Somatosensory processing of the tongue in humans

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INTRODUCTION
The tongue is essential for daily life. The tongue is an epithelial sac filled with muscles and connective tissue; these muscles can be controlled willfully and are generally referred to as skeletal muscles or voluntary striated muscles, which are divided into intrinsic and extrinsic muscles (Brand and Isselhard, 2003). In addition, the tongue has various functions: preservation of the position of the teeth and expression of feelings, speech, swallowing, and mastication. However, there has been relatively few neuroimaging and neurophysiological studies focusing on the functions of the tongue. There are several problems with studying somatosensory (tactile) processing of the tongue in humans. The first problem is the choice of a stimulator that can stimulate the tongue while the subject is under a scanning coil or electrodes without causing noise or technical problems. The second problem is that it is extremely difficult to fix the stimulator on the tongue stably, since the tongue itself is made of soft tissue and is convex in shape. The third problem is that tactile stimulation in this area frequently triggers the vomiting reflex. Taking these problems into consideration, ingenuity is required to record the brain activity associated with somatosensory processing of the tongue, and some researchers have developed devices, which are introduced in later sections, to solve these problems.

Recently, several non-invasive recording methods have been used to measure human brain activity. Among these are methods based on neurophysiology, including electroencephalography (EEG), magnetoencephalography (MEG), and methods based on neuroimaging, including functional magnetic resonance imaging (fMRI) and positron emission tomography (PET). EEG is indispensable for examining the neural activities in the human brain and offers a high temporal resolution on the order of milliseconds. EEG technology captures fluctuations in the electrical voltage of the brain through electrodes placed on the scalp in accordance with the standardized guidelines of the International 10–20 system (Jasper, 1958). EEG data also represent changes in the potential differences between different points on the human scalp and the electric field potentials that arise from excitatory and inhibitory postsynaptic potentials. MEG offers several theoretical advantages over EEG in localizing cortical sources (brain dipoles) because the magnetic fields recorded on the scalp are less affected by volume currents and anatomical homogeneity. MEG also permits the spatial and temporal localization of excited cortical areas on the order of millimeters and milliseconds (reviewed in Hari et al., 2000; Kakigi et al., 2000). fMRI, which measures the blood oxygenation level-dependent (BOLD) signal, has been used not only as a tool for mapping brain activity but also as a means of studying the dynamics of neural networks by tracking fMRI response characteristics across various spatial and temporal scales (Logothetis et al., 2001). PET has been used to measure regional cerebral blood flow (rCBF) with the intravenous radioactively labeled water (H215O) bolus technique (Decety et al., 1994). These recording methods have been used to clarify somatosensory processing of the tongue in humans.

In this review article, we examine recent non-invasive research showing the brain activity for somatosensory processing related to the tongue. To our knowledge, no systematic review of the scientific literature on somatosensory processing of the tongue is available despite much research. We argue that combining and reviewing non-invasive research will provide new insights because neurophysiological and neuroimaging methods have limitations when it comes to brain activity. We focus on three broad areas: primary somatosensory cortex (SI), secondary somatosensory cortex (SII), and neuroimaging data.

RESPONSES FROM PRIMARY SOMATOSENSORY CORTEX
Neurophysiological studies using EEG and MEG have helped to clarify the temporal dynamics of the tongue SI (Ishikawa et al., 1980; Altenmüller et al., 1990; Karhu et al., 1991; Nakamura et al., 1998; Maloney et al., 2000; Disbrow et al., 2003; Nakahara et al., 2004). The
evaluation of time-locked EEG and MEG following somatosensory stimulation (i.e., somatosensory-evoked potentials, SEPs; somatosensory-evoked magnetic fields, SEFs) constitutes one of the most useful methods for investigating the human somatosensory system. These studies showed SI activity in the postcentral gyrus, which is consistent with the homunculus reported by Penfield and Boldrey (1937).

One major problem with SEPs and SEFs is that the peak latency of the primary response has not been consistent. For instance, there is general agreement that SI responds after about 20 ms to electrical stimulation of the median nerve or finger (Hari et al., 1993; Mauguire et al., 1997; Kakigi et al., 2000). However, the response time to stimulation of the tongue ranges widely from 10 to 35 ms (Table 1). In addition, some studies reported activity in SI contralateral to the stimulation (Nakamura et al., 1998; Maloney et al., 2000; Nakahara et al., 2004), whereas others found activity in both SIs (Ishiko et al., 1980; Altenmüller et al., 1990; Karhu et al., 1991; Disbrow et al., 2003; Maezawa et al., 2008; Sakamoto et al., 2008a; Tamura et al., 2008).

