Human second somatosensory area: subdural and magnetoencephalographic recording of somatosensory evoked responses

Tatsuya Mima, Akio Ikeda, Takashi Nagamine, Shogo Yazawa, Takeharu Kunieda, Nobuhiro Mikuni, Wataru Taki, Jun Kimura, Hiroshi Shibasaki

Abstract

Objective—To investigate somesthetic functions of the perisylvian cortex. Methods—Somatosensory evoked magnetic fields (SEFs) and somatosensory evoked potentials (SEPs) of the perisylvian cortex were recorded directly from subdural electrodes in a patient with a left frontal brain tumour. Results—The most prominent SEP components after electrical stimulation of the right and left hands and the right foot were double peaked negativity recorded just above the sylvian fissure (latency 80 to 150 ms), respectively (N1a and N1b). Generator sources for the magnetoencephalographic counterparts of those peaks (N1a(m) and N1b(m)) were both localised at the upper bank of the sylvian fissure, and those of N1a(m) were more anteromedially located than those of N1b(m). Conclusions—These findings suggest the existence of at least two separate somatosensory areas within the human perisylvian cortex.

Keywords: somatosensory evoked potentials; subdural recording; magnetoencephalography; human second somatosensory area

There have been only a few studies in humans in which the somatosensory cortex in the opercular cortex, possibly corresponding to the second somatosensory area (SII) of primates, has been investigated by applying direct cortical stimulation or recording somatosensory evoked potentials (SEPs) from the cortical surface. Recording of somatosensory evoked magnetic fields (SEFs) has been found to be a useful tool for studying SII non-invasively. Until recently, however, little was known about human SII compared with the primary somatosensory area (SI). Moreover, some primate species have been reported to have multiple sensory representations within the opercular cortex. Therefore, the problem remains unsolved as to whether the human so-called SII as previously determined by the classic invasive methods and magnetoencephalography (MEG) in humans is a single and uniform area or not.

To investigate the human somatosensory areas in the perisylvian cortex, we compared SEFs and SEPs directly recorded from the cortical surface in a patient who needed chronically implanted subdural electrodes for the surgical treatment of a left frontal brain tumour, and estimated their generator sources.

Methods

PATIENT

A 29 year old right handed man was evaluated before surgical resection of a left frontal brain tumour which was discovered by CT. Clinically, he had no neurological symptoms. A Wada test disclosed that the left hemisphere was dominant for language and memory function. Functions of the cortical areas surrounding the tumour were studied by using two chronically implanted 4×5 subdural electrode grids (A, B) (fig 1A). Each subdural electrode grid (AD-Tech Co) consisted of platinum electrodes of 3 mm diameter, and the centre-to-centre interelectrode distance was 1 cm. Informed consent was obtained from the patient according to the approval criteria of the ethics committee of Kyoto University School of Medicine.

Electrical cortical stimulation with the subdural electrodes according to a previously described method showed the cortical functional map in that area (fig 1B). Repetitive electric stimuli were given to each electrode for five seconds in a separate session. The upper limit of the stimulus strength was set to the intensity just below the afterdischarge threshold, or 15 mA if no afterdischarges occurred. Stimulation of electrodes 9 and 15 elicited a negative motor response, or the inability to perform voluntary motor contraction. However, none of the electrodes produced sensory symptoms when stimulated. Actual location of the electrode grids relative to the sylvian fissure was visually determined during surgery (fig 1B).

The brain tumour was located just below the posterior central part of the plate A with a diameter of 3 cm, and was pathologically diagnosed as grade 2 astrocytoma. Brain oedema around the tumour was minimal and did not involve the brain tissue below plate B.

RECORDING METHODS

Stimulus for SEFs and cortical SEPs

Median and tibial nerves were stimulated at the wrist and the ankle, on both sides separately with an electric pulse of 0.2 ms duration and the intensity adjusted just above the motor threshold for each corresponding muscle. The interstimulus interval (ISI) was constant at 2.9 s.
A shorter ISI (0.9 s) was also used for the recording of SEFs and the right median nerve stimulation for the subdural recording of SEPs.

