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A B S T R A C T
Recently, the cortical mechanisms of tactile-induced analgesia have been investigated; however, spatiotemporal characteristics have not been fully elucidated. The insular–opercular region integrates multiple sensory inputs, and nociceptive modulation by other sensory inputs occurs in this area. In this study, we focused on the insular–opercular region to characterize the spatiotemporal signature of tactile-induced analgesia using magnetoencephalography in 11 healthy subjects. Aδ (intra-epidermal electrical stimulation) inputs were modulated by Aβ (mechanical tactile stimulation) selective stimulation, either independently or concurrently, to the right forearm. The optimal inter-stimulus interval (ISI) for cortical level modulation was determined after comparing the 40-, 60-, and 80-ms ISI conditions, and the calculated cortical arrival time difference between Aδ and Aβ inputs. Subsequently, we adopted a 60-ms ISI for cortical modulation and a 0-ms ISI for spinal level modulation. Source localization using minimum norm estimates demonstrated that pain-related activity was located in the posterior insula, whereas tactile-related activity was estimated in the parietal operculum. We also found significant inhibition of pain-related activity in the posterior insula due to cortical modulation. In contrast, spinal modulation was observed both in the posterior insula and parietal operculum. Subjective pain, as evaluated by the visual analog scale, also showed significant reduction in both conditions. Therefore, our results demonstrated that the multisensory integration within the posterior insula plays a key role in tactile-induced analgesia.

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Introduction
The analgesic effect of vibrotactile stimuli has been acknowledged as an important mechanism that can modulate pain perception (Melzack and Wall, 1965). Practically, people unconsciously rub a painful part of their body to reduce pain. Several studies have shown the analgesic effect of both tactile and vibratory stimuli in psychophysiological evaluation in healthy subjects (Mancini et al., 2014; Yarnitsky et al., 1997) and in chronic pain conditions (Lundeberg et al., 1984; Roy et al., 2003; Staud et al., 2011). Additionally, an electrophysiological study with laser-evoked potentials demonstrated the attenuation of a vertex component by concurrently applied vibratory stimulation (Kakigi and Shibasaki, 1992). Because these studies presented nociceptive and non-nociceptive stimuli simultaneously, cross-modal interaction most likely occurred in the spinal cord. On the contrary, recent somatosensory evoked potential (SEP) and magnetic field (SEF) studies (Inui et al., 2006; Mouraux and Paghki, 2007; Testani et al., 2015) have indicated a supraspinal mechanism in tactile-induced pain relief via experimental paradigms with concurrent bimodal inputs at the cortical level. However, the precise spatiotemporal profile for centrally mediated tactile-induced analgesia is largely unknown.

To date, several neuroimaging studies have demonstrated that the insular–opercular region (IOR) is consistently involved in pain processing (Apkarian et al., 2005; Peyron et al., 2000, 2002). Some reports have also suggested that this region encodes quantitative as well as qualitative aspects of pain (Iannetti et al., 2005; Oertel et al., 2012; Timmermann et al., 2001). Furthermore, the IOR is known to be involved in multisensory processing, including tactile sensation, pain, proprioception, temperature, and vestibular information (Mazzola et al., 2012; zu Eulenburg et al., 2013). Thus, we hypothesized that the IOR is a key site for multisensory interactions, possibly generating tactile-induced analgesia at the cortical level.

In the present study, we sought to elucidate the cortical representation of pain relief in a temporally and spatially precise manner. We adopted mechanical tactile stimulation (MTS) and intra-epidermal electrical stimulation (IES) for selective Aβ–Aδ interaction. MTS has higher selectivity for Aβ fibers (Onishi and Kameyama, 2011), and this is suitable for increasing the sensitivity of source separation between tactile-related and pain-related activity (Baumgärtner and Treede, 2009). IES...
is well known as a selective Aβ stimulation technique (Inui et al., 2002; Inui and Kakigi, 2012; Mouraux et al., 2010). Then, we recorded magnetoencephalographic (MEG) responses with minimum norm estimates (MNE) analysis, which is widely used as a distributed source analysis method enabling better delineation of spatial and temporal features. With respect to spatial localization, it is crucial to determine the three-dimensional segregation within the IOR, because this area has been subdivided into cytoarchitectonically-distinct areas harboring multisensory representations in fMRI studies (Eickhoff et al., 2006a,b; Kurth et al., 2010; Mazzaola et al., 2012; zu Eulenburg et al., 2013). Furthermore, an SEF study with multidipole analysis demonstrated that an IES-evoked SEF response in the insula was localized more anteriorly compared with a transcutaneous electric SEF response (Inui et al., 2003). From several lines of evidence, we decided to assess the spatiotemporal signature of tactile-induced analgesia in the IOR.