As mentioned in the Introduction, there might be several reasons why consistent results have not been recorded for tongue somatosensory stimulation. Thus, researchers have made new devices to measure stable brain responses in SI.

Ishiko et al. (1980), who were the first to investigate SEPs elicited by stimulating the tongue, applied mechanical stimulation to the right anterior region of the tongue as it protruded slightly from the mouth. The striking surface of the probe was square and flat with an area of 1 mm². Its striking strength, in terms of tension developed while tapping the tongue, was found to be 10 g.

Altenmüller et al. (1990) used a modified EEG ear clip electrode (5 mm in diameter) located on either side of the tip of the tongue with the cathode on the upper and the anode on the lower side and vice versa. To avoid synchronous electrical stimulation of the oral mucosa and the lips, the outer side of the electrode was electrically isolated. During stimulation, the tongue was held relaxed inside a slightly opened mouth.

Karhu et al. (1991) used a hand-made clip electrode consisting of two Ag plates (diameter = 5 mm), and delivered current to the anterior left side of the tongue. The outside of the electrode was insulated to avoid synchronous stimulation of the oral mucosa. The tongue was held relaxed inside the mouth which was slightly opened to allow leads to come out between the upper and lower teeth.

Nakahara et al. (1998) used an air-puff-derived tactile stimulator, which provides a light, superficial pressure stimulus to the skin. The area of contact by the circular rubber bladder was 1 cm in diameter, and the intensity of the mechanical stimulation was 40 g/cm². The rise time was 20 ms as measured at 10–90% of the intensity.

Maloney et al. (2000) used pairs of thin, stainless steel disk electrodes on modified mandibular or maxillary acrylic splints, similar to orthodontic retainers. The Mandibular splint electrodes were oriented to make contact with the under surface of the tongue along the course of the right and left lingual nerves and the maxillary splint electrodes were oriented to make contact with the hard palate bilaterally along the course of the palatine nerves.

In the MEG study of Disbrow et al. (2003), stimuli consisting of pneumatically driven mechanical taps were applied to the tongue with a balloon diaphragm 1 cm in diameter. The diaphragm was placed as far from the midline as possible, near the edge of the tongue.

Nakahara et al. (2004) used a clip electrode with a 5-mm interelectrode distance, which was attached to the tongue mucosa 5–10 mm from the lingual edge.

Maezawa et al. (2008) used a pair of pin electrodes. The interelectrode distance was 3 mm. Three points on the dorsum of the tongue were stimulated in the following order: (1) the right side (2 cm from the tip of the tongue, 1 cm from the edge), (2) the left side (symmetric to the right side), and (3) the midline (1 cm from the tip of the tongue). Before recordings were made, the three points were marked with crystal violet. A pair of epoxy resin-coated platinum pin electrodes (0.4 mm in diameter) was used for stimulation.

In the MEG study of Tamura et al. (2008), the device used for stimulation was modified from Braille cells for the visually impaired. It consisted of a piezoelectric element and stimulus pins pushed out (0.7 mm/0.4 ms) by application of a direct current to the piezoelectric element to produce tactile stimulation of the area targeted. Eight stimulus pins were aligned, with a gap of 2.4 mm between each. The stimulation was 0.18 N in force.

More recently, we fabricated an intraoral device for each individual using hydrophilic vinyl silicone impression material (EXAFAST/Putty Type, GC, Japan), and recorded SEFs (Sakamoto et al., 2008a) (Figure 1A). The subject bit bilaterally into the EXAFAST, which was mixed uniformly and formed into two blocks. The jaws of the subjects were positioned based on centric occlusion and opened about 8 mm between the upper and lower teeth to make a small space that was important to build the electrode for stimulating the tongue. These blocks were used to create a space from the right to left central incisor teeth to allow relaxation of the tongue, and to keep the jaw in a mandibular rest position. Then, we made four holes, which passed from the buccal to lingual side of the device, positioned on the lingual cusp of the canine teeth and the distolingual cusp of the second molar teeth of the mandible bilaterally.

