastic tetraplegia with marked flexor spasms of the lower limbs associated with hyperactive deep tendon reflexes, bilateral Babinski signs and an uninhibited neurogenic bladder. Touch, vibration and position sense were preserved while temperature and deep pain sensibility were absent below C6 level; pinprick sensation was partially present. The dorsal root was stimulated through silver wires, connected to a Grass stimulator (S8) with isolation unit. Monophasic pulses of 0.2 ms were used in the range of 1-0 to 5-0 V at 1 Hz. Biphasic recordings from the ventral root were amplified through a Tektronic 3A9 differential amplifier and tape recorder for later analysis and photography (figs 6-8). The recording procedure was shortened by the use of tape recording lasting about 30 minutes; anaesthesia was reduced to a lighter level using nitrous oxide and oxygen.

**Results**

In the 36 cerebral palsy cases analysed, 3840 rootlets were stimulated, either singly or in bundles, and determined to be either normal or pathological in their reflex responses (fig 1). Eighty per cent of the rootlets showed pathological responses, except in three cases only 25–30% of the rootlets produced such responses. Rootlets with the pathological responses had a lower threshold of stimulation to produce a detectable EMG response (fig 2). At or below a level of 1-0 V, 60% of the pathological rootlets were excited compared to 34% of the normal rootlets.

**Monosynaptic reflexes**

**A CEREBRAL SPASTICITY**

In the cerebral palsy patients, we studied the reflex muscle response to a just suprathreshold stimulus, applied to a dorsal root or rootlet. As the lowest threshold stimulus fires the largest fibres (Ia afferents), the EMG response results from a synchronous monosynaptic multineuronal discharge. Three
main features of the monosynaptic reflex arc have been observed:

1 Normal reflex activity: with 1 Hz DRS the reflex response is normally repeatable and without loss of amplitude (fig 3A). When the stimulation frequency is increased, the multineuronal reflex response is reduced in amplitude (fig 3A, 10 and 20 Hz). At frequencies between 20 and 50 Hz, there is an early failure to follow and the amplitude of the response is more reduced (fig 3A). At 50 Hz only the first stimulus succeeds in producing a reflex muscle contraction. Fasano et al. recording from the ventral roots showed a similar failure of transmission at 50 Hz, so demonstrating the failure at the spinal level rather than at the myoneural junction or muscle fibre. A similar decrease in amplitude of the monosynaptic response with increasing stimulation frequency and absence of the subsequent responses after the first response was observed by Decandia et al. They stimulated Ia afferents in anaesthetised cats and recorded from single filaments of the ventral roots.

In the patients with cerebral palsy, normal monosynaptic activity was found in approximately 20% of the roots or rootlets stimulated, with considerable variation from case to case.

2 Abnormal reflex activity. a Increased activity of the reflex arc. At 50 Hz, 80% of the stimulated roots or rootlets showed pathological reflex activity following stimulation, about 70% were involved in abnormal monosynaptic reflex activity as shown in figs 3–5. Some 50% of the abnormal reflexes were attributable to abnormal polysynaptic activity (see below). Further analysis of our earlier data confirmed that these segmental monosynaptic pathways transmit without failure and show three basic patterns. The first pattern is the ability of the system to transmit without any sign of failure, as seen in fig 3B. The second pattern shows ability to follow the high frequency (50 Hz) but with reduced amplitude (fig 3C). The third pattern shows a cyclic variability of the amplitude (fig 3D).

A further demonstration of this abnormality is seen when recording from two different muscle groups (fig 4, tibialis anterior A; triceps surae B) following stimulation of one dorsal root at 50 Hz. With a progressive increase of stimulus intensity from zero, the tibialis anterior shows no reflex activity while the triceps surae shows a reflex contraction proportional in amplitude to the increase in stimulus intensity. When the stimulus intensity is threshold and applied at 50 Hz the tibialis muscle, however, responds with a normal pattern in that only the first transmission occurs. But the triceps muscle contracts and abnormally follows the 50 Hz stimulation throughout with less amplitude (fig 4B).

An abnormal and in our experience unique, response was observed following repetitive stimulation at various frequencies (fig 5). Although the trains of stimuli were continuous, the reflex activity occurred only in periodic bursts of 600–800 ms. The intervals between each burst were between 800 and 1000 ms, regardless of the frequency of stimulation (5–50 Hz).

b Decreased activity of the reflex arc. At stimulation frequencies below 5 Hz, Fasano et al. described depression and failure of discharge. In further analysis of this data, circuits displaying such a feature (fig 3E) appeared to occur rarely.

