The vestibular control of balance after stroke

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Objectives: To examine vestibular control of balance in those who recovered the ability to stand after middle cerebral artery (MCA) stroke.

Methods: Sixteen patients with MCA stroke were compared with 10 age matched controls. Two additional patients were studied with isolated corticospinal tract lesions, one each at the level of the pons and medulla. Vestibular evoked postural responses were obtained using galvanic vestibular stimulation (GVS) while patients stood with their eyes closed and head facing forwards, equally loading both legs. The GVS response was characterised by measuring the amplitude of the stimulus evoked lateral forces acting through each leg and the lateral displacement of the axial skeleton.

Results: Lateral displacement and net lateral force following GVS were significantly larger after stroke. Unlike controls, the lateral forces in the stroke group were asymmetrical, being enhanced on the side of the non-paretic limb and small on the side of the paretic limb. The degree of GVS evoked asymmetry correlated with corticospinal damage assessed using transcranial magnetic stimulation. A similar asymmetrical response was seen in the patient with the pontine lesion but not the patient with the medullary lesion.

Conclusions: MCA stroke may disrupt corticobulbar projections to brainstem output pathways involved in vestibular control of balance. These projections are either collaterals of the corticospinal tract or lie close to that tract and terminate in the pons/upper medulla. This hypothesis accounts for the association between corticospinal tract damage and GVS response asymmetry, and the lack of GVS evoked asymmetry with corticospinal lesions below the rostral medulla.
which was attached to an overhead gantry capable of low friction movement in the horizontal plane.

**Single pulse, transcranial magnetic stimulation procedure**

The motor evoked potential (MEP) elicited in the preactivated tibialis anterior muscle was assessed. Subjects sat with their foot plantigrade in a custom made manipulandum. The axis of ankle dors/plantarflexion was collinear with the axis of rotation of the manipulandum. Subjects isometrically dorsiflexed the foot to a constant torque of 0.5 Nm, as measured via a strain gauge. Visual feedback of ankle torque and the required target value was provided on an oscilloscope. Muscle activity from the gastrocnemius was recorded via surface electromyography (EMG) to ensure that no co-con traction occurred during this task.

Single pulse TMS was given via a figure of eight coil placed on the vertex, with the current acting in an antero–posterior direction. Initially, the threshold for stimulation (defined as the value that gave a response in three of five stimuli) was determined on the non–paretic side during isometric dorsiflexion. Five stimuli were then recorded at ×1.5 motor threshold. After this, the paretic limb was assessed using the same coil position and stimulus intensities as for the non–paretic side with the ankle in an identical position, dorsiflexing to the same constant torque. The maximal motor response was then elicited in the tibialis anterior by stimulating the common peroneal nerve at the level of the head of the fibula, with the tibialis anterior muscle relaxed. Three stimuli were recorded for the left and the right legs. Tibialis anterior muscle responses were recorded via surface EMG (MT8 MIE; Medical Research, Leeds, UK). Signals were sampled at 2 kHz after amplification.

**Clinical measures**

The time taken and the number of paces required to walk 10 m at the patients’ normal pace were recorded. Lower limb muscle strength was manually tested and graded on a 0–5 ordinal scale. The summed scores for the hip flexors, adductors, and extensors, the knee extensors and flexors, and the ankle dorsiflexors and plantarflexors were calculated, giving a total maximum score of 35. Distal lower limb sensation was tested using the Rivermead assessment of somatosensory performance (RASP; Thames Valley Test Company Ltd, Bury St Edmunds, UK). This assessed light touch on the plantar and dorsal aspects of the foot (test 2) and proprioceptive awareness at the ankle and hallux (test 7).

**Galvanic vestibular stimulation procedure**

Subjects stood barefoot with each foot on a separate force plate with a 5 cm gap between the medial borders of the feet. A visual display 1 m in front of the subjects gave feedback in 5% steps about the percentage of body weight being taken by the right leg. After a warning tone, a central light appeared above the feedback display that indicated the point when the legs took equal weight. Once subjects had achieved equal lower limb weight bearing with the aid of the feedback display, they were instructed to close their eyes and the trial was started. After a 0.5 to two second random delay, data collection began. After a three second baseline period, a three second, 1.0 mA monaural galvanic vestibular stimulus was given, which was then followed by a further three second period. The stimulus was applied using 2.5 cm diameter electrodes; the active electrode was placed on the mastoid process with an indifferent electrode on the back at the level of T1 spinous process (PALS plus; Nidd Valley Medical, Knaresborough, North Yorkshire, UK). There were four stimulus conditions. The active electrode on the mastoid process could either be on the same or opposite side to the lesion and, by altering the stimulus polarity, it could produce a sway either towards or away from the side of the lesion (table 1). The order of stimulus presentation was randomised, with 10 stimuli being given for each condition.