Subdural recording of SEPs
SEPs were recorded from 20 subdural electrodes of plate B (Pathfinder II MEGA, Nicolet). All electrodes were referred to one of the subdural electrodes on the plate A on the frontal lobe. The passband for the EEG recording was set to 0.5-500 Hz. In each session, two blocks of 100 responses each were separately averaged. The analysis time window consisted of 25 ms before and 225 ms after stimulus onset (sampling point 256). The amplitude of each peak was measured in the group averaged waveforms from the baseline, which was determined by averaging the pre-stimulus segment for each channel.

SEFs
SEFs were recorded two weeks before implanting the electrode grids. With the subject sitting in a magnetically shielded room (NKK Co), SEFs were recorded with a 122 channel whole head first order planar SQUID gradiometer (Neuromag122, Neuromag Ltd). A vertical bipolar electro-oculogram (EOG) was simultaneously recorded by a pair of cup electrodes. The signals were filtered (0.03 to 300 Hz for MEG, and 0.07 to 320 Hz for EOG) and digitised at 0.9 kHz. Samples containing the EOG exceeding 150 µV or the MEG exceeding 3000 fT/cm were automatically excluded from the average. Two blocks of 100 responses each were averaged separately for each session. The analysis time window was 600 ms including the prestimulus baseline of 100 ms. The averaged signals were low pass filtered at 200 Hz.

The coordinate system was defined using an Isotrack digitiser (Polhemus Navigation Sciences). The x axis ran through the two preauricular points from the left to the right, the y axis passed through the nasion perpendicular to the x axis (anterior direction positive), and the z axis ran from the skull base to the top of the head. The exact location of the head within the sensors was determined by three head position indicators placed on the head.

The channel with the local extreme signal was selected for the measurement of the peak latency. To identify the cortical generator sources, the equivalent current dipole was determined by applying a least squares search, using a subset of channels (at least 20 channels for each) over the response area. The head was modelled by a conductor sphere whose centre was defined by fitting it to the curvature of the brain surface of this patient. Equivalent current dipoles explaining more than 80% of the field variance were selected for analysis.

Results
In the subdural recording, SEPs following the right and left median, and the right tibial nerve stimulation showed the maximal response at B8 electrode which was just superior to the sylvian fissure and anterior to the face motor area (fig 2). Initial cortical response to the right median and tibial nerve stimulation was a positive potential (latency 44 and 62 ms, respectively; labelled as P1 (asterisk) in fig 2, table). For the right and left median, and the right tibial nerve stimulation, double peaked negative potentials (N1a and N1b) were recognised at the same electrode (right median; 84 and 108 ms, left median; 91 and 124 ms, right
tibial; 104 and 136 ms). In all the records in which the responses were identified, N1a was more anterosuperiorly distributed (B8 and B9) than N1b (B8 and B13). For the hand stimulation, small positive potentials were also recognised in the superior electrodes: at B5 (108 ms) for the right, and at B15 (116 ms) for the left median nerve stimulation.

The planar gradiometer detects the local extreme signals just over the generator source. The generator source of the initial cortical response (N20m) was localised at the post-central cortex (hand SI). In addition to the SI source, responses to median nerve stimulation with longer ISI (2.9 s) were recognised at the bilateral temporal areas at the latency of around 100 ms. No clear responses were found at these areas for the shorter ISI (0.9 s) or after stimulation of the tibial nerve. For right and left median nerve stimulation, local cortical activity over the left perisylvian cortex was identified (latency 108 and 122 ms respectively), which was preceded by a smaller peak or notch at the nearby site (86 and 89 ms respectively) (fig

**Figure 2.** SEPs recorded subdurally from the B plate after electric stimulation of (A) the right and (B) the left median nerve at the wrist, and (C) the right and (D) the left tibial nerve at the ankle. Two sets of SEPs with ISI of 2.9 s are superimposed, SEPs with the shorter ISI (0.9 s) are also shown by interrupted lines in (A). Vertical interrupted lines show the stimulus onset. Each recognisable peak is indicated in the figure. RMN=right median nerve; LMN=left median nerve; RTN=right tibial nerve; LTN=left tibial nerve.

