Mapping the cortical representation of the lumbar paravertebral muscles

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Abstract
**Objective**

The aim of this study was to map the cortical representation of the lumbar spine paravertebral (LP) muscles in healthy subjects.

**Methods**

Transcranial magnetic stimulation (TMS) was employed to map the cortical representations of the LP muscles at 2 sites. Stimuli were applied to points on a grid representing scalp positions. The amplitude of motor evoked potentials (n=6) were averaged for each position.

**Results**

The optimal site for evoking responses in the contralateral LP muscles was situated 1cm anterior and 4 cm lateral to the vertex. Ipsilateral responses were evoked from sites lateral to the optimal site for evoking contralateral responses. Contralateral responses were also obtained from areas anterior to the optimal site.

**Conclusions**

Maps of these muscles can be produced. The results suggest discrete contra and ipsilateral cortical projections. Anterior sites at which excitation can be evoked may indicate projections arising in the SMA are involved.

**Significance**

This study provides normative data regarding the cortical representation of the paravertebral muscles and provides a technique for evaluating cortical motor plasticity in patients presenting with spinal pathologies.

**Introduction**

Transcranial magnetic stimulation (TMS) has been extensively employed to map the topographic motor cortical representation in a range of muscles (McMillan et al., 1998; Wasserman et al., 1992; Wilson et al., 1993) and to investigate plastic changes in this representation in response to injury, immobilisation and pathology (Rossini & Pauri, 2000; Schwenkreis et al., 2001; Zanette et al., 1997).

The majority of this research has focused on muscles of the extremities. This being due to difficulty in generating reliable maps of the axial musculature. However various studies have identified direct corticospinal pathways to the muscles of the trunk (Nowicky et al., 2001; Ferbert et al., 1992; Plassman & Gandevia, 1989).
The trunk muscles play a key role in the maintenance of upright posture and balance. The corticospinal drive to the trunk muscles has been considered to have a stronger bilateral hemispherical input than that to the muscles of the extremities. Ferbert et al. (1992) identified contralateral and ipsilateral projections to the erector spinae muscle using TMS. This has also been demonstrated in other proximally situated muscles such as the diaphragm (Maskill et al., 1991) and the sternomastoid (Gandevia & Applegate, 1988) and abdominal muscles (Strutton et al., 2004; Tunstill et al., 2001).

Ferbert et al. (1992) employed TMS to investigate the corticospinal projection to the erector spinae muscles of the lumbar spine. They evoked MEPs in contralateral lumbar erector spinae muscles by cortical stimulation in 5 out of 9 subjects but the methodology employed did not provide an accurate location for the cortical representation of erector spinae.

Animal and human studies have demonstrated that nerve injury and cerebral infarcts lead to associated plastic reorganisation of the motor cortex (Toldi et al. 1996, Nudo et al. 1997, Nudo 1999). Changes include shrinkage of the cortical representation synonymous with the area of injury (Toldi et al. 1996), the corresponding area in the uninjured cortex and invasion of adjacent territory by neighbouring representations (Nudo et al. 1997). This motor cortical reorganisation can be influenced by training tasks and studies have shown that functional recovery mirrors further cortical reorganisation (Nudo et al. 1996, Liepert et al. 2000, Schmidlin et al. 2004).

Cortical plasticity of the representation of the back in the somatosensory cortex has been demonstrated in chronic low back pain (Flor et al. 1997) and plasticity in motor pathways of the thoracic paravertebral muscles has been found following spinal cord injury (Cariga et al. 2002). TMS mapping of paraspinal muscles may be a useful tool in the study of plastic changes in the motor cortex following spinal pathology and in monitoring the cortical effects of therapeutic interventions for these disorders.

As no study to date has generated comprehensive maps of the motor representation of the lumbar paravertebral (LP) musculature, the aim was to address this issue, with the objective of developing a robust methodology for the investigation of plasticity in response to pathologies affecting this muscle group and clinical interventions.