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**Fig 3** Main types of monosynaptic reflex EMG responses evoked in the CP patients following repetitive DRS. A: normal reflex activity. B-D: abnormally increased reflex activity. E: abnormally decreased reflex activity.

**Fig 4** CP patient. Stimulation of a dorsal root at 50 Hz EMG recording on tibialis anterior (A) and triceps surae (B). C: stimuli. Left side of recording: 50 Hz stimulation with progressive increase in strength. Right side of recording: 50 Hz constant stimulation (liminal for monosynaptic reflex).
B SPINAL SPASTICITY

In the 21 year old patient with spastic tetraplegia undergoing myelotomy, the following spinal cord physiology study was undertaken. With both stimulating and recording electrodes approximately 70 mm from the spinal cord on the dorsal and ventral root (fig 8H), single shocks to the dorsal root (DR) evoked a triphasic ventral root (VR) potential at a constant latency of 2-0 ms (fig 6A) giving rise to a conduction velocity of about 70 m/s for the entire reflex.

The earliest component of the VR discharge appears with stimulus intensities of approximately 1-0 V and progressively increases in amplitude with stimulus intensity (fig 6 C-E). This initial synchronous low threshold response is presumably due to activation of the monosynaptic spinal reflex.\(^\text{20-21}\) It is followed by relatively prolonged activity which appears only at higher stimulus intensities (fig 6 B, D, E) at a latency of 4 ms. At the intensities required to evoke this later activity, contraction of the abdominal muscles at the umbilical level followed each stimulus. It is possible, therefore, that this later activity could be due to the resulting spindle afferent activity, since the reflex loop was intact. However, the estimated conduction distance to the site of visible muscle contraction was 40-50 cm; and impulses in Ia fibres, travelling at a rate of 100 m/s, would require 8 to 10 ms to complete the loop. Furthermore, these delayed discharges appeared while the monosynaptic response was depressed (fig 8A). The delayed responses were therefore attributed to polysynaptic reflex activity evoked by dorsal root stimulation.\(^\text{21-22}\)

Two methods were used to assess the excitability of these reflexes. Trains of repetitive stimuli at 5 V were delivered at various frequencies while monitoring the monosynaptic reflex responses (fig 7). At frequencies of 60 Hz the first and fifth stimuli evoked responses of nearly controlled amplitude, but there is a marked depression of reflex excitability during the intervening 67 ms (fig 7B). At higher stimulation frequencies (100-400 Hz), monosynaptic reflex excitability is increased for approximately 10 ms following the initial stimulus (fig 7 C-H);
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Fig 8 Excitability cycle of monosynaptic ventral root reflex. Subject with spinal spasticity. Experimental conditions as shown in H. Graph shows the peak-to-peak amplitude of the monosynaptic reflex plotted as percentage of control amplitude (ordinate), each point being a single determination. Abscissa gives the interval between the conditioning (c in A) and testing (t in A) stimuli, each being 4 V and delivered at 1 Hz from a single dorsal root electrode. A-G are sample records taken at the c-t intervals indicated at the right of each trace of 3 superimposed sweeps.

subsequent stimuli produced transmissions but at a decreased amplitude (fig 7F).

This response pattern was confirmed by a detailed examination of the excitability cycle as shown in fig 8. In this test, pairs of identical DR stimuli were delivered at various intervals at a rate of 1 Hz, each stimulus being of sufficient strength to evoke a polysynaptic DR response. It was observed that the period of increased monosynaptic reflex excitability was 10–12 ms, and it overlapped the onset and duration of the polysynaptic responses. During this period, the amplitude of the second or testing response was increased by 2–3 times the control amplitude (fig 8A, B). This confirmed the presence of a hyperexcitable state in the DR-VR synapse system following a transmission. Of considerable interest was the absence of the polysynaptic response during the facilitated monosynaptic response (fig 8A, B) and its subsequent reappearance from 12 ms onwards. The monosynaptic response failed to appear from approximately 20 ms to 60 ms after the initial transmission, recovering at 90 ms. Throughout this period of depressed monosynaptic activity, the polysynaptic activity was present and appeared normal (fig 8 C-G).

Polysynaptic reflexes

A CEREBRAL SPASTICITY

Polysynaptic activity is minimal or absent when large myelinated afferents are stimulated at a threshold liminal for the monosynaptic reflex in the anaesthetised cat and in man. Following repetitive stimulation, similar minimal polysynaptic activity is seen in the cat and in the EMG recordings of Cook and of Barolat-Romana in normal humans.