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Both MEG and EEG amplitudes were measured from the baseline determined by the prestimulus segment. The three dimensional locations of ECDs were measured from the origin of the head coordinate.
No earlier responses were recognised at these areas. The location of the equivalent current dipoles for all of these responses was at the superior bank of the sylvian fissure (fig 3B). Judging from the latency and the direction of the electrical dipole (pointing anterosuperiorly), these two responses are most likely the magnetic counterparts of subdurally recorded N1a and N1b. Therefore they were referred to as N1a(m) and N1b(m), respectively (table). Equivalent current dipole of the ipsilateral N1a(m) was located superior to that of the contralateral N1a(m), and that of the contralateral N1b(m) was anterior to that of the ipsilateral N1b(m). Irrespective of the side of stimulation, the generator sources of N1a(m) were localised anteromedial to those of N1b(m).

**Discussion**

In the present study, somatosensory symptoms were not elicited by electrical stimulation of either the electrode B8 or B9. Therefore, it is possible that SII was buried in the sylvian...
fissure. However, cortical SEPs after stimulation of each hand and right foot were recorded from the left perisylvian cortex. This is the first demonstration of SEPs to foot stimulation recorded directly from the human perisylvian cortex. Primate experiments and human MEG studies have shown somatotopy in the perisylvian cortex.⁹¹⁰ However, no somatotopy was seen in the present study, as all the cortical potentials showed maximal amplitude at B8, although the response to the foot stimulation was not detected by MEG. Anatomically, the somatosensory areas at the perisylvian cortex are rather small compared with the SI.⁴ Therefore, it is plausible that the somatotopy, if present, would be too small to be detected by the electrode grids with the interelectrode distance of 1 cm.

The present study is the first to compare subdurally recorded SEPs and SEFs in the same subject. The initial cortical SEP component was P1 to contralateral hand or foot stimulation. However, the MEG counterpart of P1 was not recognised. One probable explanation is that the direction of the generator source of P1 was not suitable for MEG recording (for example, radially oriented to the scalp). It could be also the case for the N1a and N1b after right foot stimulation of which the MEG counterpart was not clearly recorded.

In previous MEG studies, only a large component with long latency (100 to 120 ms, corresponding to N1b(m)) in the present study was analysed as the so-called SII responses. However, the reports of invasively recorded SEPs suggested the existence of two successive but distinguishable components around 100 ms (N1a and N1b in the present study). For the hand stimulation, those two components were distinctly recognised also by the present MEG recording (N1a(m) and N1b(m)). The MEG channel which showed the most negative peak for N1a(m) was different from that for N1b(m) (fig 3A). As our MEG system consists of planar gradiometers, the fact that these two peaks are separable in both space and time suggests that at least two generator sources are needed to explain these cortical activities. The equivalent current dipoles of the first were anteromedially located compared with those of the second. The distance between the source of N1a(m) and that of N1b(m) was more than 10 mm. Thus the hypothesis of multiple generator sources would be more appropriate than the model assuming the propagation of the activation within the same area. In cortical SEP waveforms, N1a was distributed more anteromedially than N1b. Therefore, in terms of the findings obtained from both SEFs and cortical SEPs, two different hand representation areas were demonstrated within the perisylvian cortex. Some primates are known to have multiple sensory representation within the operculum.¹¹ In the present study, no differences in foot somatosensory areas were detected demonstrated by MEG source localisation for the first time in a human, and confirmed by cortical SEPs. As has been documented in a previous MEG study, responses at the perisylvian cortex are extremely sensitive to the stimulus presentation rate.⁴ In the present study, SEPs to the right median nerve stimulation was almost lost when the interstimulus interval of 0.9 s was applied instead of 2.9 s (fig 2A).

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