Methods

Subjects

Eleven healthy right-handed volunteers participated in this study. The subjects had no history of neurological conditions or chronic pain. They also had no current or prior history of traumatic events or surgery involving the right arm, and were not on any medication. Written informed consent was obtained from all subjects. This study was approved by the Institutional Ethics Committee of Kyushu University.

Aβ and Aβ stimulus

We delivered noxious pain stimuli by IES with the aim of selectively stimulating Aβ fibers (Inui et al., 2002; Inui and Kakigi, 2012; Mouraux et al., 2010). The stimulus consisted of double current pulses separated by a 10-ms interval, and each pulse had a triangular waveform with a 0.5-ms rise and 0.5-ms fall, as per previous reports (Inui and Kakigi, 2012; Omori et al., 2013). The pain intensity was adjusted individually for each subject prior to the experiment. We aimed to produce a definite, but tolerable pinprick sensation, corresponding to 20–40/100 on a visual analog scale (VAS). The VAS range was presented as a horizontal line, which was anchored by “no pain” (score of 0) and “worst imaginable pain” (score of 100), where a score of 5–44 indicates mild pain (Hawker et al., 2011). Stimulus intensities were set lower than 0.3 mA to minimize the possibility of concurrent Aδ fiber stimulation. Mean stimulus intensity for IES was 0.26 ± 0.03 mA.

For the innocuous tactile stimulation, we used piezoelectric actuators (KGS, Saitama, Japan) to control the MTS. This enabled us to produce time-locked Aβ-specific afferent inputs (Onishi et al., 2010). The stimulus probe had eight pins, and each pin protruded 0.7 mm with a maximum pushing force of 0.33 N/pin, producing a sensation that was almost identical to real touch. The mechanical onset delay from the trigger to the highest pin position was less than 0.5 ms, and the stimulus duration was 10 ms. We delivered white noise binaurally throughout the experiment to mask stimulator noise.

Experimental paradigm

IES and MTS were delivered to the right forearm, slightly distal from the elbow. The stimulation points were located in close proximity (<1 cm) to one another so that the inputs from both modalities occurred within the same dermatome. We expected that this would maximize the effect of the tactile analgesia (Mancini et al., 2014). Before beginning the experiment, we estimated the cortical arrival times of the Aβ and Aδ inputs using the known conduction velocities (CVs) of each fiber and the distance from the forearm to the C1 vertebra of the spine (Inui et al., 2006). The average distance between the forearm and the spinal cord (i.e., the C7 vertebra of the spine) was approximately 50 cm, and that between C7 and C1 was approximately 10 cm. Given that the CV of the Aβ fiber is 10–15 m/s (Kakigi and Shibasaki, 1991) and that of the Aδ fiber is 60 m/s (Inui et al., 2006), we estimated that it would take 42 ms more for the Aδ impulse to reach the spinal cord (we adopted the slowest CV of the Aδ fiber, i.e., 10 m/s for the calculation). Similarly, the CVs of the spinothalamic tract and the posterior columns are 8–10 m/s and 50–60 m/s (Kakigi and Shibasaki, 1991), respectively. Therefore, we estimated the difference in conduction time within the spinal cord to be 11 ms. Thus, we predicted that the cortical arrival time of the Aδ projection would be 53 ms later than that of Aβ. We also conducted an experiment to estimate the CV of the Aβ fiber in three subjects by stimulating the right dorsal hand and shoulder using IES.

Because the inter-stimulus interval (ISI) between IES and MTS was critical in this study, we performed two experiments to determine the optimal ISI for cortical level modulation (Exp. I) and to assess cortical modulation at a fixed ISI (Exp. II). In Exp. I, we adopted four different stimulus conditions: a) control IES, b) control MTS, c) 0-ms ISI (IES and MTS were applied simultaneously), and d) three types of ISI settings (Fig. 1). Three different ISI settings (40-, 60-, or 80-ms) were tested to determine the optimal ISI for the most effective suppression at the supraspinal level. Seven subjects (three females; age, 25–49 years) participated in this experiment. The experiment comprised five sessions, and each session was separated by 3 to 4 min of rest to avoid any attenuation of responses as well as habituation of the subjective pain sensation. Ten trials were performed for each condition in each session. Therefore, each session consisted of 40 randomly ordered trials with inter-trial intervals of 4 to 5 s. According to the results of Exp. I (see Results), we used the 60-ms ISI condition for the cortical inhibitory condition in Exp. II (Fig. 1d). Thus, the 0-ms ISI (IES and MTS were applied simultaneously) was designed to determine interaction at the spinal level, while the 60-ms ISI was formulated to elucidate the presence of tactile-induced analgesia at the supraspinal level. The seven subjects from Exp. I and four further subjects were recruited for Exp. II. Thus, a total of 11 subjects (four females; age, 21–49 years) were examined. The subjects rated subjective pain intensities for each condition using a VAS prior to MEG measurements. Twenty trials (five trials for each condition) were randomly delivered using the same inter-trial intervals, and the subjects gave a VAS score for each individual stimulus.