In the cerebral palsy series, however, 50% of the dorsal roots or rootlets stimulated repetitively at a threshold for the monosynaptic reflex showed marked polysynaptic activity. Although not evident at 1 Hz stimulation, this polysynaptic activity was progressively built up as the frequency increased.

At 50 Hz stimulation the polysynaptic activity was characterised by: 1 a delayed latency of about 150–200 ms with a progressive increase in amplitude. With cessation of the stimulation, the activity continued for a few hundred ms with a decreasing amplitude (fig 9A); 2 an interferential EMG recording (fig 9A); 3 an irradiation phenomena involving multimuscular groups and multisegmental levels. Such irradiation patterns vary from patient to patient and from root to root; other muscle groups in the ipsilateral or contralateral lower extremities can be involved as well as the upper extremities and the muscles of the neck and trunk.

Two definite patterns of abnormal irradiation have been commonly observed: bilateral hip adductors contraction: with 1 Hz stimulation the ipsilateral hip adductors contract whereas at 50 Hz bilateral contraction of hip adductors occur; and triple flexion response: at 1 Hz stimulation a muscle group contracts in the lower extremities whereas at 50 Hz stimulation, contraction of hip flexors, hamstrings and tibialis anterior occur forming the triple flexion response. In fig 10 the quadriceps muscle responded monosynaptically at 1 Hz but when the stimulation frequency was raised to 50 Hz polysynaptic activity occurred in the tibialis anterior, hamstrings and biceps brachii; simultaneously a triple flexion

Fig 9 CP patients. A: Dorsal root stimulation at 50 Hz EMG recording. B: Dorsal root stimulation at 20 Hz EMG recording on tibialis anterior (upper) and hamstrings (lower).
A response was observed in the lower limb as well as flexion of the elbow.

An unusual observation is shown in fig 9B, where a root was stimulated at 20 Hz and shows polysynaptic activity starting after the burst stimuli have finished in both the hamstrings and tibialis anterior. It should be pointed out that only the hamstrings muscle responds during stimulation showing a monosynaptic response.

B SPINAL SPASTICITY

Ventral root polysynaptic activity was clearly identified following an increase in the strength of stimulation of the DR at 1 Hz (fig 6B-E). Its latency is approximately 4 ms onwards, lasting 14 ms. The time of maximum activity is about 10 ms.

Although bipolar recording does not permit a detailed analysis of the dispersed polysynaptic discharges, it is obvious from the responsiveness cycle (fig 8) that it is absent while the monosynaptic activity is initially enhanced. The polysynaptic responses return at 12–15 ms and are not depressed during the period of profound monosynaptic reflex depression (fig 8).

Clinical correlations

Correlations between the observations made during operation and the clinical features have been made in the 36 cases selected for this study. Fig 11 shows the relationship between the clinical features (topographical extent of spasticity, severity of increase of stretch reflexes and of startle reactions) and the proportion of rootlets evoking pathologic reflex activity.

Twenty-six cases clinically showed spasticity affecting the lower limbs only. The number of rootlets with abnormal reflex activity was above 75% of all the stimulated rootlets in 15 cases (57%). A similar percentage of pathologic rootlets was found in 10 subjects with spasticity affecting all four extremities.
Discrimination

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Lloyd et al. attributed low frequency (below 10 Hz) depression to presynaptic mechanisms and high frequency, (above 10 Hz) depression to motoneurone subnormality, that is to postsynaptic mechanisms. Decandia et al. later proved that failure of transmission with frequencies up to 50 Hz following repetitive activation of Ia afferents is due to presynaptic mechanisms. Two mechanisms are involved in presynaptic depression of the monosynaptic reflex. One is depletion of the available transmitter in the presynaptic terminals, and the other is presynaptic inhibition. The time course of presynaptic inhibition is such that it begins with a latency of about 5 ms, reaches its maximum at about 20 ms and lasts longer than 200 ms.

No reflex effects secondary to muscle spindle contraction seem to be involved, as similar frequency dependent depression has been observed in experiments where both the afferent nerve and the corresponding ventral roots have been sectioned.

The time course of monosynaptic reflex depression we have observed in the roots or rootlets with normal reflex activity (fig 3A) closely resembles their findings, and shows a lack of frequency-related inhibition of the monosynaptic reflex. As Delwaide stated, in cerebral spasticity all the proprioceptive afferent impulses are able to gain direct access to the motoneuronal pool and exert a motor influence. It has been postulated by Delwaide and Ashby et al that presynaptic inhibition is lacking in established spasticity. Our data (fig 3B-D) closely resembles their findings, and shows a lack of frequency-related inhibition of the monosynaptic reflex. As Delwaide stated, in cerebral spasticity all the proprioceptive afferent impulses are able to gain direct access to the motoneuronal pool and exert a motor influence. It has been postulated by Delwaide and Ashby et al that presynaptic inhibition is lacking in established spasticity. Our data suggests that there may be a lack of presynaptic mechanisms on the regulation of Ia afferents in cerebral spasticity, although other mechanisms, such as direct motoneuronal hyperexcitability and reduced postsynaptic inhibition, cannot be excluded.

An important observation from our study is that normal (fig 3A) or even reduced (fig 3E) reflex activity of the monosynaptic arc can be observed in the spinal circuits of subjects with spasticity.

A remarkable feature of the results of DRS in the subject with spinal spasticity is the ability of the monosynaptic arc to follow stimuli to frequencies as high as 400 Hz (fig 7). This correlates with the marked increase (up to 200–300\% of the conditioning response) of the test reflex response during the early

![Diagram of reflex activity](image)

Fig 12 Intraoperative findings of the distributions of reflex activity following 1 Hz and 50 Hz DRS in 3 cases of CP. HF: hip flexors; A: hip adductors; Q: quadriceps; TA: tibialis anterior; TS: triceps surae; HA: hamstrings; UL: upper limb; NT: neck and trunk. Enclosure of muscle groups indicates abnormal reflex responses.

hamstrings) of the lower limbs; bilateral irradiation of the response occurred with right L4 and left L5 stimulation (fig 12).

Case 3 clinically showed a severe spastic tetraparesis mostly affecting hip adductors and hamstrings, with some athetoid features. With 50 Hz stimulation, the pathologic circuits mostly involved the hip adductors, the hamstrings and tibialis anterior; irradiation of the upper limbs, trunk and neck muscles occurred (fig 12).

Discussion

**A. NEUROPHYSIOLOGICAL MECHANISMS**

1. Monosynaptic reflexes: In the anaesthetised cat, early observations by Jefferson et al., Lloyd et al. and, more recently, by Decandia et al., showed that a depression occurs when a monosynaptic reflex pathway is subjected to repetitive stimulation. Similar observations have been made in normal humans.

Lloyd et al. attributed low frequency (below 10 Hz) depression to presynaptic mechanisms and high frequency, (above 10 Hz) depression to motoneurone subnormality, that is to postsynaptic mechanisms. Decandia et al. later proved that failure of transmission with frequencies up to 50 Hz following repetitive activation of Ia afferents is due to presynaptic mechanisms. Two mechanisms are involved in presynaptic depression of the monosynaptic reflex. One is depletion of the available transmitter in the presynaptic terminals, and the other is presynaptic inhibition. The time course of presynaptic inhibition is such that it begins with a latency of about 5 ms, reaches its maximum at about 20 ms and lasts longer than 200 ms.

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facilitatory phase (fig 8). Early facilitation can normally reach 120–160% of the conditioning shock and is normally present only after a supramaximal afferent volley. Such an increase of this reflex activity with supramaximal stimuli is likely to reflect hyperexcitability of the synaptic system or reduced postsynaptic inhibition or both. The remaining part of the excitability cycle following paired supramaximal stimuli (fig 8) shows that early recovery starts at 60 ms and reaches 100% by 90 ms. Studies in normal subjects show that, with strong volleys, recovery from early depression does not begin before 150 ms and does not reach 50% of the conditioning volley. Studies in subjects with spinal spasticity have similarly shown earlier recovery and increased facilitation following early depression.

Mechanisms involved in early depression and subsequent facilitation of the H reflex excitability cycle are complex and still controversial. Presynaptic mechanisms seem to play an important role in depression, while subsequent facilitation has been attributed to long-loop facilitatory reflexes or to an increase of the central excitatory state of alpha motoneurones. Alterations in the first hundred ms of the excitability cycle (fig 8) can therefore be attributed to defective presynaptic mechanisms or excessive activity of the later facilitatory mechanisms or both.

2 Polysynaptic reflexes: DRS in patients with cerebral spasticity has shown an abnormal activation of widespread polysynaptic pathways following repetitive volleys on large myelinated afferents. It is interesting to note that polysynaptic activity is usually absent with low frequency stimulation, and builds up only when the frequency is increased. This means that mechanisms normally preventing diffusion through polysynaptic pathways are reduced and that they are particularly susceptible to temporal summation.

It is to be noted that polysynaptic irradiation can occur in synaptic circuits which do not respond monosynaptically (fig 10). As there is a correlation between the topographical extension of spasticity and the amount of polysynaptic irradiation seen on stimulation of dorsal roots, such a widespread reflex activation of polysynaptic pathways can be responsible for global synkinesias and lack of selective voluntary movement. Similar observations have been made by Dimitrijević et al in spinal spasticity. They, in fact, observed that after a tendon tap, the antagonist may show a prolonged polysynaptic activity even though no activity was induced in the agonist except for its first synchronous discharge of motor units.

It must be pointed out that no uniformity exists in the spinal circuits of spastic subjects at different segmental levels. Different roots or subgroups of rootlets, even in the same subject, can be involved in normal or abnormal reflex activity.

Lack of frequency-related depression of the monosynaptic reflex and abnormal spreading through polysynaptic pathways are two independent variables. This means that either abnormality or both can be present, although to a different extent in the same subject. The great variability of the spinal circuits found in the patients with cerebral palsy derives from the various intermingling of these abnormal reflex activities.

The polysynaptic response observed in the excitability cycle of the subject with spinal spasticity (fig 8) is largely of cutaneous origin and should be normally depressed for over 300 ms after a single conditioning cutaneous volley primarily because of presynaptic inhibition. Our investigations do not permit a quantitative analysis of this discharge, but records show no signs of the pronounced depression which might be expected. As shown by Van Harreveld et al, Murayama et al and Gelfan, asphyxiation of the spinal cord for several minutes leads to profound loss of interneurones and considerable attenuation of polysynaptic activity. The presence of a polysynaptic reflex as shown here indicates the presence of sufficient functional interneurones to mediate a ventral root response, but the absence of an obvious postdischarge depression might reflect the functional loss of interneurones mediating presynaptic inhibition.

This hypothesis must, of course, stand the test of more extensive observation and experimentation than recorded here, but is especially attractive in view of the well known clinical observation that the flexor spasms seen in these patients are frequently triggered by cutaneous stimuli, the spinal input most effectively influenced by presynaptic inhibition.

B CLINICAL CORRELATIONS

The data shown in the paragraph on clinical correlations shows the relationship that exists between the characteristics of the spinal circuits and the clinical features in subjects with cerebral spasticity. The main correlations can be summarised as follows:

1 The distribution of abnormal reflexes produced by lumbar root stimulation correlate closely, in the majority of cases, with the distribution of spasticity seen in the lower extremities.

2 The proportion of lumbar rootlets with abnormal responses is related to the severity of increase of stretch reflexes and not to the topographical extent of spasticity (fig 11). This means that patients with markedly hyperactive stretch reflexes, even if confined to the lower limbs only, have a larger number of pathologic rootlets than subjects with mildly in-
increased stretch reflexes involving all four extremities. 3 Similarly, there is a direct correlation between the number of rootlets with abnormal responses and the severity of increase of startle reactions (fig. 11). 4 There is a correlation between the irradiation of the reflex response to the upper limbs, as elicited at surgery, and the spasticity topography as judged clinically preoperatively. As presented above, eight out of nine cases with polysynaptic irradiation to the upper limbs had a spastic involvement of the upper extremities in the preoperative evaluation.

A main conclusion is that in the reflex motor organisation of each subject with cerebral palsy, the overall pattern of abnormal motor responses can be revealed by activation of spinal circuits at different segmental levels. In the normal subject there are alternatives in the pattern of response to stimuli. The spastic patient particularly with cerebral palsy has, however, limited alternatives because of a stereotype response pattern which controls the segmental reflex activities. In case 1 (fig 12), for instance, which clinically showed extensor hypertonia of the lower extremities, the quadriceps was constantly activated throughout DRS, independently of the segmental level of stimulation.

These observations are similar to what Dimitrijevic et al observed in spinal spasticity. They concluded that two principles govern the involvement of reflex polysynaptic pathways in this condition. They found, in fact, that once certain motoneurones are active, any stimulation will facilitate these motoneurones; at the same time, pathways to certain motoneurones are more easily available, and all stimulation tends to activate the same muscle groups.

It is therefore important to realise that when electrophysiological observations are used to investigate systematically complex reflex activities, continuous correlation with the clinical features is of paramount importance to the understanding of mechanisms of spasticity.

We thank Professor VA Fasano, Director of the Institute of Neurosurgery, University of Torino, for allowing the investigation to be undertaken and for his guidance and assistance, Mr A Squazzi for his extensive assistance, and Dr Ken Casey, University of Michigan, for his assistance with the spastic quadriplegia patient.

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