Axial displacement was measured using a three dimensional motion analysis system (CODA MPX30 system; Charnwood Dynamics, Rothley, Leicestershire, UK) via markers attached to a helmet, semirigid belt, and to the back at the level of the C7 and L3 spinous processes. Lateral reaction forces were obtained from the force plates under each foot (Kistler 9281B, left leg; Kistler 9287, right leg). Throughout the experiment, subjects wore a safety harness,

### Table 1  Stimulus conditions

<table>
<thead>
<tr>
<th>Side of stimulation</th>
<th>Ipsilateral to lesion</th>
<th>Contralateral to lesion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sway direction</td>
<td>Anode</td>
<td>Cathode</td>
</tr>
<tr>
<td></td>
<td>Ipsilateral to lesion</td>
<td>Contralateral to lesion</td>
</tr>
</tbody>
</table>

The table indicates the relationship between the site and polarity of the active electrode and the stimulus evoked sway direction. The site of stimulation and the stimulus evoked sway direction are given relative to the side of the lesion.

**METHODS**

**Clinical details**

Patients with an MCA stroke were recruited from the National Hospital for Neurology and Neurosurgery, London, UK, and from a local stroke self help group. Subjects were recruited if they were able to walk 10 m independently, with or without the use of a gait aid, at the time of our study, although all had been unable to stand or walk independently in the acute stages after the stroke. Exclusion criteria were the presence of a previous stroke or other neurological or orthopaedic impairments. Age and sex matched healthy subjects with no history of neurological or orthopaedic impairments were recruited as controls. In addition, we assessed two subjects who had discrete lesions predominately affecting the pyramidal tract, one within the pons and the other in the medulla, to elucidate further the role of corticofugal projections to brainstem pathways. All subjects participated with informed consent and the approval of the local ethics committee according to the Declaration of Helsinki.
For each trial, the mediolateral postural sway was measured over the baseline period. This was defined as the standard deviation of the mediolateral velocity of the C7 marker following 20 Hz low pass filtering. The lateral reaction forces and the lateral displacement of the axial markers were used to measure the GVS evoked postural response. Any baseline drift in the axial displacement or reaction force in each trial was removed by subtracting the slope over the baseline period, estimated by a least squares linear regression, from the complete trace. Secondary analysis was then performed on the average of the 10 trials for each condition.

The initial lateral force response was characterised by determining the change in impulse (force × time) from 320–500 ms after stimulus onset. The average lateral displacement from 520–700 ms post stimulus onset of the head, C7, and pelvic markers were calculated (see grey bars in fig 1A). The initial lateral reaction force response averaged across all stimulus conditions was also used to calculate an asymmetry index, which was defined as: (P − NP)/(P + NP), where P and NP refer to the paretic and non-paretic leg responses, respectively. In some subjects with stroke, the paretic leg and NP refer to the paretic and non-paretic leg responses, respectively. The asymmetry index was defined as: (P − NP)/(P + NP), where NP indicates the amplitude of the non-paretic leg and P indicates the amplitude of the paretic leg. As with the GVS asymmetry index, this is a bounded measure taking values from −1 to 1, where 0 indicates symmetrical responses between the two limbs.

**Statistical analysis**

The parameters measured were analysed using a between groups general linear model in SPSS (version 11). The response to the four GVS conditions was analysed using the stimulus side relative to the lesion (two levels—ipsilateral or contralateral) and the sway direction relative to the lesion. The mean (SD) is presented, except for the measures of muscle strength and sensory function, where the median and interquartile range is given. The total scores indicate the maximum score possible for this test. NP and P indicate the scores for the non-paretic and the paretic legs, respectively.

