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Sensorimotor alpha activity is modulated in response to the observation of pain in others

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INTRODUCTION

When seeing a football player receive a painful tackle we cringe and might even grasp our own knee in affective resonance with the victim’s painful state. This phenomenon of vicarious pain experience is explained in the perception–action account of empathy: “The attended perception of the object’s state automatically activates the subject’s representation of the state, situation, and object, and the activation of these representations automatically primes or generates the associated autonomic and somatic responses, unless inhibited” (Preston and de Waal, 2002).

The perception–action account of empathy for pain has been greatly expanded upon in the last decade, resulting in a nuanced neuroscientific framework that integrates knowledge of affective and perception–action processes with an understanding of the influences of social context, expectation, and attention (for a comprehensive review on its evolutionary basis and social expression). This framework involves a network of affective brain regions, such as the anterior cingulate, paracingulate gyrus, and anterior insular, that are also activated during the first-person experience of pain (Ingvar, 1999; Rainville, 2002). Furthermore, responses in these regions have been found using a wide variety of paradigms, from presenting abstract cues of other people in pain (Singer et al., 2004; Jackson et al., 2005; Saarelva et al., 2007), to pictures of body parts being pinpricked (Lamm et al., 2007), and painful facial expressions (Jabbi and Keysers, 2008). Empathetic responses in these brain regions have, furthermore, been shown to be influenced by social contexts such as group membership and perceived fairness (Hein et al., 2010) as well as task demands and knowledge about the reality of the stimulus (Gu and Han, 2007). Often, but less consistently, primary somatosensory regions are found to be involved, which seems to depend...
on the availability of the sensory information about the painful stimulus. In their meta-analysis of nine fMRI experiments, Lamm et al. (2011) showed that vicarious activation of the somatosensory cortex seems only to occur when visual details of the flesh-and-bone aspect of the painful situation are observed, not when these are inferred from abstract cues. The authors propose, however, that this somatosensory activation reflects unspecific co-activation elicited by the visual display of body parts rather than a specific matching of the other’s somatosensory and nociceptive state. This is in line with their aforementioned characterization of empathy as, first and foremost, an affective state. Other authors, however, have argued for the functional importance of primary sensory cortices as part of the empathetic (pain) response, subserving pain intensity, and location coding (Keysers et al., 2010). Interestingly, in this respect individual propensities differ greatly, as shown by Osborn and Derbyshire (2010). They reported that those people that respond to painful images by experiencing a “real” sensation of pain, show activation of somatosensory cortices, while these regions are not activated in those that do not have such first-person experiences. Further evidence for the involvement of sensory cortices in pain observation comes from electroencephalography (EEG) studies. Bufalari et al. (2007) recorded a reduction of early sensory-evoked potentials after medial nerve stimulation when subjects were watching movies of limbs in painful situations. Pain systems are also tightly linked to action systems (Ingvar, 1999; Saitoh et al., 1999; Juottonen et al., 2002; Farina et al., 2003; Wager et al., 2004), and it has been repeatedly shown that observation of painful movies strongly inhibits corticospinal excitability specific for the muscle that was observed being pinpricked (Avenanti et al., 2005, 2006, 2009). These findings together suggest that primary sensorimotor regions are indeed involved in the empathic response in ways that entails more than aspecific increases in arousal.