Data acquisition

SFs were recorded using a 306-channel whole-head MEG system (Neuromag; Elekta, Helsinki, Finland). Prior to recording, we attached four head position indicator (HPI) coils to the scalp, and used a 3D digitizer (Fastrack; Polhemus, Colchester, VT, USA) to measure the anatomical landmarks (nasion and left and right pre-auricular points), the HPI positions, and the head surface. We conducted HPI measurement immediately before the first session, and the subject maintained their initial head position throughout the recording. The subjects were seated in an upright position and instructed to keep their eyes open. Their vigilance was carefully monitored by a video camera positioned in a shielded room. The MEG signals were digitally sampled at 1000 Hz with a band-pass filter of 0.3–330 Hz. Structural MRI was obtained using a 3.0-T high resolution MRI scanner (Achieve; Philips N.V., Eindhoven, The Netherlands) for MEG-MRI co-registration and subsequent source localization analysis (repetition time 8.2 ms; echo time 3.8 ms; flip angle 8°; 190 sagittal slices; 1.0 mm isotropic voxels with no gap).

Data analysis

We processed all data sets using the spatiotemporal signal separation method (Taulu et al., 2004) to filter out noise signals from outside of the head. We applied a bandpass filter in the 1–57 Hz range, and trials above a rejection threshold of 5000 fT/cm for gradiometer channels were removed before averaging. The artifact-free data were averaged based on the onset of each stimulus in the control condition, where
averaging was triggered by the onset of IES for each ISI setting (40-, 60-, or 80-ms). Ten trials multiplied by five sessions (50 trials) were conducted. In Exp. I, 12 data sets [4 conditions ((a) control IES, (b) control MTS, (c) 0-ms ISI, and (d) delayed- ISI settings (40-, 60-, or 80-ms))] were obtained and group averaged for seven subjects after applying source reconstruction and transformation into the standard brain during source analysis. In Exp. II, four data sets [four conditions ((a) control IES, (b) control MTS, (c) 0-ms ISI, and (d) 60-ms ISI)] were obtained and group averaged for 11 subjects.

Using FreeSurfer software (Dale et al., 1999; Fischl et al., 1999), we reconstructed the cortical surface for each subject using T1-weighted images. We used MNE-suite (http://www.martinos.org/mne/) (Dale and Sereno, 1993; Dale et al., 2000; Hämäläinen and Ilmoniemi, 1994), which is a distributed source model for MEG analysis, to ensure rich spatiotemporal resolution for current sources over the cortical surface in our source localization analyses. The extracted MRI contour was then imported to co-register with the MEG head coordinate system using the scalp surface points obtained by the 3D digitizer. The general procedures for MNE and dynamic statistical parametric mapping (dSPM) (Dale et al., 2000; Molins et al., 2008) were based on our previous studies (Hironaga et al., 2014; Kanamori et al., 2013; Miyaji et al., 2014). For noise normalization in dSPM, we created a noise covariance matrix from −200 to −100 ms in the pre-stimulus period.

Fig. 2 shows a schematic image of the methodological procedure. First, we defined a region of interest (ROI) for the contralateral IOR on the standard brain (MNI305; provided by FreeSurfer software, MNI = Montreal National Institute (Collins et al., 1994), and then projected this onto each individual brain (Fig. 2a). Within the IOR, the parietal operculum is commonly referred to as the secondary somatosensory cortex (SII) in reference to its functional aspect. Nevertheless, the definition and localization of SII varies from study to study (Eickhoff et al., 2006b), hence we decided to use the parietal operculum as the anatomical description for SII in this study. For optimal spatial segregation between the activities elicited by the control IES and control MTS, we defined a restricted ROI (rROI). Fig. 2b shows the encircled rROI from the activation patterns of group-averaged data. We did this by narrowing down the activated vertices to reveal a region where the dSPM value was larger than 90% of the highest values in a 20-ms time window centered at a peak latency:

\[
\text{rROI} = \{ \nu_i : d_{\text{dSPM}}(\nu_i) > \max_i(d_{\text{dSPM}}(\nu_i, t)) \times 0.9; \nu_i \in \text{ROI}, t \in t_p \pm 10_{\text{ms}} \}
\]

where \(d_{\text{dSPM}}(\nu_i)\), ROI, and \(t_p\) indicate a vertex index, dSPM value of the vertex \(\nu_i\), the original ROI (i.e., IOR), and a peak latency, respectively. The peak latencies for each modality were initially identified in terms of root-mean-square waveforms for approximately nine pairs of gradiometers located at a contralateral temporal area. This area was first located by visual inspection, and we then confirmed that these latencies corresponded to the peak latencies of source activation curves (source waves) obtained from dSPM values within the original ROI. We used weighted averaging (Hironaga et al., 2014) to determine the center of the vertices within the rROI as follows:

\[
\text{center} = \frac{1}{\sum d_{\text{dSPM}}(\nu_i)} \sum d_{\text{dSPM}}(\nu_i)P(\nu_i)
\]

where \(P(\nu_i)\) represents the vertex position. After calculating the center coordinates for each modality, we transformed the coordinates in the individual MRIs (expressed in terms of the Right Anterior Superior (RAS) coordinate system) onto the standard brain using the MNI-Talairach coordinate system. We also created group-averaged MNE

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**Fig. 1.** A schematic image of the experimental procedure for Experiments I and II. Red bars indicate intra-epidermal stimulation (IES), while blue bars represent mechanical tactile stimulation (MTS). IES was delivered to the right forearm, while MTS was applied in close proximity to IES. In the control conditions, IES (a) and MTS (b) were given independently as unimodal stimulations. In the bimodal conditions, IES and MTS were applied simultaneously to assess the spinal-level interaction (i.e., 0-ms ISI) (c), whereas IES was applied prior to MTS to assess the cortical-level interaction (40-, 60-, or 80-ms ISI) (d). Each of these four conditions was delivered randomly with 4–5 s inter-trial intervals.
activation maps after transformation into the standard brain for summed control and actual bimodal conditions with standardization. Standardization by dividing by the maximum values of two conditions, control and bimodal, was performed to retain the activation ratio in each individual. This process was applied for two ISI settings separately.

To investigate tactile-induced interactions in pain processing, we focused on the region most relevant to pain-related activity, namely, the rROI for the control IES. To this end, we marked all activities identified as rROIs for the control IES (Fig. 2c), and reconstructed source waves for each stimulus condition (Fig. 2d). Here, we defined the source wave amplitudes from the 0-, 40-, 60-, and 80-ms ISI conditions as the amplitudes of the actual bimodal stimulation, and compared these with the sum of the waveforms obtained in the control IES and MTS conditions (Gondan and Röder, 2006; Hsieh et al., 1995; Torquati et al., 2003; Wühle et al., 2011). Since the MTS was delivered later than the IES in all ISI conditions, we shifted the waveform of the control MTS by +40, 60, or 80 ms to simulate these conditions before summation of waveforms. We used the following equation to calculate the interaction ratio (IR) converting into percentage. This allowed us to express the degree of amplitude reduction that occurred in each bimodal condition (Ishibashi et al., 2000):

\[
IR(ISI) = \frac{d_{SPM}(cIES) + d_{SPM}(cMTS) - d_{SPM}(Bi)}{d_{SPM}(cIES) + d_{SPM}(cMTS)} \times 100
\]  

where cIES, cMTS and Bi denote the control and bimodal conditions (Bi signifies the 0-, 40-, 60-, or 80-ms ISI condition), and terms that are parenthesized and include \(d_{SPM}\) with an overbar represent the averaged \(d_{SPM}\) value in a 50-ms time window centered at a peak latency in each condition (see Results for a detailed account of how we set the time-window of interest).

Statistical analysis

Originally, to determine the optimal ISI for cortical pain modulation, we evaluated the difference in IRs among 40-, 60-, and 80-ms ISI conditions using analysis of variance (ANOVA) with a randomized block design and stimulus condition as a factor. We compared the center coordinates of the control IES and control MTS conditions using ANOVA with a randomized block design and stimulus modality as a factor. To analyze the difference in VAS, we used an ANOVA with a randomized block design and stimulus condition as a factor. An arbitrary contrast was analyzed for evaluation of the tactile-induced analgesic effect by applying contrast coefficients \((1, -0.5, -0.5)\) for each condition (control IES, 0-ms ISI, 60-ms ISI, respectively). We employed an ANOVA with a randomized block factorial design and two factors; stimulus condition (0-ms ISI or 60-ms ISI) and measurement method (actual bimodal or summed), to evaluate source wave amplitudes. The least square means were used in Tukey’s Honestly Significant Difference post-hoc test for VAS and the source wave amplitudes. We evaluated the difference in IRs between the 0-ms ISI and 60-ms ISI conditions using a paired \(t\)-test. All data are mean ± standard error (SE), and \(P < 0.05\) was considered significant.