Methods

12 healthy volunteers, 2 male, 10 female aged between 19 and 51 years (mean 25 years) were investigated with the approval of the local Ethics committee. All volunteers gave informed consent prior to participation. 10 right hand dominant subjects with no previous history of low back pain or neurological disease were investigated.

The vertex (Cz) was determined using the international 10/20 system and was marked on a latex swimming cap. A line running laterally through Cz from the left to the right
antitragus and a line running sagittally from the external occipital protuberance to the
nasion was drawn. From these reference lines a grid was drawn over the left cortex in
1cm steps antero-posteriorly from 1cm posterior to Cz to 5 cm anterior, and 2cm steps
laterally from Cz to 8cm lateral.

Single stimuli were delivered using a repetitive biphasic Magstim Rapid TMS system
(Magstim Co. Wales) in single pulse mode with a 70mm figure of 8 coil. Stimulation was
applied with the coil handle oriented 45 degrees from the sagittal plane so that the
induced current flowed in an anteromedial direction as suggested by Eguchi et al. (1995)
and Ziemann et al. (1999). Ferbert et al.(1992) used 100% of maximum stimulator output
to elicit responses from these muscles but reduced the stimulus intensity in subjects with
readily indentifiable responses. In the current study pilot work mapping at a range of
stimulator intensities indicated that the most reliable responses were obtained in most
subjects using 100% of maximum stimulator output. Mapping was therefore performed
using this intensity in all subjects.

Surface electromyography was recorded bilaterally from 2 sites: 3cm lateral to the L3
spinal process (recording site 1) and 1cm lateral to the spinous process of L5 (recording
2) using self-adhesive electrodes(Arbo, circular 28mm, Henleys Medical
Supplies, Welwyn Garden City, UK). These sites have been used previously to
successfully record EMG from the paravertebral muscles (Lariviere et al 2003). The
ground electrode was placed over the centre of the sacrum.

EMG signals were recorded from the muscles (Quad 1902 signal conditioner, Cambridge
Electronic Design (CED), Cambridge UK), filtered (2nd order Butterworth, 12dB/octave,
20Hz-2kHz), amplified (1000x), and sampled (4kHz) using an A/D acquisition system
(micro1401, CED) and personal computer running recording and analysis software
(Signal for windows version 2.15 CED). All signals were averaged (n=6) and rectified for
subsequent determination of MEP area. Correct electrode placement was confirmed by
recording EMG whilst subjects performed a resisted back extension movement.

Pilot work identified difficulty in evoking repeatable responses with the muscles at rest.
Subsequently subjects contracted the muscles by sitting forward in a chair and
maintaining extension of the lower back. Biofeedback was given using a light box
calibrated to a subject’s maximum voluntary contraction. Subjects were instructed to
maintain 30% of this maximum during stimulation.

With the subject performing this manoeuvre the average MEP response from 6
stimulations was obtained at each point. Stimuli were delivered with an inter-stimulus
interval of 2-5 seconds. Stimuli were delivered starting at the midline and moving
laterally for each latitude. The subjects were given rest periods between sets of stimuli to
reduce the potential impact of fatigue.

Since facilitation was used, MEP amplitude was recorded after subtraction of the mean
pre-stimulus EMG. The values of the average MEP area for each stimulation site were
calculated and entered into a grid corresponding to the map on the subjects’ cortex. All
responses were normalised by expressing them as a percentage of the peak response obtained from the specific recording site in each subject.

Following the first stage of testing, 3 subjects (all male age 33-64, mean age 46) were recruited to be mapped across both hemispheres. Subjects were mapped using a grid in 1cm steps antero-posteriorly from Cz to 5 cm anterior to Cz, and in 1cm steps laterally from Cz to 4cm lateral to Cz over the left and the right cortex to heighten resolution across the transverse axis. To eliminate possible current spread to the opposite hemisphere due to asymmetrical coil orientation the coil handle was oriented posteriorly in the sagittal plane. As before subjects were stimulated at 100% of maximum stimulator output with facilitation.

**Results**

MEPs were obtained from the contralateral muscles tested in 10 of the 12 subjects. In 2 subjects cortical stimulation evoked no measurable MEPs from the recording sites. It was possible to construct cortical maps for the LP muscles. Figure 1 illustrates the average map obtained for each muscle using the initial mapping protocol.

As in the Ferbert et al. (1992) study, latency of MEPs displayed marked variability both within and between subjects. Table 1 displays latency data for contralateral and ipsilateral responses.

The maps from both recording sites are similar and exhibit considerable overlap. The optimal site for evoking responses was situated 1cm anterior and 4 cm lateral to Cz. The map of contralateral recording site 2 but not recording site 1 reveals another area which produced large responses located 4cm anterior and 2 cm lateral to Cz. On the maps obtained of ipsilateral responses using the initial mapping protocol, large responses were found on the midline with smaller responses evoked from more lateral stimulation sites. Since these responses may have been due to unintentional stimulation of the right hemisphere with the coil orientation of 45°, 3 subjects were mapped as described across both hemispheres using a sagittal coil orientation. Figure 2 illustrates the maps obtained using this protocol.

This mapping protocol produced minimal responses from midline stimulation sites. In the contralateral muscles 2 separate hotspots can be seen at both recording sites. Using this protocol small differences are seen in the optimal site for evoking MEPs with the optimal site located at 2cm anterior and 2cm lateral to Cz for right sided muscles and 1cm anterior and 3 cm lateral for left sided muscles. A separate anteriorly situated hotspot can also be clearly seen. Ipsilateral responses were evoked from lateral stimulation sites.

**Discussion**
The results indicate that it is possible to map the cortical representation of the lumbar paravertebral muscles using TMS. The range of MEP latencies found is consistent with those found in other studies (Ferbert et al., 1992; Nowicky et al., 2001; (Plassman and Gandevia 1989) indicative of a fast-conducting corticospinal pathway.

Analysis of the contralateral maps illustrates that the hotspot at both sites was the same. Due to the multiple peaks found on some maps centre of gravity calculation was not deemed appropriate.

The results support previous findings (Ferbert et al., 1992), of an ipsilateral projection to trunk muscles from the cortex. As with the findings of Ferbert et al. (1992) in the erector spinae and Ziemann et al. (1999) in hand and arm muscles, ipsilateral responses were evoked when stimulating at lateral sites. The midline ipsilateral responses seen using the original mapping protocol may be the result of unintentional stimulation of the opposite cortex since they disappear when mapping using a sagittal coil orientation. Maskill et al. (1991) found ipsilateral responses in the diaphragm which were stronger near the midline but accepted the possibility that these may be due to stimulation of the opposite hemisphere. The current results support this conclusion.

The location of ipsilateral responses and the coil orientation used in this experiment makes it unlikely that they are the result of current spread to the opposite hemisphere. Ipsilateral responses had a longer latency than contralateral responses of around 4 msecs. These findings are in agreement with those of Ziemann et al. (1999) who demonstrated slower ipsilateral responses in upper limb muscles obtained more prominently from sites lateral to the optimal stimulation site for contralateral responses.

The small differences observed between the hotspots using the different mapping protocols may be the result of the alteration in coil orientation and thus current direction. The increased resolution of the map grid in the bilateral protocol may also have contributed towards this.

The presence of an excitable area of the cortex from anterior stimulation sites is an interesting finding. The bilateral mapping protocol demonstrates that this anterior site is only seen clearly in contralateral muscles. Large responses evoked from relatively anterior stimulation sites may indicate excitation of the supplementary motor area (SMA). Studies have confirmed a somatotopic organization of the pre-motor area (PMA) and the SMA (Fink et al., 1997; Godschalk et al., 1995). Sharshar et al. (2004) demonstrated a discrete anterior site for evoking MEPs in the diaphragm and suggest that this anterior area may represent the SMA. In the current study the latency of large responses obtained from anterior sites was not significantly different to that found at the hotspot. As with the findings of Sharshar et al. (2004) the short latency responses found from anterior stimulation sites in the current study suggest a direct projection from the SMA. Imaging studies would be of use in determining whether these anterior spots do represent the SMA.
The variability seen in the latencies of MEPs may reflect the activation of differing pathways to these muscles due to the necessary use of high stimulation intensity with active motor facilitation. The difficulty in determining onset latency in the presence of background EMG may mitigate this variability. The shorter latency responses may reflect a fast-conducting direct motor pathway. Longer latency responses may indicate an oligosynaptic pathway.

The wide distribution of responses observed and the finding of multiple peaks in some maps limits the use of CoG in accurately ascertaining map position. Map area was not analysed for similar reasons. This is primarily due to the chosen protocol of stimulating all subjects at 100% of stimulator output with voluntary activation of the muscle. The bilateral mapping methodology using a sagittal coil orientation produced good quality maps from which ipsi and contralateral responses could be clearly distinguished. It is therefore recommended that future mapping studies for these muscles adopt this approach.

It is important to consider what influence the use of a biphasic stimulator may have had on the results. Studies have shown that biphasic stimulation may excite larger populations of neurons than monophasic stimulation and that the resting motor threshold for biphasic stimulation is lower than that for monophasic stimulation (Arai et al. 2005, Sommer et al. 2002). Thus maps obtained with monophasic stimulation may vary from those found in the current study. Since biphasic stimulation presents a more powerful stimulus it may be more successful in elicting MEP’s from these muscles from a wider cortical area. Given the technical difficulties encountered in obtaining repeatable responses from these muscles at lower stimulus intensities it appears unlikely that monophasic stimulation would provide more information at these intensities. The influence of different stimulation waveforms on the results of cortical mapping is an interesting direction for further study.

Using this methodology the parameters of hotspot location, response latency and MEP amplitude may provide parameters by which plastic changes may be investigated and monitored in response to pathology and therapeutic intervention.

References


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Plassman BL, Gandevia SC. Comparison of human motor cortical projections to abdominal muscles and intrinsic muscles of the hand. Exp Brain Res 1989;78:301-308


Strutton PH, Beith ID, Theodorou S et al. Corticospinal activation of internal oblique muscles has a strong ipsilateral component and can be lateralized in man. Exp Brain Res 2004;158:474-479


Figures and Tables

Figure 1. Averaged cortical maps (n=10) for the 4 recording sites using the initial mapping protocol of stimulating the left cortex with the coil handle oriented at 45°. %MR= amplitude expressed as a percentage of the maximum response obtained from that recording site.
Figure 2. Averaged cortical maps (n=3) for the 4 recording sites using the second mapping protocol of stimulating the left and right cortex with the coil handle oriented in the sagittal plane.

Table 1. Mean (±Standard Deviation) and range of MEP latencies (msecs) obtained from the optimal site for obtaining responses at each recording site in each subject, and for all responses obtained in each subject.

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<tr>
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<th>Contralateral Responses</th>
<th>Ipsilateral Responses</th>
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<tbody>
<tr>
<td>Mean latency at optimal site</td>
<td>19.53 ± 4.78</td>
<td>24.19 ± 3.77</td>
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<tr>
<td>Range of latencies at optimal site</td>
<td>12.10 – 27.70</td>
<td>18.60 – 31.53</td>
</tr>
<tr>
<td>Mean Latency all responses</td>
<td>19.72 ± 4.41</td>
<td>24.3 ± 5.97</td>
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<tr>
<td>Range of latencies all responses</td>
<td>12.05 - 30.48</td>
<td>14.06 - 36.45</td>
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