Next, we made a concentric bipolar electrode, which could be set consisting of stainless steel (3.5 mm in diameter and a gap between the anode and cathode of about 0.9 mm). To extend the line, the

### Table 1 | The peak latency of the primary response after tongue stimulation in previous studies (EEG and MEG).

| Reference            | Recording | Latency of primary response (ms) | Recorded site |
|----------------------|-----------|---------------------------------|---------------|
| Ishiko et al. (1980) | EEG       | 13                              | Contr, Ipsil  |
| Altenmüller et al.  (1990) | EEG   | 21                              | Contr, Ipsil  |
| Karhu et al. (1991)  | MEG       | 55                              | Contr, Ipsil  |
| Nakamura et al. (1998) | MEG | 36                              | Contr         |
| Maloney et al. (2000) | EEG   | 13                              | Contr         |
| Disbrow et al. (2003) | MEG | 10                              | Contr, Ipsil  |
| Nakahara et al. (2004) | MEG | 55                              | Contr         |
| Tamura et al. (2008) | MEG       | 14                              | Contr, Ipsil  |
| Maezawa et al. (2008) | MEG       | 23                              | Contr, Ipsil  |
| Sakamoto et al. (2008) | MEG | 19                              | Contr, Ipsil  |

Contr, contralateral hemisphere to the stimulation; Ipsil, ipsilateral hemisphere to the stimulation.
Current dipole (ECD), which was estimated in MEG studies, is also listed in Table 1, and some studies showed the inferior-posterior direction (Karhu et al., 1991; Nakamura et al., 1998; Nakahara et al., 2004; Maezawa et al., 2008). All these studies indicated the primary response at more than 20 ms. On the other hand, Tamura et al. (2008) and Sakamoto et al. (2008a) showed the primary response within 20 ms, and the superior-anterior or inferior-anterior directions (Sakamoto et al., 2008a; Tamura et al., 2008). After stimulating median nerve, the primary response was recorded at about 20 ms, and the ECD demonstrated anterior direction (Karhu et al., 1991; Wasaka et al., 2003; Huttunen et al., 2006). Several previous studies also have provided evidence that the primary response obtained within 20 ms indicated anterior direction after stimulating face and oral regions, such as buccal (Tamura et al., 2008), lip (Tamura et al., 2008), and hard palate (Bessho et al., 2007). Therefore, we inferred that the true primary response, which was recorded within 20 ms after stimulation of the tongue, should show the anterior direction in ECD, and the second and sequential responses show the posterior direction.

As a characteristic of somatosensory processing of the tongue, neural activation was found in contralateral and ipsilateral hemispheres to the stimulation. Seven in 10 previous studies reported the bilateral activities (Table 1), and some studies showed the inferior-posterior direction (Karhu et al., 1991; Nakamura et al., 1998; Nakahara et al., 2004; Maezawa et al., 2008). All these studies indicated the primary response at more than 20 ms. On the other hand, Tamura et al. (2008) and Sakamoto et al. (2008a) showed the primary response within 20 ms, and the superior-anterior or inferior-anterior directions (Sakamoto et al., 2008a; Tamura et al., 2008). After stimulating median nerve, the primary response was recorded at about 20 ms, and the ECD demonstrated anterior direction (Karhu et al., 1991; Wasaka et al., 2003; Huttunen et al., 2006). Several previous studies also have provided evidence that the primary response obtained within 20 ms indicated anterior direction after stimulating face and oral regions, such as buccal (Tamura et al., 2008), lip (Tamura et al., 2008), and hard palate (Bessho et al., 2007). Therefore, we inferred that the true primary response, which was recorded within 20 ms after stimulation of the tongue, should show the anterior direction in ECD, and the second and sequential responses show the posterior direction.

As a characteristic of somatosensory processing of the tongue, neural activation was found in contralateral and ipsilateral hemispheres to the stimulation. Seven in 10 previous studies reported the bilateral activities (Table 1). For example, MEG studies showed that the difference of the latency between the contralateral and ipsilateral hemisphere was about 1 ms for P40m in Maezawa et al.
In addition to the tongue SII, the neural activities and the somatopic representation of the tongue SII should be clarified. Compared with SI, SII has been speculated to serve a higher level of cognitive function in somatosensory processing, such as attention, decision-making, object recognition, and the integration of nociceptive and non-nociceptive inputs (Mima et al., 1998; Steinmetz et al., 2000; Romo et al., 2002; Torquati et al., 2003; Inui et al., 2004; Nakata et al., 2004; Wasaka et al., 2005). The notion that SII is higher than SI in hierarchy was proposed on the basis of their anatomical relationships: SI sends projections to SII, while SII projects back to the superficial layers of SI (see a review, Iwamura, 1998). Studies in monkeys also showed that a unilateral lesion of SII impaired tasks of tactile learning and retention (Ridley and Ettlinger, 1976, 1978; Garcha and Ettlinger, 1978), and patients with lesions of SII had tactile agnosia (Caselli, 1993). Thus, investigating the neuronal activities of the tongue SII is important to understand the characteristics of somatosensory processing of the tongue, based on several aspects such as source location, source orientation, functional features and comparison with other somatic SII responses (Schnitzler et al., 1999).

To our knowledge, two studies have examined the neural activities of the tongue SII using MEG. In one previous study, Disbrow et al. (2003) showed the responses of the tongue SII after stimulating the edge of the tongue. They also recorded the lip SII, but did not compare the latency, location, and strength of brain responses directly and statistically between them.

In our study, we used individual intraoral devices and a concentric bipolar electrode, and recorded SEFs after stimulating four body sites, the left antero (LA) and postero (LP) lateral margins of the tongue, left median nerve at the wrist (hand), and left tibial nerve at the ankle (foot) (Sakamoto et al., 2008b). Neural activities were recorded from bilateral SII in both hemispheres after the four sites were stimulated. The activity of the tongue SII was recorded 80–110 ms after the stimulation (Figure 4). The tongue SII for LA and LP was located close to the hand SII and significantly more anterior than the foot SII, and there was a significant difference in the location of dipoles between the LA and LP areas of the tongue SII (Figure 5). The tongue SII was located very close to the hand SII, and significantly apart from the foot SII, showing a more anterior location than the foot SII (Figure 6). These findings concerning the locations of ECDs for SII imply the existence of an antero-posterior and infero-superior arrangement of the body surface in SII, which was consistent with some previous studies using MEG (Maeda et al., 1999; Nguyen et al., 2005), fMRI (Del Gratta et al., 2000, 2002; Disbrow et al., 2000; Ruben et al., 2001), and monkeys (Burton and Carlson, 1986; Krubitzer et al., 1986; Cusick et al., 1989). A rough somatotopic representation exists within SII regions, that is, the face, upper limb, and lower limb are located from the lateral to medial region in this order. The mean peak latencies of the tongue
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...bending pressure, and compared the cortical activation with toes and fingertips. They showed the cortical activation, which was located on the contralateral postcentral gyrus, and organized mediolaterally in the order of toes, fingertips, and tongue tip.

Pardo et al. (1997) stimulated the right or left side of the protruding tongue at a rate of 1 Hz with a wooden stick, and rCBF was measured using PET. Stimulation of the right side of the tongue produced a contralateral response in SI, while left side stimulation activated bilateral SI.

FIGURE 4 | MEG signals for the left anterior (LA) stimulation of the tongue in a representative subject. The upper figure shows the SEF waveforms over 204 planar coils from the top of the head. Lower-left figures indicate enlarged waveforms recorded in three areas, A, B, and C. Lower-right figures show each magnetic field pattern at 43.8, 106.7, and 115.5 ms, respectively. The patterns are shown on the sensor array viewed from right (SI and cSII) and left (iSII). The arrows indicate the orientation of the ECD. Comparison between A (SI) and B (cSII) revealed clearly different field distributions and that the ECD of A (SI) was directed posteriorly, while the ECDs of B (cSII) were directed superiorly. L, left; R, right; A, anterior; P, posterior; SI, primary somatosensory cortex; cSII, secondary somatosensory cortex contralateral to the stimulation; iSII, secondary somatosensory cortex ipsilateral to the stimulation. Adopted from Sakamoto et al. (2008b).

