Motor neurone excitability in back muscles assessed using mechanically evoked reflexes in spinal cord injured patients

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Objective: The clinical and functional assessment of back muscles in human spinal cord injury (SCI) has received little attention. The aim of this study was to develop a method to assess the level of thoracic spinal cord lesion based on the reflex activation of back muscles.

Methods: In 11 control subjects and in 12 subjects with clinically complete thoracic SCI (T2–T12), either a spinous process or an erector spinae muscle was prodded to elicit short latency reflexes recorded electromyographically at the spinal level of stimulation. An electromagnetic servo, attached to a blunt probe, applied stimuli at a frequency of 1 Hz and amplitude of 3 mm. Two trials of 50 mechanical prods were conducted at each site.

Results: Reflexes were evoked in control subjects in 82% of trials when the spinous process was prodded, and in 80% of trials when the muscle was prodded. In contrast, reflexes in SCI subjects could be elicited in 90–100% of trials two segments either above or below the lesion. Reflex responses in control subjects had a mean (SEM) latency of 5.72 (0.53) ms when the spinous process was prodded, and 5.42 (0.42) ms when the muscle was prodded. In the SCI subjects, responses had slightly (but insignificantly) longer latencies both above and below the lesion to either stimulus. The amplitude of reflex responses, expressed as a percentage of the background EMG, was on average 2–3 times larger at the three vertebral levels spanning the lesion in SCI subjects than at sites above or below the lesion or at any level in control subjects.

Conclusion: We propose that the size of these mechanically evoked reflexes may be useful in determining the level of thoracic SCI. Furthermore, the reflexes might provide a valuable tool with which to monitor recovery after an intervention to repair or improve function of a damaged spinal cord.

Spinal cord injury (SCI) can be classified based on the level of lesion and on the completeness (density) of lesion. American Spinal Injury Association (ASIA) assessment scores are widely used to assess the level and density of SCI, but do not provide an accurate picture, especially within the thoracic cord. In addition, the ASIA assessment relies only on the sensory scores to classify thoracic lesions and does not attempt to assess muscles of the trunk. Quantification of treatment outcome is an important feature of assessment scores, and ASIA assessment scores lack the sensitivity to monitor treatment outcome accurately. The need for accurate classification and prediction of outcome of spinal cord injury has become increasingly important with the advent of new cellular and molecular interventional therapies in pre-clinical studies. Recovery from spinal cord injury is influenced by a number of factors, and it is difficult to demonstrate the effect of a single treatment. To identify the various components of recovery, specific neurophysiological tests are required. A number of such neurophysiological tools for assessing spinal cord injury have been assessed recently. In this paper, we investigate the reflex activation of back muscles as the basis for a neurophysiological test for assessment of level of a thoracic lesion in chronic SCI patients that may also be a potential indicator of treatment outcome. The work relates to a previous study, in which it was speculated that the amplitude and latency of short latency reflexes elicited in trunk muscles could prove to be a useful diagnostic tool for evaluating nerve root compressions.

Spinal reflexes are subject to a number of supraspinal controlling influences, with descending inhibition being crucial for the prevention of hyper-reflexia. Disruption to the descending inhibition, for example by a spinal cord lesion, will release the reflex arc from supraspinal control. The mechanism appears to involve reciprocal Ia nerve inhibition, the level of which is related to the amount of spasticity and recovery of function in SCI. This disinhibition of spinal reflexes, resulting in hyper-reflexia or even spasticity, should therefore be useful as an indicator of the level of lesion and also as a tool for monitoring recovery from spinal cord injury. Spinal reflexes can be elicited by activating either the mechanoreceptors present in the interspinous ligaments, or by activating the muscle spindles in paraspinal muscles. We used a mechanical method to preferentially stimulate either mechanoreceptors (interspinous ligaments) or muscle spindles (in ES muscles) at different thoracic levels, and recorded short latency reflex responses in spinal cord injured and control subjects.