Questions of when empathetic responses occur have been investigated using EEG scalp recordings. Fan and Han (2008) found larger early (140–380 ms) frontal event-related potentials (ERPs) amplitudes in response to pictures of limbs in painful situations compared to those in control situations. These early empathic responses were influenced by contextual reality (real pictures versus cartoons). Later (380–500 ms) central–parietal effects of pain that were prominent in a pain judgment task, were greatly reduced when subjects only had to count the number of limbs. This shows that the empathetic response can be modulated at different times, due to different task contexts, in line with a model of empathy that permits modulation of an automatic perception–action response at multiple stages (de Vignemont and Singer, 2006). Frequency analysis of EEG and magnetoencephalography (MEG) recordings has also been a particularly successful tool in studying the involvement of the sensorimotor system in action, touch, and pain observation. Alpha (∼10 Hz) and beta (∼20 Hz) oscillations originating from the sensorimotor cortex (Hari and Salmelin, 1997) have been implicated in action observation (Hari et al., 1998; Cochin et al., 1999; Jarvelainen et al., 2001; Rossi et al., 2002; Muthukumaraswamy and Johnson, 2004; Muthukumaraswamy et al., 2004; Nakamura et al., 2004; Pineda, 2005; Kilner et al., 2006; Caetano et al., 2007; Oberman et al., 2007; Holz et al., 2008; Koelewijn et al., 2008; van Elk et al., 2008) recognizing point-light biological motion (Ulloa and Pineda, 2007), as well as in understanding communicative gestures (Nakamura et al., 2004). Muthukumaraswamy and Johnson (2004) were the first reporting a reduction of the beta rebound after medial nerve stimulation when subjects concurrently observed a hand being brushed or pricked, but not when only movement was observed. Cheng et al. (2008) also observed reduced alpha rebound after medial nerve stimulation while people watched static pictures of limbs in painful situations. Although these studies point to a modulation of the somatosensory cortex, they do not show how somatosensory oscillations respond to the observation of pain in the absence of actual somatosensory stimulation. To investigate such a visual-to-somatosensory process, three EEG studies investigated effects on ongoing alpha oscillations after observing images of painful situations versus control images, without a contingent transcranial or median nerve stimulation. Two of these studies showed more sensorimotor alpha suppression in response to pain than in response to control images (Yang et al., 2009; Perry et al., 2010), while a third study showed reduced alpha suppression (Mu et al., 2008). One reason for these contradictory outcomes might have been the fact that volume conduction makes it difficult to separate sensorimotor alpha (or mu-rhythm) from posterior alpha sources in EEG scalp recordings (Hari and Salmelin, 1997). Since the strongest modulation of alpha power typically involves alpha-blocking in response to visual stimulation (Pfurtscheller et al., 1996), this activity might have confounded the interpretation of alpha activity from central sources that was found by Mu et al. (2008). Indeed, Perry et al. (2010) only found increased suppression by pain observation on fronto-central but not on posterior sensors, while Yang et al. (2009) only found increased central alpha suppression. Although these findings strongly suggest alpha suppression in response to the observation of pain in others, they suffer from a lack of spatial resolution needed to univocally establish a sensorimotor origin. These studies also suffered from underspecified or confounded time windows of interest. In Perry et al. (2010) alpha suppression was calculated over the full 2 s post-stimulus, from stimulus-onset to stimulus offset, while in Yang et al. (2009) the first 1.3 s directly after stimulus-onset were used. In both cases stimulus-onset evoked responses were therefore included, making it ambiguous to what degree their observations can be interpreted exclusively as a modulation of ongoing alpha activity and to what degree evoked responses contributed (Steriade et al., 1990; Lopes da Silva, 1991; Pfurtscheller and Lopes da Silva, 1999; Mazaheri and Jensen, 2010). Mu et al. (2008) did perform an analysis on separate time windows and reported modulation of the alpha suppression only between 200 and 400 ms after stimulus-onset. They also tried to minimize the effect of phase-locked activity on the power estimate by subtracting average ERP in response to stimulus-onset. Since images were presented for only 200 ms, the power estimation might have still been confounded by transients in response to stimulus offset. Finally, the evaluation of every stimulus in Mu et al. (2008) might have resulted in motor-preparation which previously has been shown to interact with sensorimotor oscillations in response to pain (Babiloni et al., 2008). Concerns about the mixing of central and posterior sources, evoked activity and motor-preparation aside, the differences in the direction of alpha modulation might also point to the interesting possibility that different functional processes were involved. Historically, alpha activity has been interpreted as reflecting a
non-functional “cortical idling” state (Pfurtscheller et al., 1996). This view has recently been challenged, and a more functional interpretation of alpha has been formulated that describes a mechanism of gating-through-inhibition (Klimesch, 1999; Neuper and Pfurtscheller, 2001; Schack and Klimesch, 2002; Jensen and Mazaheri, 2010). According to this view, task-irrelevant regions are inhibited through an increase of alpha oscillations, routing information to task-relevant regions. For instance, it has been demonstrated that alpha activity over visual areas increases in motor tasks and vice versa (Pfurtscheller et al., 1996). A similar mechanism seems to function when attention is directed within the visual or somatosensory domain. For instance, when covert attention is directed to one hemifield (e.g., the left), alpha decreases in the contralateral (right) hemisphere but increases in the ipsilateral (left) hemisphere (Worden et al., 2000; Thut et al., 2006; Rihs et al., 2007; Kelly et al., 2009; van Gerven and Jensen, 2009; Handel et al., 2010). Alpha activity was also shown to decrease in the primary sensorimotor cortex contralateral to the engaged hand while it increased in the ipsilateral hemisphere during a somatosensory working memory task. In line with such a view, we propose that the findings of Yang et al. (2009) and Perry et al. (2010) show how observation of pain in others induces a disinhibition of the somatosensory cortex through alpha suppression. This would create the optimal cortical context in which somatosensory processes such as location and intensity coding of the observed pain (Keysers et al., 2010) can be performed. Such an account might also tentatively explain the increase in alpha reported by Mu et al. (2008). The short (200 ms) presentation of images, together with the task of evaluating these on their painful content, could have resulted in an increase of functional inhibition of the somatosensory cortices for the purpose of reducing interference during the evaluation of the somatosensory (pain) representation. Similar processes have indeed been shown during the retention interval in a visual long term memory task (Meeuwissen et al., in press) as well as during as somatosensory working memory task (Haegens et al., 2011) where distraction in the visual or somatosensory modality was inhibited.