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**Table 2 Individual clinical details**

<table>
<thead>
<tr>
<th>Subject</th>
<th>Age (years)</th>
<th>Sex</th>
<th>Time since lesion (months)</th>
<th>Lesion location</th>
<th>GVS AI</th>
<th>TMS delay (ms)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>47</td>
<td>M</td>
<td>90</td>
<td>L MCA infarct, fronto–temporo–parietal</td>
<td>−1</td>
<td>NA</td>
</tr>
<tr>
<td>2</td>
<td>19</td>
<td>M</td>
<td>22</td>
<td>R MCA haemorrhage, fronto–temporo–parietal</td>
<td>−1</td>
<td>NA</td>
</tr>
<tr>
<td>3</td>
<td>76</td>
<td>M</td>
<td>32</td>
<td>L thalamic and IC infarct</td>
<td>−1</td>
<td>15.5</td>
</tr>
<tr>
<td>4</td>
<td>65</td>
<td>M</td>
<td>12</td>
<td>L anterior artery and anterior MCA infarct, fronto–parietal</td>
<td>−1</td>
<td>12.5</td>
</tr>
<tr>
<td>5</td>
<td>62</td>
<td>F</td>
<td>25</td>
<td>R anterior and posterior limbs of IC and caudate</td>
<td>−0.70</td>
<td>1.5</td>
</tr>
<tr>
<td>6</td>
<td>66</td>
<td>M</td>
<td>24</td>
<td>R anterior watershed territory infarct</td>
<td>−0.64</td>
<td>7</td>
</tr>
<tr>
<td>7</td>
<td>33</td>
<td>M</td>
<td>29</td>
<td>L MCA, fronto–parietal</td>
<td>−0.57</td>
<td>5</td>
</tr>
<tr>
<td>8</td>
<td>55</td>
<td>M</td>
<td>24</td>
<td>R MCA, temporo–parietal infarct</td>
<td>−0.56</td>
<td>7</td>
</tr>
<tr>
<td>9</td>
<td>46</td>
<td>M</td>
<td>28</td>
<td>L focal lesion at the junction of the thalamus and the posterior limb of the IC</td>
<td>−0.46</td>
<td>7.5</td>
</tr>
<tr>
<td>10</td>
<td>40</td>
<td>F</td>
<td>12</td>
<td>Left lentiform/head of caudate and anterior and posterior IC infarct</td>
<td>−0.41</td>
<td>3.5</td>
</tr>
<tr>
<td>11</td>
<td>66</td>
<td>M</td>
<td>25</td>
<td>R caudate nucleus infarct extending into the corona radiate</td>
<td>−0.35</td>
<td>10</td>
</tr>
<tr>
<td>12</td>
<td>68</td>
<td>M</td>
<td>19</td>
<td>R posterior limb IC</td>
<td>−0.32</td>
<td>6.5</td>
</tr>
<tr>
<td>13</td>
<td>53</td>
<td>F</td>
<td>46</td>
<td>R MCA temporo–frontal infarct extending into basal ganglia</td>
<td>−0.28</td>
<td>4</td>
</tr>
<tr>
<td>14</td>
<td>53</td>
<td>M</td>
<td>24</td>
<td>R posterior right lentiform nucleus and the adjacent IC infarct</td>
<td>−0.25</td>
<td>7</td>
</tr>
<tr>
<td>15</td>
<td>55</td>
<td>M</td>
<td>37</td>
<td>R MCA</td>
<td>−0.21</td>
<td>2.5</td>
</tr>
<tr>
<td>16</td>
<td>70</td>
<td>M</td>
<td>45</td>
<td>R caudate and putamen infarction with secondary degeneration of the right cerebral peduncles and pons</td>
<td>0.23</td>
<td>−3.5</td>
</tr>
</tbody>
</table>

F, female; GVS AI, galvanic vestibular stimulation asymmetry index; IC, internal capsule; L, left; M, male; MCA, middle cerebral artery stroke; NA, not assessed; R, right; TMS, transcranial magnetic stimulation.