**Fig. 2.** Methodological procedures. (a) A region of interest (ROI) in the contralateral insular-opercular region (IOR) in the brains of the two subjects: subject 1 (S1, upper) and subject 2 (S2, lower). A reference ROI for the contralateral IOR was first marked on the standard brain, and then projected onto each individual brain. (b) Selection of the restricted ROI (rROI) (purple outline). dSPM values larger than 90% of the highest value within the ROI (a) were extracted (Eq. (1)) and encircled. (c) Marked rROI (purple) of the control IES for the analysis of tactile-induced interactions; (b) and (c) are the enlarged views of the white rectangles (a). (d) source waveforms for the control IES (red line) and control MTS (blue line) conditions obtained from the encircled rROI. The a.u. (arbitrary unit) in the ordinate denotes dSPM values. (e) Label annotation based on FreeSurfer software (inside the brackets is original FreeSurfer annotation), 1) insula = \((G_{circular\_insula\_sup} + G_{insular\_short} + G_{ins\_lg\_and\_S\_cent\_ins})\), 2) subcentral cortex = \((G_{and\_S\_subcentral})\), 3) Inferior parietal cortex = \((G_{pariet\_inf\_supram})\), and 4) lateral fissure and retroinsular area = \((Lat\_Fis\_post)\).
Results

Physiological measures and behavioral results

SEFs elicited by IES stimulation were recorded in three subjects in a separate experiment to confirm selective stimulation of Aδ fibers. The estimated CV based on peak latencies (149 ± 14.2 ms and 98 ± 6.2 ms for the right dorsal hand and shoulder, respectively) and distances (mean distance; 56 ± 4 cm) in the three subjects was 11.5 ± 1.4 m/s, which was reasonable for selective Aδ stimulation.

The mean VAS ratings for the control IES, 0-ms ISI, and 60-ms ISI conditions were 22.5 ± 3.2, 16.2 ± 2.4, and 16.6 ± 2.6, respectively. We found a significant main effect of stimulus condition (F (2, 20) = 11.7; *P* < 0.001), and the arbitrary contrast revealed a significant difference among the control IES and the two bimodal conditions (i.e., 0-ms ISI and 60-ms ISI) (F (1, 20) = 23.3; *P* < 0.001). This implied that the MTS had an inhibitory effect on subjective pain ratings in both bimodal conditions. There was no significant difference between the two bimodal conditions, indicating that the magnitudes of reduction in the VAS ratings for the two conditions were not statistically different.

Comparison of different ISI settings

Fig. 3 shows the temporal profile of mean source waveforms (n = 7) elicited by actual bimodal stimulation (blue) and summed control responses (red) in three different ISI settings (40-ms (a), 60-ms (b), and 80-ms (c)), and related statistical results (d) in Exp. I. Pain inhibition was observed in the 40-ms and 60-ms conditions, but was not evident in the 80-ms ISI condition. In addition, SEs in the 80-ms ISI condition (Figs. 3c and d) showed large inter-subject variability. We were unable to observe pain inhibition in three out of seven subjects. Accordingly, IRs were similar between the 40- and 60-ms conditions (30.0 ± 9.2% and 26.0 ± 9.4%, respectively, see also Fig. 3d), but IRs were considerably smaller in the 80-ms condition (5.8 ± 18.9%). No significant main effect was found for the condition (F (2, 12) = 0.76; *P* = 0.49). Considering that the calculated difference in cortical arrival time between IES- and MTS-delivered stimuli was 53 ms, it is reasonable to assume that pain inhibition in the 40-ms ISI condition could be affected at the spinal level. Concerning the 60-ms and 80-ms cortical conditions, no rationale was provided for employing the 80-ms ISI condition because of small IRs with considerable inter-subject variability. Thus, we determined that the 60-ms ISI condition was optimal for later analysis of cortical inhibition.

Pain modulation of the source waveforms and MNE activity

The suppression of pain activity by simultaneous tactile stimulation (0-ms ISI) was compared with the delay paradigm (60-ms ISI). Figs. 5a and b show the temporal profile of the mean source waveforms elicited by actual bimodal stimulation (blue) and summed control responses (red). The color-matched transparent areas represent the positive and the negative SEs of the mean. As shown in Figs. 5a and b, the peak responses of the group-averaged data were reduced by bimodal stimulation. Moreover, the small SEs suggested low variability across subjects. We compared the mean amplitudes of the summed source waves with those obtained from actual bimodal stimulation within the separate time windows (Figs. 5a and b, gray shadings), because peak latencies of the group-averaged data in both conditions were shifted slightly. A significant main effect of the measurement method (F (1, 30) = 17.3; *P* < 0.001) was found, which indicated that both amplitudes in the actual bimodal conditions (0-ms ISI and 60-ms ISI) were lower than those of the summed control responses where no interaction occurred. There was no significant effect of the interaction between the stimulus condition and the measurement method (F (1, 30) = 0.7; *P* = 0.40), which implied that the two actual bimodal conditions caused suppression of the activity to the same degree. Although the IRs...
focused on in this study. The effect was not statistically significant. The percentage difference was 10.9% (9.9%) compared with the 60-ms ISI condition (16.7 ± 9.9%), this difference was not statistically significant (P = 0.13). Taken together, these statistical results indicated the cortical level interaction was equipotent to the spinal level interaction in our experimental condition.

Pain modulation within the IOR

Figs. 5c and d show spatial profiles of group-averaged MNE results for summed control and actual bimodal conditions under the 0-ms ISI and 60-ms ISI conditions with standardization. Summed activation was located within the posterior insula (Fig. 5c-R1 and Fig. 5d-R3) and the parietal operculum (Fig. 5c-R2 and Fig. 5d-R4) in the 0-ms and 60-ms ISI conditions, and no clear difference was observed between the conditions. Consistent with source-wave amplitude reductions (Figs. 5a and b), source activation maps indicated a reduction in brain activity in the 0-ms and 60-ms ISI conditions. In the 0-ms ISI condition, a strong reduction in activity was observed in the posterior insula (Fig. 5c-R1) and parietal operculum (Fig. 5c-R2). In the 60-ms ISI condition, reduction in activity was predominant in the posterior insula with a modest reduction in the most posterior part (Fig. 5d-R3) and the parietal operculum (Fig. 5d-R4).

Discussion

The present study clearly demonstrated pain relief by tactile input at the spinal and cortical level, showing the spatiotemporal signature of tactile-induced analgesia. Our results demonstrated the difference in cortical source locations between pain and tactile processing with modality-selective stimulation. The observed temporally-specific neural activities underpin the previous fMRI findings, which segregated multisensory representations within the IOR. Most importantly, posterior insular activity (Fig. 5d-R3) was suppressed by cortical-level pain relief.

Differential representation of Ao and Aβ processing in the IOR

We found a clear functional localization of Ao and Aβ inputs within the IOR. Using MNE, the two modality-specific activities were successfully identified as anterior-to-posterior distinct patterns of activation (Fig. 4a), in agreement with previous fMRI studies (Mazzola et al., 2012; zu Eulenburg et al., 2013), which consolidated differential multisensory representations within the IOR. An argument may be made that MNE is less sensitive for determining focal activities compared with other techniques such as dipole analysis or beamforming techniques. Nonetheless, the two activation clusters were clearly grouped and we could identify the center of the cluster, in accord with our previous study on laryngeal somatotopic organization (Miyaji et al., 2014). Hauk et al. (2011) also suggested that estimation of the insula using dSPM showed maximum activation around the true source location in a simulation study.

In the light of localization results, progress is required on anatomical details. Both IES and MTS-elicited activity was distributed in the proximity of the border between the posterior insula and parietal operculum (Fig. 4a), yet the x-direction showed no significant difference (Fig. 4b). Both IES and MTS are recent techniques, and detailed research, including intracranial studies, using these techniques is lacking. Thus, our study provides novel information regarding their spatial segregation. IES localization results were within the range evaluated by an intracranial recording of laser-evoked potentials (Frot et al., 2014), while MTS was localized deeply within the parietal operculum (Eickhoff et al., 2006a). Unlike electrical Aβ stimulation, MTS induces natural touch-like sensations, but weaker evoked responses. This difference has a significant influence on source localization accuracy. In fact, single-dipole
analysis could not segregate the contralateral opercular responses (Onishi et al., 2010; Onishi and Kameyama, 2011). In contrast, our MNE-based approach successfully segregated opercular activity elicited by MTS. Subsequently, we were able to clearly distinguish the source location for $\alpha$-$\alpha$ interactions associated with natural-touch like sensation.

