SHORT REPORT

Multimodal electrophysiological studies including motor evoked potentials in patients with locked-in syndrome: report of six patients

Claudio Bassetti, Johannes Mathis, Christian W Hess

Abstract

Clinical and electrophysiological findings in six patients with locked-in syndrome are reported. Motor evoked potentials (MEPs) after magnetic stimulation of the motor cortex were absent in four patients, none of whom recovered clinically. In two patients, MEPs could be obtained from the severely paretic limbs and almost full motor recovery followed. Somatosensory evoked potentials were altered in four of the patients, and brainstem auditory evoked potentials were altered in two of four patients examined, showing a clinically unsuspected segmental involvement. The EEG showed a predominance of reactive alpha activity in all patients, documenting a preserved consciousness. It is concluded that a multimodal electrophysiological approach, in addition to clinical assessment, can be helpful in diagnosing locked-in syndrome, estimating the extension of the underlying brainstem dysfunction, and predicting functional outcome.

(J Neurol Neurosurg Psychiatry 1994;57:1403-1406)

The term “locked-in” was coined in 1966 by Plum and Posner1 for a condition first described by Alexandre Dumas in his book The Count of Monte Cristo. The locked-in syndrome is characterised by tetraplegia, paralysis of all cranial motor functions except vertical eye movements, and preserved consciousness that can be assessed by means of communication through a blinking code. Ventral pontine vascular lesions are the most common cause of locked-in syndrome. During the past few years, an increasing number of reports allowed the recognition of different varieties of locked-in syndrome with respect to the clinical picture (classic, incomplete, and total forms), topography of the lesion (mesencephalic and pontine lesions), aetiology (vascular and non-vascular causes), and evolution (transient and chronic forms) of the disorder.1-4

The preserved consciousness can be ascertained by a normally reactive alpha rhythm in the EEG.9 Involvement of segmental brainstem structures in locked-in syndrome is not uncommon and has been documented in several patients by means of somatosensory and evoked potentials (SEPs) and brainstem auditory evoked potentials (BAEPs).10 Conversely, dysfunction of the pyramidal tract has been shown electrophysiologically with the help of motor evoked potentials (MEPs) only once.11

The outcome of locked-in syndrome is generally poor. In a review of 139 cases, Patterson and Grabois7 reported a mortality of 67% in vascular and of 41% in non-vascular cases. Over the past years, however, patients with good recovery have been reported,1-4,8 stimulating a search for prognostic indicators. Because of the lack of knowledge about the diagnostic and prognostic value of a multimodal electrophysiological approach we decided to study six consecutive patients with locked-in syndrome by means of a systematic protocol including clinical assessment, EEG, SEPs, BAEPs, and MEPs.

Methods

Transcranial magnetic stimulation of the motor cortex and transcutaneous high voltage electrical stimulation of the cervical motor roots (for the upper limb muscles) were performed as described elsewhere.13 For magnetic stimulation a specially designed powerful magnetic stimulator with a total capacitance of 800 μF and a total maximal charging energy of 2500 J was employed. A circular stimulating coil—mean diameter 8 cm and total inductance 30 μH—was centred over the vertex for the upper limb muscles and placed 2-4 cm anteriorly for the lower limb muscles with the current flowing anticlockwise as viewed from above (as conventionally defined) to excite the right sided muscles and vice versa. The MEPs were recorded from the abductor digiti minimi, biceps brachii, and tibialis anterior muscles of both sides from surface electrodes placed over the belly and tendon of the target muscles. Stimulus intensity was increased stepwise until reproducible responses were obtained, and three to four compound muscle action potentials (CMAPs) were recorded from each
target muscle. Recordings were made with a four channel EMG device (Medelec sensor) with amplifier bandpass filters from 3 Hz to 3 kHz. In earlier studies on normal subjects it was always possible to obtain unequivocal responses from those target muscles even without facilitation by voluntary contraction. When no response could be evoked until 100% of stimulus output strength, painful stimuli were applied to the respective limb, sufficient to induce some reaction (in the face) just before the cortical stimulus to facilitate the responses. The onset latencies of the CMAPs obtained from cortical and cervical stimulation were measured. The central motor conduction time (CMCT) to the abductor digiti minimi, biceps brachii, and tibialis anterior muscles was defined as the difference between the onset latencies after cortical and motor root stimulation.