FIGURE 5 | (A) ECD locations for LA, LP, Hand and Foot superimposed on 2D MR images in two representative subjects. Source locations of Subject 1 are superimposed on the coronal plane and those of Subject 2, on the axial plane. The ECDs for bilateral SII responses were located in the upper bank of the Sylvian fissure in the left and right hemispheres. White and gray squares indicate the locations for LA and LP respectively. White and gray circles indicate the locations for Hand and Foot, respectively. (B) Schematic drawing of spatial relationships of the ECDs for SII among each stimulation point. Upper figures depict the medial-lateral and anterior–posterior directions. Lower figures illustrate the superior-inferior and anterior–posterior directions. LA and LP are located most lateral, anterior and inferior, while the ECD for Foot is located most medial, posterior, and superior. Bars indicate standard error (SE). Adopted from Sakamoto et al. (2008b).

NEUROIMAGING STUDIES

Recently, several neuroimaging studies using fMRI and PET have reported human brain activities evoked by somatosensory stimulation of the tongue, to clarify its somatotopic representation (Sakai et al., 1995; Pardo et al., 1997; Miyamoto et al., 2006; Minato et al., 2009). As compared with research on somatosensory processing of the hand and foot, a typical electrical stimulation can not be performed during fMRI recordings, when the tongue area is stimulated, because a magnetic body must not be put into the MR gantry. Thus, some ingenuity is needed to stimulate the tongue.

In an fMRI study, Sakai et al. (1995) stimulated the tip of the tongue from the medial side to the right side (2-cm long) using a cotton swab (stick diameter, 2 mm) with a 30-g bending force and 10-g/mm² bending pressure, and compared the cortical activation with toes and fingertips. They showed the cortical activation, which was located on the contralateral postcentral gyrus, and organized medially-to-laterally in the order of toes, fingertips, and tongue tip.

Pardo et al. (1997) stimulated the right or left side of the protruding tongue at a rate of 1 Hz with a wooden stick, and rCBF was measured using PET. Stimulation of the right side of the tongue produced a contralateral response in SI, while left side stimulation activated bilateral SI.
Miyamoto et al. (2006), who used fMRI, made a long stick with a grooved rubber at its tip. The stick used for stimulation was fixed on a table that was set on both edges of the scanner bed to avoid it touching the subject’s body. The stick was allowed to rotate around its long axis to minimize the possibility of touching the surrounding structures. The oscillating movement of the stick provided oscillating strokes of ~5 mm at the contact zone. The anterior part of the tongue, 1 cm to the right of the midline, was stimulated. Stimulation was provided by the same well-trained experimenter to minimize the variability of stimuli across the subjects. They identified the somatotopic representation of the lips, teeth and tongue in S1, and examined the rostro-caudal changes in the somatotopic organization in S1 in terms of the overlap between each sensory representation. In the rostral portion of the postcentral gyrus, the representation of teeth was located significantly superior to that of the tongue and inferior to that of the lip, consistent with the classical “sensory homunculus” proposed by Penfield, while this somatotopic representation became unclear in the middle and caudal portions of postcentral gyrus. The overlap between each representation in the middle and caudal portions of the postcentral gyrus was significantly greater than that in the rostral portion of the postcentral gyrus.

Minato et al. (2009) also used fMRI, and delivered somatosensory stimulation to either side of the tongue at a constant frequency of 2 Hz with acrylic balls (diameter: 8 mm) attached to the ends of two plastic sticks (inner cylinders) that were incorporated in two plastic tubes (outer cylinders) stabilized at the occlusal surface of the bilateral posterior regions of the splint. The proximal end of the 100-cm extension was attached to the inner cylinders of the mandibular splint, while the distal end was equipped with a stopper at 15 mm, which allowed constant displacement of the acrylic balls against the posterior edge of the tongue on each side. With this set-up, a well-trained experimenter could stimulate a specific targeted region without touching surrounding structures. They investigated whether the pattern of hemispheric cortical activation by tactile tongue stimulation differed, with special attention to the preferred chewing side. As the results, the number of activated voxels in S1 contralateral to the preferred chewing side was significantly greater than that in S1 contralateral to the non-preferred chewing side.