**Temporal features of pain–tactile interaction**

On the basis of distinct spatial source localization for each stimulus modality, we investigated the timing of the pain–tactile interaction. A previous SEF study (Torquati et al., 2003) of the somatosensory gating effect demonstrated that the simultaneous presentation of painful and non-painful electric stimuli resulted in a smaller SII response compared with the sum of the responses elicited by each stimulus separately. Other studies also used the same analytical method to assess multisensory interactions with event-related potentials (e.g. Gondan and Röder, 2006). If no interference exists between the two different sensory modalities, the amplitude obtained by summation of the evoked responses to the separate-modality stimuli should be equal to that obtained by combined-modality stimulation (Hsieh et al., 1995; Ishibashi et al., 2000). To date, few studies have demonstrated that the pain–tactile interaction occurs strictly at the supraspinal level (Inui et al., 2006; Mouraux and Plaghki, 2007; Testani et al., 2015). This is likely because some of former studies did not avoid interference at the spinal cord. We arranged a time delay by adjusting the timing of tactile input to ensure nociceptive input reached the cortex earlier than tactile input, or at least simultaneously. Therefore, the amplitude difference for the bimodal stimulation condition in the present study effectively represents tactile-induced analgesia at the spinal cord and IOR. The source waveform analysis enabled us to specifically investigate the temporal features of pain modulation occurring within the IOR. The source waveform analysis enabled us to specifically investigate the temporal features of pain modulation occurring within the IOR. The statistically significant reduction of source waveforms of bimodal stimulation was clearly observed in the group-averaged data (Figs. 5a and b). In both spinal and cortical conditions, the suppression of waveforms was observed in the 100–200 ms time window. The finding that peak latency of the bimodal source waveforms in the 0-ms ISI condition was relatively close to peak latency in the tactile control condition suggests that most cortical activity could be elicited from tactile input, due to pain inhibition at the spinal cord level.

In contrast, the peak latency of the bimodal stimulation condition in the 60-ms condition (Fig. 5b) was approximately 150 ms, which differed from latencies obtained from the tactile control condition. If

![Fig. 5. Comparison of spatiotemporal profiles between the two bimodal conditions. Group-averaged (n = 11) source waveforms extracted from the rROI (Fig. 2c) in the two conditions ((a) 0-ms ISI condition (b) 60-ms ISI condition) for the actual bimodal stimulation (blue lines) and the summed control waveforms (red lines). Color-matched transparent areas represent the SEs of the mean. Gray shaded areas in (a) and (b) indicate the evaluation of interaction ranges where reduction in amplitude occurred in the selected time windows: 95–145 ms (a) and 110–160 ms (b) ($P < 0.001$). Group-averaged source localization maps of the ranges of the gray shaded area in the two conditions for (c) summed and actual bimodal activations in the 0-ms ISI condition, and (d) in the 60-ms ISI condition. Dotted white circles in (c) and (d) (R1–4) are the regions that we focused on in this study. The figures show an enlarged view of the area surrounded by the white rectangle (right). Each label depicts the same structures as in Fig. 2e.](image-url)
nociceptive inputs were inhibited at the spinal cord (i.e., overridden by tactile input), the peak latency for the 60-ms ISI condition could be the same as the 0-ms ISI condition; shifted +60 ms rightward, i.e., around 185 ms. In short, nociceptive input predominates when cortical processing is initiated, and after this period, activity begins to be inhibited. Based on this assumption, it can be concluded that peak latency for the 60-ms ISI condition reflects integration of pain and tactile information in the posterior insula, whereas reduction in nociceptive input with less cortical modulation occurs in the 0-ms ISI condition.

Spatial features of pain–tactile interaction

In addition to the temporal characteristics of pain modulation, the difference in spatial features between the spinal and cortical suppression of pain were clearly observed within the IOR (Figs. 5c and d). Torquati et al. (2003) reported a suppressive effect for bimodal (painful and non-painful) simultaneous stimulations in the contralateral SII region. The authors suggested that the spatial overlap in nociceptive and non-nociceptive neural populations of SII could explain this effect. In addition, they argued that the functional connection between pain and tactile neuronal populations should be considered an inter-modality integrative effect in SII. In the present study, the source localization results of the 60-ms ISI condition showed clear suppression of activity in the posterior insula (see comparison of R3 in Fig. 5d), which suggests involvement of the posterior insula in nociceptive and non-nociceptive integration, i.e., cortical pain modulation.

The insula plays a key role in pain-related brain activity; however, the anterior and posterior insula have been postulated to behave differentially. Several lines of evidence support the importance of the posterior insula for sensory aspects of pain processing (Lannetti et al., 2005; Oertel et al., 2012; Timmermann et al., 2001; zu Eulenburg et al., 2013). Direct recording from the insular cortex using stereotactic electrode implantation also proved that sensory information processing is predominant in the posterior insula (Frot et al., 2007, 2014; Isnard et al., 2011). In contrast, the anterior insula encodes emotional-affective qualities of pain (Gu et al., 2012; Singer et al., 2004). It is likely that the equivocal activation of the anterior insula compared with the posterior insula is due to the relatively weak pain intensity in our experimental design, by which an aversive response was not induced. Specifically, our pain stimuli delivered a pain sensation without unpleasantness. Thus, along with the role of the posterior insula in cortical pain modulation, our findings mainly represent sensory interaction in quantitatively- and qualitatively-discriminative aspects of pain rather than affective aspects.

Spinal and cortical mechanisms of tactile-induced analgesia

The multi-receptive, i.e., wide dynamic range (WDR), neurons in the dorsal horn of the spinal cord receive information from both nociceptive and non-nociceptive inputs, and are believed to be involved in spinal level pain modulation (Le Bars, 2002; Mancini et al., 2014; Saltar and Henry, 1990). IES input should attenuate at the spinal cord due to reduced firing of shared WDR neurons, and therefore, only tactile-elicited impulses reached the cortex in the 0-ms ISI condition. Given that most IES input was attenuated at the spinal cord as a result of tactile-induced inhibition, such spinal-level inhibition may be more effective than cortical-level inhibition. We found that source amplitude decreased with decreasing VAS scores in the 0-ms ISI condition. However, no significant difference between the spinal and cortical pain-relief conditions was found. One possible explanation for this is that the relatively weak pain intensity, as demonstrated by lower VAS scores, resulted in a modest analgesic effect at the spinal level.

As discussed above, spatial overlap of the convergent neural population in the posterior insula could be a candidate mechanism underlying cortical pain modulation. The existence of a convergent area for multisensory processing is also suggested by an intracerebral recording study that compared the spatial distribution of laser-SEPs with those of electrical SEPs in the insular-opercular cortex (Frot et al., 2001). The posterior insula, particularly the granular area (Ig), has both anatomical and functional connections with SII and limbic areas, and the SII-Ig-limbic pathway is important for both pain and tactile processing (Augustine, 1996; Friedman et al., 1986; Schneider et al., 1993). In addition to this pathway, the insula receives direct input from the ventral medial thalamic nucleus (VMpo), which is involved in the cortical processing of pain (Willis et al., 2002). Moreover, single-unit recording studies in monkeys have also identified WDR neurons within the insula (Robinson and Burton, 1980; Zhang et al., 1999), and the Ig has been recognized as one of the main regions containing this type of neuron (Robinson and Burton, 1980). Because the insula projects directly to the VMpo, it is possible that the posterior insula, rather than SII, predominates in inter-modality integration. As shown in our result, the localized brain area, i.e., the posterior insula, could further consolidate convergent pain and tactile processing with the interaction between the two modalities. Thus, the present study suggests that the posterior insula is an important site for pain inhibition by tactile input.

Limitations of this study

We limited stimulus intensity to less than 0.3 mA to minimize contamination with Aβ inputs from IES stimulation (Inui and Kakigi, 2012; Mouraux et al., 2010). This enabled us to independently evaluate modality-specific interactions, albeit with relatively low VAS ratings. We conducted VAS ratings in a separate session to reduce the length of already long experiments, which may have led to a decrease in the concentration of the subjects. A simultaneous recording of MEG and behavioral data is a methodological option we will consider in future studies, as this would enable better quantification of the causal relationship between source amplitude reduction and subjective pain relief. Ipsilateral responses were not analyzed in the present study. Although previous EEG/MEG studies analyzed bilateral responses (Inui et al., 2003; Lim et al., 2012; Timmermann et al, 2001), contralateral activity is generally more reproducible (Mima et al., 1998; Peyron et al., 2002). This was partly true in the present study, as ipsilateral pain-related activity was not clearly obtained and had considerable amplitude variability. Consequently, we focused on consistent contralateral responses to analyze the pain inhibitory process, while minimizing inter- and intra-individual variability.

Conclusion

Pain–tactile interaction at the cortical level was investigated using carefully timed sensory modality-specific stimuli, in combination with cortical source-based neuromagnetic analysis. We were able to clearly distinguish source location from pain and tactile inputs associated with natural-touch like sensation. Our study clearly demonstrates that the spatial feature of cortical modulation by nociceptive input was different compared with spinal modulation. The current study also provides new spatiotemporal evidence that the posterior insula is a critical area of the cortical mechanisms underlying tactile-induced analgesia in humans. Therefore, our findings contribute to the detailed electrophysiological exploration of central mechanisms underlying cross-modal sensory interactions.

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