The BAEPs were elicited by monaural clicks with an intensity of 75–95 dB above sensory level delivered at 10/s while masking the contralateral ear with white noise. Recording was done with a Medelec sensor device and amplifier bandpass filters from 1 Hz to 1 kHz. The recording needle electrodes were placed at Cz' (indifferent) and A1/A2 (different) according to the 10–20 nomenclature.

The SEPs were obtained by electrical stimulation of the median nerve at the wrist with square wave pulses of 0.2 ms duration and an intensity of 4 mA above motor threshold of the thenar muscle delivered at 10/s while masking the contralateral ear with white noise. Recording was done with a Medelec sensor device and amplifier bandpass filters from 3 Hz to 3 kHz with the recording needle electrodes placed according to the 10–20 nomenclature: (a) ipsilateral Erb's point to Fz; (b) spinous process of C7 to Fz; (c) contralateral handfield (P3/P4) to the ipsilateral ear (A1/A2); (d) P3/P4 to Fz. Two samples each of 512 to 1024 sweeps were averaged on either side.

The EEG recordings were taken with an eight channel machine for at least 30 minutes with both bipolar and unipolar (Wilson) reference. At the same time reaction to pain, passive eye opening, and acoustic stimuli were tested.

The results of the MEPs, BAEPs, and SEPs were compared with the normative data of our laboratory. A result was judged abnormal when it was more than 2.5 SD beyond the mean of our normal data.

### Results

The table summarises the principal clinical, radiological, and electrophysiological features of the six patients studied.

### Clinical Findings

All patients (three men and three women, age ranging from 16 to 71 years) presented with the locked-in syndrome as defined by Plum and Posner. Locked-in syndrome appeared after successful resuscitation after cardiac arrest in patient 4, and after general anaesthesia for osteosynthesis in patient 5. In the other four patients, locked-in syndrome was preceded by symptoms and signs of vertebrobasilar ischaemia. In the classification proposed by Bauer et al., patients 4 and 6 had a partial locked-in syndrome because of partially preserved horizontal eye movements. In the remaining patients the syndrome was complete. In all patients tetraparesis was severe. Patient 2 and 3 died within the first week after admission. Patient 1 and 5 evolved to a chronic locked-in syndrome (follow up of four years and one year respectively). Patients 4 and 6 made a good functional recovery (independent in everyday activities) within four months of onset of locked-in syndrome.

### Neuroradiology

All patients had brain CT, which was considered to be normal in four. In two patients (5 and 6) brainstem infarction was suspected. Brain MRI was carried out in three patients (1, 5, 6), confirming the presence of a bilateral ventral pontine infarction in all three (fig 1). Patient 6 had also had a right ventral mesencephalic infarction. Conventional angiography or MR angiography was performed in three patients and showed a basilar thrombosis in two (1, 5).

### Electrophysiology

An EEG showed normally distributed, reactive alpha activity in all patients. In patient 5, the EEG was reactive to photostimulation but not to pain or acoustic stimuli. Amplitude and latency of early (N20/P25) cortical responses of SEPs were altered bilaterally in four patients. The BAEPs were altered unilaterally or bilaterally in two of four patients.

In the acute stage, MEPs could be recorded in two patients (4, 6) both of whom recovered clinically. Of the four patients with initially absent MEPs, two died and two

---

**Clinical, radiological and electrophysiological findings in six patients with locked-in syndrome**

<table>
<thead>
<tr>
<th>Patient No</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>36</td>
<td>M</td>
<td>F</td>
<td>F</td>
<td>M</td>
<td>F</td>
</tr>
<tr>
<td>Sex</td>
<td>M</td>
<td>M</td>
<td>F</td>
<td>F</td>
<td>M</td>
<td>F</td>
</tr>
<tr>
<td>Aetiology</td>
<td>Brainstem stroke</td>
<td>Brainstem stroke</td>
<td>Brainstem stroke</td>
<td>Brainstem stroke</td>
<td>Cardiac arrest</td>
<td>Brainstem stroke</td>
</tr>
<tr>
<td>EEG</td>
<td>7–10 Hz reactive</td>
<td>9 Hz reactive</td>
<td>ND</td>
<td>10–12 Hz reactive</td>
<td>7–10 Hz reactive</td>
<td>7–8 Hz reactive</td>
</tr>
<tr>
<td>SEP</td>
<td>Altered bilateral</td>
<td>Altered bilateral</td>
<td>Altered bilateral</td>
<td>Normal</td>
<td>Normal</td>
<td>Normal</td>
</tr>
<tr>
<td>BAEP</td>
<td>Absent*</td>
<td>Absent</td>
<td>Absent</td>
<td>Absent</td>
<td>Absent</td>
<td>Absent</td>
</tr>
<tr>
<td>MEP</td>
<td>Normal</td>
<td>Normal</td>
<td>Normal</td>
<td>Normal</td>
<td>Normal</td>
<td>Normal</td>
</tr>
<tr>
<td>CT</td>
<td>Normal</td>
<td>ND</td>
<td>ND</td>
<td>Good recovery</td>
<td>Absent*</td>
<td>Absent</td>
</tr>
<tr>
<td>MRI</td>
<td>Pontine infarct</td>
<td>Death</td>
<td>Death</td>
<td>Pontine infarct</td>
<td>Pontine infarct</td>
<td>Pontine infarct</td>
</tr>
<tr>
<td>Evolution</td>
<td>Chronic locked-in syndrome</td>
<td>Chronic locked-in syndrome</td>
<td>Chronic locked-in syndrome</td>
<td>Chronic locked-in syndrome</td>
<td>Chronic locked-in syndrome</td>
<td>Chronic locked-in syndrome</td>
</tr>
</tbody>
</table>

*Reappeared on follow up; †follow up of four years; ‡follow up of one year. ND = not done.
Electrophysiological studies in patients with locked-in syndrome

Figure 1  Patient 1; Sagittal (A) and axial (B) T1 weighted MRI on day 5 in hospital showing a coil shaped gadolinium enhancing, bilateral infarction at the basis of the pons.

Figure 2  Patient 1; MEPs (A) after transcranial magnetic stimulation of the motor cortex at three days and (B) at six months. In the acute phase no MEPs could be recorded from any of the examined muscles (A; only abductor digit minimi responses are shown). In the chronic phase MEPs were recorded from all muscles examined (B; abductor digit minimi (ADM), biceps brachii (BB), tibialis anterior (TA)), although with a prolonged latency and a low amplitude (ADM right: central motor conduction time (CMCT) 20.4 ms, amplitude 0.9 mV (12% of amplitude obtained by stimulating at the spinal roots); ADM left: CMCT 8.8 ms, amplitude 0.3 mV (4%); BB right: CMCT 12.8 ms, amplitude 2.4 mV (25%); BB left: CMCT 12.8 ms, amplitude 2.3 mV (25%); TA right: corticomuscular latency* 38.6 ms, amplitude 3.3 mV; TA left: corticomuscular latency* 36.8 ms, amplitude 3.3 mV). *Stimulation of the lumbar motor roots was not done. Patient 6; MEPs after transcranial magnetic motor cortex stimulation at two days. Normal responses were recorded from the left ADM (C). Responses from the other target muscles examined (ADM right (C) and both TAs (D) had a prolonged latency and reduced amplitude (ADM right: central motor conduction time (CMCT) 9.9 ms, amplitude 0.4 mV (14% of amplitude obtained by stimulating at the spinal roots); ADM left: CMCT 7.3 ms, amplitude 1.5 mV (26%); TA right: CMCT 21.1 ms, amplitude 0.5 mV (12%); TA left: CMCT 20.5 ms, amplitude 0.2 mV (6%).
evolved to a chronic locked-in syndrome. In the two patients with chronic locked-in syndrome MEPs could only be recorded a few months later in the presence of minimal motor recovery (fig 2).