In our MRI study, we compared the brain activities following stimulation of the postero-lateral part of the tongue with those following stimulation of the antero-lateral part (Sakamoto et al., 2010). To stimulate different areas of the tongue, we fabricated an intraoral device, which was the same as used in MEG studies (Sakamoto et al., 2008a,b). In addition, the jaws of the subject were positioned based on centric occlusion and opened about 5 mm between the upper and lower teeth to make a small space that was important to build the projection for stimulating the tongue. These blocks were used to create a wide space from the right to left canine teeth to allow comfortable frontal movement of the tongue. Then, we made four grooves on the lingual side of this device, which were positioned on the lingual cusp of the first premolar of the lower jaw and the distal-lingual cusp of the second molar of the lower jaw bilaterally. Next, we made a projection with polymethylmethacrylate on each groove. The projection has an elliptical shape and is 3 mm in diameter and 3 mm in height. This projection is easily attached and detached, but very stable during experiments.

As a control task, the subjects were required to perform the tongue-protruding movement while no projection was set on the device. In the other four tasks, the subjects performed the movement with a projection. The projection was set at four positions to stimulate specific areas of the tongue: the left antero-lateral, left postero-lateral, right antero-lateral, and right postero-lateral areas (Figure 6). In each task, only one projection was set for the target area. After each session, this projection was replaced with another groove; which was attached and detached by the operator out of the MR gantry. To investigate only the somatosensory-related activation by removing the motor-related activation and somatosensory-related activation for the device and/or intraoral structures, we analyzed subtraction images obtained from the contrasts as follows: the task in which the projection was on the left antero-lateral of the tongue (left antero-lateral) minus Non-projection task (Control) (LA), left postero-lateral minus Control (LP), right antero-lateral minus Control (RA), and right postero-lateral minus Control (RP).

Stimulation of the left and right postero-lateral parts of the tongue induced significant activity in the SI and Brodmann area 40 (BA 40) in the right hemisphere and the anterior cingulate cortex (ACC) (Figure 7). In contrast, antero-lateral stimulation produced activity only in the right SI (Figure 8). The activated region in SI was significantly larger following stimulation of the posterior than anterior part. These results indicate that a clear difference exists in somatosensory processing between stimulation of the antero-lateral and postero-lateral parts of the tongue, and the right hemisphere is dominant for the stimulation of both antero-lateral and postero-lateral areas. As anatomical data in humans, the anterior two-thirds of the tongue are innervated by the afferent fibers that travel in a branch of the trigeminal nerve (V) called the lingual nerve. The
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study, the peripheral parts of the lingual and hypoglossal nerves were dissected in forty human specimens obtained from 25 adult subjects. We speculate that the same connection is formed between the lingual nerve and glossopharyngeal nerve. Doty et al. (2009) also suggested that branches of the glossopharyngeal nerve extend anteriorly beyond the sulcus terminalis and circumvallate papillae, with extensions occurring along the lateral lingual margin anterior to the foliate papillae. In addition, anastomoses were identified between the glossopharyngeal nerve and the lingual nerve, raising the possibility of functional interactions between the trigeminal nerve and the glossopharyngeal nerve. This notion was supported by a recent anatomical study (Zur et al., 2004). Based on these findings, we inferred that the activated region in SI was larger following stimulation of the posterior than anterior part.

In addition to SI, BA 40 immediately lateral to SI was activated during the postero-lateral stimulation. We assumed that the activation of BA 40 included neuronal activation associated with visceral sensation following the stimulation of the posterior tongue. Previous studies demonstrated that the cortical representation of visceral organs differs from that of the somatosensory system, using fMRI (Aziz et al., 2000; Hobday et al., 2001; Lotze et al., 2001; Strigo et al., 2003; Eickhoff et al., 2006; Ladabaum et al., 2007), and MEG (Schnitzler et al., 1999). For instance, Hobday et al. (2001) observed activation in the inferior part of SI and BA 40 following visceral rectal stimulation, but only in SI following somatic anal stimulation. Previous studies demonstrated that the cortical representation of visceral organs differs from that of the somatosensory system, using fMRI (Aziz et al., 2000; Hobday et al., 2001; Lotze et al., 2001; Strigo et al., 2003; Eickhoff et al., 2006; Ladabaum et al., 2007), and MEG (Schnitzler et al., 1999). For instance, Hobday et al. (2001) observed activation in the inferior part of SI and BA 40 following visceral rectal stimulation, but only in SI following somatic anal stimulation. Ladabaum et al. (2007), who recorded brain activity following gastric distension, found no evidence of activation of SI, but found activation in a broad region of BA 40. Taking these studies into consideration, it is likely that BA 40 was activated following visceral stimulation, but more studies will be necessary to determine the relationship between the neural activity of BA 40 and visceral sensation and to clarify the characteristics of this area with respect to tongue stimulation.