**Table 3 Group clinical details**

<table>
<thead>
<tr>
<th>Group</th>
<th>M/F</th>
<th>Age (years)</th>
<th>Mass (kg)</th>
<th>Height (m)</th>
<th>10 m timed walk</th>
<th>Muscle strength NP (n = 35)</th>
<th>Muscle strength P (n = 35)</th>
<th>Sensation test 7P</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Time (s)</td>
<td>Steps</td>
<td>(n = 35)</td>
<td>(n = 35)</td>
</tr>
<tr>
<td>R stroke</td>
<td>8/2</td>
<td>57.0 (13.9)</td>
<td>72.9 (12.4)</td>
<td>1.65 (0.26)</td>
<td>24 (2.3)</td>
<td>22.4 (15.4)</td>
<td>23.3 (8.5)</td>
<td>34 (2)</td>
</tr>
<tr>
<td>L stroke</td>
<td>5/1</td>
<td>51.2 (16.2)</td>
<td>79.6 (16.7)</td>
<td>1.69 (0.12)</td>
<td>23 (5)</td>
<td>23.5 (21.6)</td>
<td>24.6 (11.8)</td>
<td>33.5 (2.5)</td>
</tr>
<tr>
<td>Controls</td>
<td>7/3</td>
<td>57.3 (13.2)</td>
<td>85.7 (15.3)</td>
<td>1.72 (0.10)</td>
<td>24 (0)</td>
<td>7.8 (0.8)*</td>
<td>14.3 (1.8)*</td>
<td>35 (5)</td>
</tr>
</tbody>
</table>

The mean (SD) is presented, except for the measures of muscle strength and sensory function, where the median and interquartile range is given. The total scores indicate the maximum score possible for this test. NP and P indicate the scores for the non-paretic and the paretic legs, respectively.

*The stroke group (left and right sided lesions combined) was significantly different from the control group, as measured using a two tailed unpaired t test. †The stroke group was significantly different from the control group, as measured using a Mann-Whitney test.

L, left; R, right.
Results
Clinical details
An ischaemic stroke occurred in 15 patients and a haemorrhagic stroke occurred in one patient. Patients were seen on average 30.9 (SEM, 4.6) months after stroke onset and had a mean age of 54.6 (SEM, 3.8) years. Both cortical and subcortical lesions were present as defined on computed tomography or magnetic resonance imaging (MRI) scans (table 2). There was no difference between the stroke and control groups in terms of age (two tailed unpaired t test, t(24) = ± 0.37; p > 0.05), mass (t(24) = ± 1.7; p > 0.05), or height (t(24) = ± 0.6; p > 0.05; table 3).

All patients had recovered the ability to walk in the period one to three months after stroke, although as indicated in table 3, deficits in walking speed were still apparent. The patients with stroke also presented with lower limb weakness and sensory deficits (table 3) compared with the control subjects. No patient had signs of spatial neglect as assessed by a psychologist at the time of initial admission.

Response to GVS
The legs were equally loaded over the baseline period. The paretic leg was loaded by 48.9% of body weight in the stroke group (SEM, 1.1%) and 50.7% (SEM, 0.5%) in the control group (t(24) = 1.3; p > 0.05). There was a tendency for stroke patients to shift their weight slowly on to the non-paretic leg over the baseline period. However, this effect was small, and there was no significant difference in the average load taken by the paretic limb over the first 200 ms and the last 200 ms of the baseline period. The degree of mediolateral postural sway over the baseline period, as measured from the C7 marker, was larger in the stroke group (two tailed t test, t(24) = ± 2.8; p < 0.05).

Net response characteristics
In response to GVS, all subjects swayed away from the active electrode if it was a cathode or towards the active electrode if it was an anode. The displacement of the axial markers following GVS was significantly larger in the stroke group (group factor: head, F(1,24) = 5.9; p < 0.05; C7, F(1,24) = 5.7; p < 0.05; pelvis, F(1,24) = 6.8; p < 0.05). The total summated lateral reaction force acting on the body...
that is, the sum of the left and right leg responses) was also higher in the stroke group (group factor: F(1,24) = 10.4; p < 0.005). However, there was no significant effect of stimulus side or sway direction on either of these measures. Figure 1 summarises the effect of GVS on the summated lateral reaction forces and axial displacement after averaging across all stimulus conditions.

Individual forces
The GVS evoked lateral forces acting through each leg were asymmetrical after stroke. The initial lateral impulse (measured from 320–500 ms after stimulus onset) was higher on the non-paretic side than on the paretic side, whereas such consistent asymmetries were not seen in the control group (group x leg interaction, F(1,24) = 6.87; p < 0.05; fig 2A and B). There were no effects of stimulus side, sway direction, or other interactions. This indicated that the stroke group's asymmetrical response was present regardless of the stimulus condition. Therefore, further post hoc analysis was performed on the lateral forces after averaging across all stimulus conditions.

Figure 2 Differences between groups in the galvanic vestibular stimulation (GVS) evoked lateral forces acting through each leg. (A) The grand average response of the individual lateral reaction forces is shown averaged across all stimulus conditions for the stroke (n = 16) and control (n = 10) groups. (B) Mean (SEM) lateral impulse acting through each leg 320–500 ms after stimulus onset. NP, non-paretic; P, paretic.

Response to single pulse TMS
Only 14 patients in the stroke group participated in this part of the experiment. Two were unable to participate because of a history of epilepsy and the presence of intracranial metal. The mean (SEM) stimulation threshold in terms of maximum output of the stimulator was 36.8% (1.9%) for the stroke subjects and 34.0% (2.9%) for the control subjects. The amplitude of the response revealed a significant group x leg interaction (F(1,22) = 5.0; p < 0.05). Post hoc analysis showed no difference between groups on the non-paretic side (unpaired t test, t(24) = −0.57; p < 0.05), whereas the response was significantly smaller for the stroke group on the paretic side (unpaired t test, t(24) = −2.7; p < 0.05; fig 3A). The asymmetry index of response amplitude was different for the two groups (stroke: mean, 0.29; SEM, 0.06; control: −0.03; SEM, 0.09; unpaired t test, t(22) = 2.7; p < 0.05).

In the stroke group, the TMS response was delayed on the paretic side by 7.1 ms (SEM, 1.3; table 2), compared with 0.85 ms (SEM, 0.5) for the control subjects (unpaired two tailed t test, t(22) = 4.5; p < 0.0005; fig 3B).

Correlations between the responses seen after GVS and TMS
There was a significant correlation between the asymmetry index seen after GVS and the relative delay in the MEP after TMS (R² = 0.61; p < 0.001; fig 3C). The linear regression model was not significantly improved by using measures of TMS response amplitude asymmetry, muscle strength, or hemiplegic limb sensory deficit as predictors. Highly
asymmetrical responses, with a GVS asymmetry index more negative than −0.8, were seen even when clinical testing of proprioception at the ankle and hallux was intact. There was no observed effect of lesion location (cortical versus subcortical) or period since stroke on the response asymmetry seen with GVS.

**Deficits in the GVS response after discrete pyramidal lesions**

To investigate further the relation between corticospinal tract damage and the asymmetrical response to GVS, two additional patients with lesions predominately affecting the pyramidal tract were assessed. Importantly, these pyramidal tract lesions were either rostral or caudal to the vestibular nuclei (fig 4).

The first patient was a 29 year old man who had had a left anterior pontine lacunar infarct 33 months previously (fig 4A). He initially presented with a right upper motor neurone facial palsy and a right sided hemiplegia. There was no history of vertigo and ocular movements were normal. Lower limb sensation was intact during clinical testing in the immediate period after stroke and when tested at the time of our study. The response to GVS was similar to that seen after MCA stroke (fig 5A and B). The postural sway in response to GVS was enhanced, whereas the individual lateral reaction forces were asymmetrical, as highlighted by an asymmetry index (star symbol in fig 3C).

This initial period was before appreciable movements of the head and axial skeleton. Therefore, the initial response is probably purely vestibular in origin, and uncomplicated by movement relatedafferent feedback or the passive biomechanical effects of positional change.

Moreover, the signal to noise ratio of the force response was superior to that of an EMG response, requiring the averaging of far fewer trials to obtain a reasonable response estimate, which is an important consideration when studying disabled subjects. The reaction forces were measured over a short period from 320–500 ms after the onset of the stimulus. This initial period was before appreciable movements of the head and axial skeleton. Therefore, the initial response is probably purely vestibular in origin, and uncomplicated by movement related afferent feedback or the passive biomechanical effects of positional change.

The main finding of our study was an abnormal inter-leg response asymmetry to GVS in the stroke group. We characterised the response primarily by measuring the change in the lateral force produced through each leg, rather than the electromyographic response of individual muscles. There are some advantages to this approach because the force represents the net result of all the distributed muscle activities and can readily be related to function. Furthermore, the signal to noise ratio of the force response is superior to that of an EMG response, requiring the averaging of far fewer trials to obtain a reasonable response estimate, which is an important consideration when studying disabled subjects. The reaction forces were measured over a short period from 320–500 ms after the onset of the stimulus. This initial period was before appreciable movements of the head and axial skeleton. Therefore, the initial response is probably purely vestibular in origin, and uncomplicated by movement related afferent feedback or the passive biomechanical effects of positional change.