In the current experiment all of the previous concerns were dealt with for the purpose of unequivocally identifying sensorimotor alpha suppression in the observation of pain in others. Subjects viewed images of limbs in pain and no-pain situations in a passive task that required no evaluation or motor responses to the stimuli. To reduce concerns of mixing sources from different regions we recorded brain activity using MEG and applied a beamformer technique for source estimation. We did a time–frequency analysis over the whole post-stimulus interval, but restricted our statistical analysis on the non-evoked period (>400 ms). We hypothesized greater sensorimotor alpha suppression in response to painful images than in response to the observation of control images.

MATERIALS AND METHODS

PARTICIPANTS

Twenty-one healthy participants (15 female, mean age 26.6 years, range: 20–49) enrolled after providing written informed consent and were paid in accordance with guidelines of the local ethics committee (CMO Committee on Research Involving Humans subjects, region Arnhem-Nijmegen, the Netherlands). Two participants were excluded from the analysis due to excessive eye or movement artifacts. One subject fell asleep during the experiment and was excluded as well. The experiment was in compliance with national legislation as well as the code of ethical principles (declaration of Helsinki).

STIMULI

A series of 128 digital color pictures showing right hands and right feet in painful and non-painful situations were used. These stimuli were previously used and validated in behavioral and fMRI studies (Jackson et al., 2005, 2006) and one MEG study (Cheng et al., 2008). All pictures depicted familiar events that can happen in everyday life involving mechanical, thermal, and pressure pain. The neutral pictures involved the same settings without any painful component. All pictures were edited to the same size (600 × 450 pixels).

SUBJECTIVE EMPATHY INDEX

Within 2 weeks before the experiment commenced subjects filled in the Interpersonal Reactivity Index, a self-report questionnaire measuring different factors related to empathy (Davis, 1983).

EXPERIMENT

While seated in the MEG system, the stimuli were projected on a screen about 80 cm in front of the subject. These were all presented in random order for 1.5 s per trial, interleaved with gray fixation screens of 1.5 s (Figure 1). The procedure was repeated over three
blocks resulting in a total experimental time of ~45 min. Subjects were instructed to remain relaxed and not move their limbs, their compliance observed by the experimenter using infrared camera. To make sure subjects paid attention to the stimuli, 10 percent of presentations showed a short twisted movement, created by short (~500 ms) presenting, within one stimulation, the same picture modified with a twirl filter (Photoshop, Adobe Systems Inc.). