SHORT REPORT

Seizures arising from the inferior parietal lobule can show ictal semiology of the second sensory seizure (SII seizure)

J Yamamoto, A Ikeda, M Matsuhashi, T Satow, M Takayama, S Ohara, R Matsumoto, N Mikuni, J Takahashi, S Miyamoto, W Taki, N Hashimoto, H Shibasaki

A 52 year old right handed man presented with medically intractable partial seizures consisting of numbness on the left upper back spreading to the left upper as well as lower limbs. Head computed tomography and magnetic resonance imaging showed a round calcified lesion in the depth of the superior ramus of the right sylvian fissure. Ictal electrocorticographic recording with chronically implanted subdural electrodes showed low voltage fast activities starting exclusively from an electrode located on the right inferior parietal lobule. No apparent ictal activities were observed from the depth electrodes inserted in the parietal operculum. Somatosensory evoked potentials of 75 ms to 145 ms latency were recorded from the ictal onset zone, which was 2 cm caudal to the perisylvian area corresponding to the second somatosensory area. Seizures arising from the inferior parietal lobule including the angular and supramarginal gyri can produce partial seizures whose ictal semiology and scalp electroencephalography are indistinguishable from the ones originating from the second somatosensory area.

The ictal symptoms of the second somatosensory seizure (SII seizure) (seizures involving the second somatosensory area) by Penfield and Jasper are characterised by bilateral and/or more widespread, axial numbness, tingling or pain whose localisation in the body is different from the somatotopy of the primary sensory cortex, and several similar cases have been reported. In humans, SII was rather indistinctly defined to be located in the parietal upper bank of the sylvian fissure, mainly based on the results of direct electric cortical stimulation. This notion has been supported by somatosensory evoked potential (SEP) findings, magnetoencephalographic studies, and more recently by neuroimaging studies. However, probably because of interindividual variability of the anatomical location of SII, and also because of a limited number of cases studied by intracranial recording, it is still unclear which part of the parietal cortex is responsible for the so-called SII seizure. We evaluated a patient in whom the initial sensory symptoms were considered to be consistent with SII seizures and the ictal onset zone was finally defined in the parietotemporal perisylvian areas, clearly caudal to SII. Neurophysiological data of this patient were described elsewhere for an entirely different purpose (M Matsuhashi, et al, 8th international conference on functional mapping of the human brain, 2002. Abstract available on CD ROM).

CASE REPORT

A 52 year old right handed man was investigated because of medically intractable partial seizures. He began to have intractable partial seizures at the age of 20 years. The seizures always consisted of numbness at the left upper back that spread to the left upper extremity and the left hip, and then to the left lower extremity. This simple partial seizure (SPS) was sometimes followed by loss of consciousness and automatism involving the hands and feet. With various combinations of antiepileptic drugs including phenytoin, carbamazepine, and valproic acid, his complex partial seizures (CPSs) occurred with an average frequency of one to three times per month, the longest seizure free interval being one month and the maximum seizure frequency of one to two times a day.

On physical examination, he had no focal neurological abnormality. Brain computed tomography (CT) showed a round calcified lesion in the depth of the right sylvian fissure. FLAIR-MRI of the brain showed high intensity signal abnormality around the calcified lesion (fig 1A). Interictal FDG-PET showed a small area of hypometabolism around the calcified lesion.

The patient was studied by continuous video EEG monitoring for 13 days. Interictal spikes were infrequently observed at T6 and O2 (International 10–20 system). Although ictal EEGs recorded from the scalp electrodes did not clearly localise the seizure onset zone precisely, the ictal symptoms and the calcified lesion in the depth of the right sylvian fissure strongly suggested that the habitual seizures in this patient originated from the right parietal perisylvian area, especially from SII.

To further localise the epileptogenic and symptomatic areas before surgical treatment, the patient underwent implantation of two subdural grid electrodes (2×5), one each on the right frontal and temporal lobes (fig 1B), and one depth electrode in the right parietal operculum around the calcified lesion. Informed consent for this investigation was obtained from the patient (clinical protocol number 79 approved by the ethical committee of Kyoto University Graduate School of Medicine). The electrodes were 3 mm in diameter and placed with 1 cm interelectrode distance (centre to centre). The depth electrode had six contacts, each 2.3 mm of length and 1 cm of centre to centre distance (AD-TECH, Racine, WI, USA).

During invasive monitoring, the patient experienced more than one hundred SPSs and five CPSs. In all the recorded habitual seizures, low voltage fast activities of about 20 Hz started exclusively from electrode 14 (fig 1C). In almost all the seizures, ictal activities started almost simultaneously with the clinical symptoms. Once ictal electrocorticographic activities propagated to the adjacent electrodes, the patient developed CPSs. No ictal activities were observed in the depth

Abbreviations: SEP, somatosensory evoked potential; CT, computed tomography; MRI, magnetic resonance imaging; SII, second somatosensory area.
electrodes in the recorded four seizures, which included two habitual SPSs and two CPSs.

For functional mapping, high frequency electric cortical stimulation through subdural electrodes was used as previously described (SEN-7203 and ss-102J, Nihon-Koden, Japan). Electric cortical stimulation between electrodes 11 and 16 produced contraction of the right oral angle. None of the other electrode pairs produced clinical symptoms even when stimulated with the maximum intensity of 15 mA (fig 1B).

For cortical SEP recording, median and tibial nerves were stimulated at the wrists and ankles, respectively, on each side separately with an electric pulse of 0.3 ms duration, and the intensity was adjusted just above the motor threshold (Pathfinder II, Nicolet Biomedical, WI, USA). All electrodes were referred to a scalp electrode placed on the skin over the mastoid process contralateral to the side of implantation. The interstimulus interval and the band pass were set to 3.3 s and 0.5–100 Hz, respectively. Cortical SEPs were identified from electrodes 12 and 14 with different latencies and somatotopic representations from each other (table 1). At electrode 14, cortical responses of 80–100 ms of latency were recognised after both the median and tibial nerve stimulation of either side, whereas at electrode 12, cortical responses of 140 ms were recorded only to the left and right median nerve stimulation.

Precise location of the electrode grids relative to the sylvian fissure as well as the central sulcus was confirmed by the observation during surgery and by 3D reconstructed MRI taken after implantation of the electrodes. Based on these findings, electrode 14 was situated just below the superior ramus of the sylvian fissure in the inferior parietal lobule where the supramarginal and angular gyri were located. Electrode 12 was located on the postcentral gyrus.

The patient underwent lesionectomy and resection of the cortical area covered by electrodes 13, 14, and 15. Histologically the calcified lesion was a psammomatous meningioma and its surrounding tissue was cortical dysplasia. The patient had been free of seizure for one year after the resection, and afterwards he gradually started having simple partial seizure consisting of numbness at the left upper back again, but with lesser frequency as compared with the presurgical state.

**DISCUSSION**

In this patient, the ictal symptoms consisting of numbness on the left upper back rapidly spreading to the left hand and foot and the existence of a calcified lesion in the depth of the right sylvian fissure (fig 1A) strongly suggested the right SII as the epileptogenic or symptomatogenic zone.

We recorded more than 100 habitual seizures during the invasive monitoring. In most of those seizures, ictal activities started almost simultaneously with the ictal symptoms. Therefore, the area underlying electrode 14 was considered to be both the ictal onset and symptomatogenic zone, and thus electrode 14 was thought to be at or close to SII. However, electrode 14 was located at the area just below the superior ramus of the sylvian fissure that corresponded to the inferior parietal lobule. The arrow indicated the onset of the clinical symptoms.

**Table 1** Latency and amplitude of cortical SEP at electrode 12 and 14 after median and tibial nerve stimulation of each side

<table>
<thead>
<tr>
<th>Electrode 14</th>
<th>Electrode 12</th>
</tr>
</thead>
<tbody>
<tr>
<td>Latency (ms)</td>
<td>Amplitude (mV)</td>
</tr>
<tr>
<td>Left median nerve</td>
<td>80</td>
</tr>
<tr>
<td>Right median nerve</td>
<td>80</td>
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<tr>
<td>Left tibial nerve</td>
<td>100</td>
</tr>
<tr>
<td>Right tibial nerve</td>
<td>100</td>
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Seizures and the inferior parietal lobule

Although human SII has been defined to be located in the parietal upper bank of the sylvian fissure by direct electric cortical stimulation, SEPs are poorly defined. Moreover, only a few cases of SII seizures were studied by intracranial recordings, and none of them described anatomical location of the ictal onset zone in detail. Therefore, the responsible area for the so-called SII seizure may actually include not only SII itself but also the adjacent cortical area, such as the angular and supramarginal gyrus.

From electrodes 12 and 14, cortical SEPs of different latencies and somatotopic representations were recorded. Long latency (80–140 ms) and bilateral representation (Table 1) are characteristic features of SEPs arising from SII. At electrode 12, however, only median nerve electric stimulation produced clear, long latency SEPs, while at electrode 14 prominent SEPs were recorded after both median and tibial nerve stimulation. It may suggest that SEPs at electrode 12 reflect the somatotopic organization of SII as described in the previous reports, which showed that the hand area was most laterally and the foot area was most medially located along the upper bank of the sylvian fissure and the parietal operculum. Thus based on the present findings of cortical SEPs and anatomical data, electrode 12 is considered to be at or very close to SII.

Electrode 14 is outside the conventionally accepted SII region. However, the generation of long latency SEPs in this area after bilateral median and tibial nerve stimulation indicate that this particular area also plays a certain part in the sensory processing different from that at electrode 12 (M Matsuhashi, et al, 8th international conference on functional mapping of the human brain, 2002). According to the primate studies, area 7b, the retinotopic and posterior auditory areas, and the granular portion of the insula may account for the bilateral and/or diffusely contralateral sensory symptoms. Therefore, the area covered by electrode 14 could be responsible for the generation of partial seizures whose ictal semiology is indistinguishable from that of SII origin.

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