METHODS
Subjects
For the study, 11 healthy control individuals (three women, eight men; age range 24–46 years) were recruited. None had ever suffered from a neurological or psychological disorder and none was taking medication. The patient group consisted of 12 subjects (four women, eight men) with spinal cord injury. The level of lesions ranged from T2 to T12. All of these SCI subjects had sustained their injury at least 12 months prior to the investigation, and all were diagnosed with clinically complete lesions. None of the patients was on anti-spasticity medication at the time of investigation. All subjects gave informed written consent. The study conformed to the Declaration of Helsinki, and ethical approval for the study was obtained from the ethics committees of Riverside Research and Stoke Mandeville Hospital NHS Trust.

Abbreviations: ASIA, American Spinal Injury Association; ES, erector spinae; SCI, spinal cord injury
Test sites
For the control subjects, reflexes were tested at neurological levels T2, T7, T11, and L4. For the SCI subjects, at least four sites were tested, one of which was at the level of lesion (according to ASIA assessment). The other sites included levels both above and below the level of lesion. For high (T1–T6) thoracic levels, we chose two sites below and one above the lesion, and for low (T7–T12) thoracic lesions we chose two sites above and one below the level of lesion.

Electromyographic recordings
Surface EMG recordings were made from the right erector spine (ES) muscles at the vertebral level of stimulation. Firstly, the thoracic and upper lumbar spinal processes were identified by palpation and by counting down from the more prominent seventh cervical process. The appropriate levels were marked with a felt tipped pen for easy identification and placement of EMG electrodes. Two self adhesive Ag/AgCl surface EMG electrodes (Arbo Neonatal Blue; 2 cm diameter) were marked over the belly of the muscle (identified using palpation) at positions 3 cm and 6 cm lateral to the spinous process. Four pairs of electrodes were placed lateral to the four chosen spinal processes. EMGs were filtered (~3 dB below 100 Hz and above 2 kHz) and amplified (×10 000) before being sampled (4 kHz) by a computer for storage and analysis (Cambridge Electronic Design 1401; SIGNAL software; IBM compatible PC).

Mechanical stimulation and protocol
Subjects were seated comfortably in a low backed chair or wheelchair, with their arms resting on a table in front of them, allowing their back to remain supported and vertical. A blunt perspex probe with an end diameter of 5 mm was attached to an electromagnetic servo and could be angled to provide mechanical stimulation to a particular spinal process or over the midline of the ES muscle. Stimulation was applied over the spinous process (primarily to activate receptors in interspinous ligaments) or over the ES muscle (primarily to activate muscle spindles) just lateral to the process on the same side as the EMG recording. The electromagnetic servo applied the probe at a frequency of 1 Hz and a throw of 3 mm. Two trials of 50 mechanical prods were conducted at each of the eight probe sites; four on the spinal process and four on the right ES muscle. Electromyographic reflex responses in ES from the recording site adjacent to the prod (that is, at the same vertebral level) were full wave rectified and averaged with respect to the stimuli. To check the degree of mechanical spread of the stimulus, an accelerometer was attached to the adjacent muscle while the spinous process was prodded at the same spinal level. The movement (derived from the accelerometer signal) at the adjacent muscle was <10% of that measured next to the probe; likewise, the movement at the spinous process was <10% of that at the adjacent muscle when the muscle was prodded.

Measurement and analysis
In each of the two trials at each segmental level tested, the 50 EMG responses at the level of stimulation were full wave rectified and averaged. Presence or absence of short latency reflexes was noted in each trial. If necessary, a paired t test comparing pre-stimulus EMG with that recorded at the time of the expected reflex was used to check for presence of a reflex, with p<0.05 taken as statistically significant. Area and latency of the rectified average response was measured for each trial, and an average value noted for each segmental level tested in each subject. The incidence of a reflex at any given level was scored as either 1 (reflexes evident in both trials), 0.5 (reflexes present in only one of the two trials), or 0 (reflexes absent on both trials). Data from above the lesion, at the lesion level, and below the lesion were compared with controls using one way analysis of variance on ranks with Dunn’s correction. Within the patient group, reflex sizes at and below the level of the lesion were examined to see whether they fell outside two standard deviations of the mean reflex size above the lesion.

RESULTS
Raw data
Fig 1 illustrates examples of averaged, rectified reflex responses elicited above, at the level of, and below the lesion in two spinal cord injured patients (both ASIA grade A complete). Panel A shows reflex responses from a patient with a clinical level of lesion level at T4 when the spinous processes were stimulated, and panel B shows reflex responses from a patient with a clinical level of lesion level at T7 when the adjacent muscle sites were stimulated.

Incidence and latency of reflexes
In control subjects, the overall incidence of a reflex (at all levels tested) was 81.8% in response to prodding the process and 79.5% in response to prodding the muscle. In general, the SCI subjects showed a greater incidence of reflex responses compared with controls when the process was prodded above or below the lesion (80–100%) with a similar incidence (80%) at the level of lesion (fig 2A). When the muscle was prodded (fig 2C) reflexes were seen in almost every trial and were only absent in a few cases when tested at one vertebral level below the lesion.

The overall mean (SEM) latency of the response in controls (all four levels tested) was 5.72 (0.53) ms when prodding the spinous process and was 5.42 (0.42) ms while prodding the muscle. Slightly, but insignificantly (p>0.05), longer latency reflexes were recorded SCI patients in response to both types of stimuli at all levels. When the process was prodded (fig 2B) the latency of responses 2–5 segments below the lesion (6.34 (0.3) ms) was no different (p>0.05) to responses at the level of the lesion (6.04 (0.1) ms) or 2–7 segments above the lesion (6.00 (0.25) ms). Likewise, when the muscle was prodded (fig 2D) the latency of responses below the lesion (6.12 (0.57) ms) was no different (p>0.05) to responses at the level of the lesion (6.14 (0.17) ms) or above the lesion (5.93 (0.21) ms).

Size of the reflex responses
Fig 3 compares the size of reflex responses to prodding the spinous process (A) or the muscle (B) as a percentage of an equivalent period of background EMG at different vertebral levels in relation to the spinal cord lesion. The levels tested are aligned with respect to the lesion level. For example, in a patient whose lesion level was T6 and the sites T2, T6, T8, and T10 were prodded, the reflex response from T2 is included in the white bar four segments above the lesion, the response from T6 in the black bar at the lesion level, and below the lesion were compared with controls using one way analysis of variance on ranks with Dunn’s correction. Within the patient group, reflex sizes at and below the level of the lesion were examined to see whether they fell outside two standard deviations of the mean reflex size above the lesion.
Overall, responses at the lesion level, and one segment either side (268 (60%) were significantly larger (p < 0.05) than the control responses (173 (18%), but the pooled responses above (126 (10%) or below (148 (18%) the lesion were no different (p > 0.05) to the controls.

There was a broadly similar result when the muscle was prodded (fig 3B), with hyper-reflexia evident at and around the lesion level. The mean reflex response was 588 (226)% at one segment above the lesion, and 566 (282)% at one segment below the lesion, but rather smaller (239 (75)%), although still hyper-reflexive, at the clinical level of lesion. As for the process prodding, we calculated the mean size of reflexes above the lesion and added two standard deviations of that mean to represent the 95% confidence limits. Five of the 11 patients showed reflexes around the level of lesion that exceeded (p < 0.05) the 95% CI of the responses above the lesion. When compared with control subjects (148 (13%)), the responses at and around the lesion (446 (112%)) were larger (p < 0.05), but there was no difference above (158 (26%)) or below (299 (90%)) the lesion level.

Relationship of reflexes to background EMG in control subjects

Figure 4 plots the size of the reflex response for each spinal level tested against the pre-prod EMG in each control subject while prodding spinous process (A) and the muscle (B). Only

![Figure 1 Reflex EMG responses to prodding a spinous process (A) or the paraspinal muscle (B) in complete SCI patients. Each record represents an average of 50 rectified responses recorded at the same vertebral level as the mechanical stimulus. The stimulus triggered the EMG recording at time zero. (A) Reflex responses evoked by prodding a spinous process from a patient with a clinical level of lesion at T4. (B) Reflex responses evoked by prodding the paraspinal muscle from a patient with a clinical level of lesion at T7. In both cases, the size of the response is clearly larger at the clinical level of the lesion than either above or below the lesion.](http://jnnp.bmj.com/)

![Figure 2 The incidence (A, C) and latencies (B, D) of reflex responses above, at, and below clinically complete spinal cord lesions. (A, B) Reflexes elicited by prodding the spinous process. (C, D) Reflexes elicited by prodding the muscle. In A and C, white bars represent the incidence of responses one or two levels above the lesion, black bars represent responses at level of lesion and grey bars represent responses one or two levels below the lesion. In B and D, white bars represent latencies 2–7 segments above the lesion, black bars represent the lesion level plus one level either side, and grey bars represent 2–5 segments below the level of lesion. The solid lines represent the mean control latencies and the dashed lines the SEM.](http://jnnp.bmj.com/)

![Figure 3 Size of reflex responses at vertebral levels relative to clinical level of lesion. Mean (SE) size of reflex responses as a percentage of the background EMG when prodding the spinous process (A) or the muscle (B) for the SCI subjects. Black bars show reflexes at or within one vertebral level of the lesion; open bars are the reflexes above the lesion; grey bars are the reflexes below the lesion. The mean size of reflexes evoked in control subjects is indicated by the solid line and dashed lines indicate SEM.](http://jnnp.bmj.com/)
while prodding the muscle at T11 was there a significant (linear regression analysis; p<0.05; r² = 0.77) positive correlation between reflex size and the level of pre-prod EMG. At all other spinal levels and at T11 while prodding the spinous process, there was no correlation (p>0.05) between the size of a reflex response and the background EMG. These data, taken alongside the results in fig 3, suggest that the hyperreflexia observed around the level of the lesion in SCI is not simply a consequence of increased resting tone in the muscle; the reflex area, as a percentage of background EMG, is far greater around the level of lesion than at other levels.

DISCUSSION
It has been shown that reflex control of paraspinal musculature is mediated by sensory receptors in the numerous viscoelastic spinal structures such as the intervertebral disc and the spinal ligaments. Short latency reflex activation of the paraspinal muscles can be achieved by stimulating the mechanoreceptors in the passive structures such as ligaments. Reflex activation may also be achieved by stimulating the muscle spindles, giving rise to a monosynaptic stretch reflex. Reflexes in the paraspinal muscles have been studied extensively in the lumbar region, because it has been thought that damaged spinal structures could activate the paraspinal muscles continuously, leading to muscle spasms giving rise to back pain. This hypothesis has been minimally explored in the thoracic levels. In the present study, we have documented the pattern of reflex activation in the paraspinal muscles by stimulating vertebral mechanoreceptors and muscle spindles in the thoracic region in a normal adult population and a group of spinal cord injured patients.

The stimuli applied in this study are not likely to be specific for either mechanoreceptors in the ligaments (spinous process prod) or muscle spindles (muscle prod). When mechanical stimulus spread was investigated using an accelerometer, it was found that movement at the muscle (to spinous process prod) or at the spinous process (to muscle prod) was only 10%, or less, of that recorded at the site of prod. Thus, it is likely that the spinous process prod predominantly activates receptors in the intranspinous ligaments and that muscle prod predominantly activates muscle spindles.

Hyper-reflexia around the level of lesion
Exaggerated stretch reflexes in upper motor neurone disorders have been attributed to hyper-excitability of the motor neurone pool resulting from loss of supra-spinal inhibition. As might be predicted therefore, hyperexcitability has been seen at and around the clinical level of a spinal cord lesion in this study (fig 3). However, while prodding the muscle, the hyper-reflexia was not so marked at the segment reflecting the clinical level of lesion compared with one segment above or below. This could be due to motor neurone loss together with reduced descending inhibition leaving the surviving motor neurones hyper-reflexive. There was considerable variability between SCI patients but, in general, reflex responses were similar to the controls when recorded three or more vertebral segments either above or below the level of the lesion.

Background EMG
It has been clearly established in previous studies that the level of background EMG has a direct relationship to the excitability of the motor neurones. In agreement, there was a weak relationship between the size of the reflex and the amount of background EMG in the control population (fig 4). However, despite the influence of background EMG, the hyper-reflexia is independent of activity in the muscle. Even when the size of the reflexes was expressed as a percentage of the background EMG there is pronounced hyper-reflexia around the level of lesion (fig 3) that cannot be accounted for by the weak relationship observed in the control recordings (fig 4).

Incidence of reflexes
In general the incidence of reflexes in the patients was greater than in the controls (fig 2 A,C). Hyper-reflexia around the level of the lesion in the patients would enhance all responses, making otherwise small reflexes easier to identify; this might partly account for the higher incidence of reflexes. The only exception to the higher incidence of reflexes in SCI was at the clinical level of lesion when the spinous process was prodded (see fig 2). A possible cause for this might be damage to the mechanoreceptors in the spinal ligaments. Ligaments, being passive inelastic structures, are more susceptible to damage in the event of trauma.

Latency of reflex responses
The latencies of the reflexes in spinal cord injury patients were not significantly different from control subjects and did...
not change significantly with respect to the level of lesion (Fig 2). However, on average, the mean latency of response in SCI, at all levels, was slightly longer than in the controls and it is worth considering possible reasons for this. Assuming that there was no physical loss of motor neurones, any increase in latency could be due to an alteration in axonal conduction velocity as observed in other pathologies.29 31 Another consideration is that the erector spinae muscles are supplied by both fast and slow conducting nerve fibres,26 and the recruitment order of motor neurones is correlated with the conduction velocity.7 It has been shown that disuse of the muscles can lead to death of motor neurones, and that the first to succumb are the fast conducting neurones.26 The loss of fast conducting motor neurones below the level of lesion caused by disuse atrophy of the erector spinae muscles7 of the spinal cord.

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The consistent but insignificant increase in reflex latency above the level of the lesion remains unexplained.

Clinical use of the reflexes
The distinct pattern of hyper-reflexia emerging from this study would be a useful tool to monitor the recovery and re-inervation after therapeutic intervention. Regeneration of descending axons following such intervention might re-establish descending inhibition and bring the reflexes back to normal level. Nevertheless, cautious interpretation is required, because at the level of lesion the damage to the inter-vertebral disc, joints and ligaments and loss of motor neurones cannot be reversed. As a result of possible damage to the mecha-noreceptors in the interspinous ligaments, mild hyporeflexia can be expected after re-establishment of descending inhibition.

The erector spinae muscles receive a multi-segmental innervation, but nevertheless show distinct vertebral levels of hyper-reflexia in SCI. A previous study in our laboratory showed that motor evoked potentials in ES (to cortical stimulation) paradoxically appear to occur below the level of a clinically complete lesion,32 and this may be due to the multi-segmental innervation of paravertebral muscles from above the lesion. Although appearing to negate the usefulness of cortically evoked responses in determining the level of SCI, a pattern of increasing cortical magnetic thresholds to transcranial magnetic stimulation was observed related to the level of injury. The short latency reflexes examined in this study show a more definite pattern of change around the clinical level of the lesion, despite this multi-segmental innervation.

CONCLUSION
This study provides evidence that mechanically evoked segmental reflexes in back muscles can be used as a diagnostic tool to help identify the level of complete thoracic spinal cord lesions. In the future, such techniques may prove invaluable for serial monitoring of the status of spinal cord lesions in patients following interventional therapies to repair the spinal cord.

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