In general, the ACC plays an important role in sensory, motor, cognitive, and emotional information (Bush et al., 2000) and pain processing (Schnitzler and Ploner, 2000; Vogt, 2005; Qiu et al., 2006).
Our results demonstrated that the ACC was activated only during the postero-lateral stimulation. Some studies showed the ACC to often be concerned with visceral sensation. For example, Hobday et al. (2001) noted that the ACC was activated by visceral stimulation, not by somatic stimulation, and it appears that their results are consistent with our findings. Thus, we considered that our ACC activation reflected the attributes of the viscera, because the viscera have a complex peripheral nervous system that allows for a wide variety of autonomic functions (Ness and Gebhart, 1990).

Our fMRI results showed a cortical representation in the right hemisphere, but not left hemisphere. By contrast, in our MEG studies (Sakamoto et al., 2008a,b) and neuroimaging studies (Pardos et al., 1997; Minato et al., 2009), bilateral activations were observed, not showing a right dominant response. There are three possible explanations for the discrepancy between our neuroimaging findings and some previous studies including our own MEG studies. The first possibility is that somatosensory processing includes asymmetric neural activation. That is, as several neuroimaging studies already showed (Perlmutter et al., 1987; Fox and Applegate, 1988; Naito et al., 2005; Nihashi et al., 2005; Eickhoff et al., 2008), the brain’s response should be stronger in the right hemisphere than the left for somatosensory processing. We believe that the present study also indicated this asymmetric neural activation. Indeed, our method of stimulation may be unable to elicit clear activation in the left hemisphere, compared to general electrical stimulation. If so, it might be difficult to detect the response in the left hemisphere. The second possibility is a negative motor effect on the left somatosensory areas. Some neuroimaging studies have also provided evidence that activation of the sensorimotor cortex representing the oral and facial regions during voluntary swallowing and mastication showed left hemispheric preference (Martin et al., 2004, 2007; Shinagawa et al., 2004). From these studies, there is a possibility that active movement of the tongue affects SI activity in the left hemisphere. Indeed, many studies have investigated somatosensory-motor integration by recording SEPs during voluntary movement. Characteristically, the amplitudes of short-latency components are attenuated, while those of long-latency are enhanced (Giblin, 1964; Kakigi, 1986; Hoshiyama and Sheean, 1998; Rossini et al., 1999; Valeriani et al., 2001; Nakata et al., 2003), and this phenomenon is termed “gating”. This gating effect has been also researched by recording SEFs, and similar results were found regarding the cortical responses. That is, the early responses generated from SI were attenuated during voluntary movement, whereas the late responses in SII were strengthened (Rossini et al., 1989; Kakigi et al., 1995, 1997; Huttunen et al., 1996; Forss and Jousmäki, 1998; Lin et al., 2000). Such modulation also occurred in an fMRI study (Hinkley et al., 2007). A third explanation is that the above two possibilities may be interrelated.

CONCLUSION

The present study showed the human brain’s response after stimulation of the tongue. Because of technical difficulty in stimulating the tongue, researchers have had to make special devices to clarify the neural mechanisms for somatosensory processing. In the somatotopic representation of SI, the face and oral regions as well as the hand area occupied larger regions (Penfield and Boldrey, 1937), indicating an important role in the human somatosensory system.

Future studies must resolve several issues pertaining to somatosensory processing of the tongue. First, extant studies have focused mainly on the earliest component of SEPs/SEFs, generated from area 3b in SI. They did not analyze other components, which were recorded on stimulation of the hand, such as the P25, N35, P45, N60, and frontal N30 (Nakata et al., 2003; Kida et al., 2004). These components should be analyzed. Second, the characteristics of the tongue SII should be clarified because few studies have investigated the neural mechanisms involved (see Responses from Secondary Somatosensory Cortex). Third, the devices for stimulating the tongue have depended on individual researchers. Therefore, standard methods need to be established.

We believe that our findings provide valuable information on the neural mechanisms relating to the oral and maxillofacial regions.

ACKNOWLEDGMENT

This article was supported by grants from the Japan Society for the Promotion of Science for Young Scientists to Kiwako Sakamoto.

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