**Discussion**

Our findings suggest potential diagnostic and prognostic value for a multimodal electrophysiological approach in patients with locked-in syndrome. In four of our six patients MEPs after magnetic stimulation of the motor cortex were abolished in the acute stage. All of them had a pontine stroke with an unfavourable outcome: two of them died within a week, and two remained severely paretic evolving to a chronic locked-in syndrome. In the remaining two patients, who presented with locked-in syndrome after cardiac arrest and a pontomesencephalic stroke, normal or only slightly abnormal MEPs were obtained from the limbs despite near-complete paralysis. Both of them had a favourable outcome with functionally satisfactory recovery of speech and motor functions. Thus the absence of MEPs confirmed the interruption of the pyramidal tract and heralded a poor outcome. Conversely, preservation of MEPs indicated at least partial structural integrity of corticospinal pathways and correctly suggested a potential for recovery.

The finding of preserved MEPs despite poor or absent motor functions is interesting from a neurophysiological point of view. A similar discrepancy between clinical and electrophysiological findings has been reported previously in patients with brainstem stroke as well as in those with traumatic spinal cord lesions. This could be explained on the basis of some spared corticospinal fibres, which are not accessible or too few to become effective at the motoneuron level when activated by voluntary effort as opposed to cortical stimulation. Since the work of Evarts, we know that spontaneous pyramidal activity in the waking monkey is continuous and regular, and increases its frequency during motion. Conversely, single cortical stimuli sufficient to excite limb muscles induce repetitive impulses in the fast pyramidal tract neurons in primates and in humans. It is thus conceivable that in our patients the strong magnetic stimuli induced abnormally long lasting bursts of impulses in the few functioning pyramidal fibres, which were sufficient to bring some motoneurons to firing level. Likewise, recovery of few pyramidal tract fibres from acute ischaemic conduction block could explain the reappearance of MEPs in the subacute stage of locked-in syndrome despite minimal clinical recovery, as also reported by Facco et al. Alternatively, the not uncommon presence of tegmental dysfunction evokes the possibility that clinical paresis may be due not only to an efferent deficit but also to an insufficient cortical motor arousal, which can be overcome by magnetic stimulation.

Four of our six patients were thought to have clinically normal sensory functions, as usually assumed in locked-in syndrome. BAEPs were, however, found to be altered in two of four patients and SEPs were altered in four of six patients examined. Similarly, Towle et al. reported abnormal SEPs in eight of nine patients. These findings accord with some necropsy studies, which documented a frequent and often clinically unsuspected tegmental involvement in locked-in syndrome. Objective evidence of a disruption of afferent pathways in a tetraparetic patient without speech can be helpful for a correct topographic diagnosis, especially when brain CT is normal as in four of our six patients and brain MRI is not available. From a prognostic point of view, SEPs in locked-in syndrome seem less helpful than MEPs. In our experience, as in other reports, normal SEPs can be followed by both a good or a bad functional recovery.

We thank P Huber, MD for performing the radiological studies and K Rößler, MD and E Markus, MD for helpful comments.

Multimodal electrophysiological studies including motor evoked potentials in patients with locked-in syndrome: report of six patients.

C Bassetti, J Mathis and C W Hess

*J Neurol Neurosurg Psychiatry* 1994 57: 1403-1406
doi: 10.1136/jnnp.57.11.1403

Updated information and services can be found at:
http://jnnp.bmj.com/content/57/11/1403

These include:

**Email alerting service**

Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/