The use of monaural stimulation allowed us to investigate whether there was any difference in response processing when the stimulus was applied either ipsilateral or contralateral to the lesion. In addition, by varying the polarity of stimulation we were able to distinguish whether there were any differences in the response when the subject swayed towards or away from the side of the lesion. However, regardless of the side of stimulation or the subsequent sway direction, we found no difference in the pattern or amplitude of the inter-leg response asymmetry. Thus, stroke was associated with a lateralised deficit in the motor output stage of vestibular processing, rather than in the sensory or spatial processing stages.

**DISCUSSION**

Figure 4 Lesion location in two patients with discrete corticospinal lesions. (A) Horizontal magnetic resonance imaging (MRI) scan indicating a left anterior pontine lacunar infarct in subject 1 (arrow). (B) Horizontal MRI of the medulla indicating a right vertebral artery aneurysm compressing the right pyramid in subject 2 (arrow).
Previous studies of standing balance after stroke have found deficits in the motor control of the paretic leg. In our present study, there was no significant difference in the response amplitude of the stroke group’s paretic limbs compared with the healthy limbs of control subjects. The obvious difference was in the behaviour of the non-paretic leg, which appeared to over respond. However, there are two factors that complicate this interpretation. First, the feet were placed 5 cm apart to enable the stroke subjects to stand independently with their eyes closed. In control subjects, relatively small increases in stance width, compared with when the feet are together, have the effect of greatly reducing the amplitude of the response. Under these circumstances, it may be difficult to establish a reduction in paretic limb response size when the control response is already very small. On top of this, the stroke group’s baseline body sway was greater than the control group, indicating that they were less stable. It is well established that conditions that increase instability and background body sway, such as the removal of visual or tactile cues, or neurological disease, are associated with enlarged responses to GVS. Again, the effect of this would be to mask any differences between an underresponding paretic limb and a healthy limb, while at the same time accentuating the apparent over-responsiveness of the non-paretic limb. What is clear, however, is that the overall whole body response was abnormally asymmetrical in the stroke group; if a small response was seen in a limb of a control subject then it occurred symmetrically in both legs.

Our results suggest that a chronic balance deficit can arise as a consequence of stroke. The two legs no longer contribute equally to the balance process, even when they are taking an equal amount of body weight. The responsibility for balance control seems to be shifted away from the paretic leg and towards the relatively unaffected leg. The correlation data suggest that this phenomenon is not simply a reflection of muscle weakness or sensory loss contralateral to the lesion. However, we did find that the GVS asymmetry index correlated with the relative delay in MEP onset latency after TMS. Experimental models of stroke in the monkey suggest that the delay in MEP onset predominately reflects the number of fibres that have been damaged. Ischaemia causes preferential damage of the larger, faster conducting corticospinal neurones, resulting in a population with an overall reduced conduction velocity and a delay in onset latency. Thus, the relative delay in TMS onset after stroke may serve as an indicator of the degree of damage to motor cortical output neurones. It has previously been suggested that the balance response to GVS may be mediated via brainstem nuclei. We suggest that MCA stroke produces its effect by disrupting projections from the cortex of one hemisphere to these brainstem motor centres. The results of the two patients with discrete pyramidal lesions support this view. It predicts that such corticobulbar projections may be affected by a lesion rostral to the lower pons, but not by a lesion in the medulla below the level of the vestibular nuclei, which was found to be the case. Thus, for the patient with the pontine lesion, the responses to both GVS and TMS were asymmetrical to a degree similar to that seen after an MCA lesion. In contrast, for the patient with the medullary lesion, the response to GVS was symmetrical, and of a similar magnitude to that seen in control subjects, whereas the TMS response was asymmetrical.

The proposed corticobulbar projections could be either collaterals of the corticospinal tract or separate neurones that lie spatially close to it, explaining the association between the GVS evoked asymmetry and the degree of corticospinal tract damage. Such a spatial proximity between the corticospinal tract and corticobulbar projections is seen for example in the projections to the facial nucleus. Corticobulbar connections from the cortex to the reticular or vestibular nuclei have been found in non-human species. For example, in the cat and primate, collaterals from the corticospinal tract, in addition to separate pathways, arise bilaterally from layer V of the primary and premotor cortices and project to the pontomedullary reticular formation. Similarly, cortical projections
from the contralateral premotor cortex to the lateral vestibular nucleus have been identified in old and new world monkeys, and are thought to modulate vestibulomotor reflex arcs. Cortical projections to the vestibular nuclei also arise from the parietal lobe. Lesions to the parietal multisensory cortex can result in deficits in balance and the perception of verticality. Therefore, these projections may also play a role in the vestibular control of balance, in addition to their known role in the control of ocular movements. The deficits in balance and verticality perception seen after parietal lesions are often associated with signs of visuo-spatial neglect, a symptom that was absent in our present cohort of patients. Furthermore, none of our patients showed signs of pushing away from the non-parietal side, a symptom often associated with acute temporoparietal lesions. Nevertheless, the patients with large GVS asymmetry indices had lesions that affected both the frontoparietal and premotor cortex (table 2), so it is possible that these proposed corticobulbar projections could arise from either the frontal or parietal lobes.

Disruption of the proposed corticobulbar connections could indirectly result in a reduced response on the parietic side. Reciprocal connections between brain stem centres each side of the neuraxis could explain the response asymmetry seen between the two sides. For example, the vestibular nuclei are reciprocally connected via inhibitory commissural connections. Here, a lesion resulting in a decrease in activity on one side could disinhibit the opposite side and thereby increase its excitability. Such asymmetries are seen after a unilateral vestibular nerve lesion. Therefore, this proposal provides a possible explanation for the asymmetrical response to GVS seen in the stroke group. Because the degree of asymmetry would depend upon the extent of the damage to cortical output pathways, it may also explain the association between the GVS evoked response asymmetry and the TMS response delay.

There are other possible pathophysiological mechanisms that could explain the asymmetrical GVS response. Vestibular stimuli directly activate an interconnected circuit of cortical areas within the parietal, temporal, and frontal lobes. Therefore, the postural response evoked by GVS could be mediated through a trans cortical pathway, whose output is via the corticospinal tract. This could explain the GVS asymmetry seen with an MCA lesion and the association with corticospinal tract damage. Another possibility is that the spinal output is altered by the loss of cortical inputs and so responds abnormally to activity in other descending pathways. However, the finding of a symmetrical response to GVS after a corticospinal tract lesion at the level of the medulla argues against these possibilities.

In conclusion, we have shown that MCA stroke disrupts the vestibular channel of balance control. We propose that stroke may produce this effect by interrupting corticobulbar modulation of brainstem balance centres. We assume that the neurones responsible lie spatially close to the corticospinal tract, such that damage to the corticospinal tract acts as a marker of damage to the proposed corticobulbar pathway. The present data do not indicate whether the vestibular motor changes are immediate consequences of stroke or are the result of longer term compensatory processes. This can only be answered by studying subjects at much shorter intervals after stroke. Nevertheless, the asymmetry that we have described may contribute to the asymmetries in standing balance often reported in stroke patients.

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REFERENCES
Cannabis may cause psychosis in young people

A large prospective population based study in young people has confirmed that using cannabis risks psychosis developing later—greatly for those susceptible to psychosis and moderately for others. It also provides evidence that using cannabis might cause psychosis and is not just a consequence of it.

Young people using cannabis at baseline in the study were more likely to have psychotic symptoms four years later (odds ratio (OR) 1.67), after adjustment for a slew of confounding factors. The likelihood of having any psychotic symptoms rose with frequency of use in a dose–response way, from 0.99 for use less than once a month to 2.23 for almost daily use. The adjusted difference in risk of psychosis with cannabis use was 23.8% for those susceptible to psychosis and is not just a consequence of it.

The study analysed data on 2437 young people aged 14–24 years who were part of the random regional representative population sample in the prospective early developmental stages of psychopathology study (EDSP) in Munich, Germany.

Whether using cannabis causes psychosis has been disputed, some arguing that predisposition to psychosis may be the driving force to taking up the drug, rather than cannabis causing psychosis to be expressed. This—the first prospective study to research the issue—suggests that cannabis may be the culprit.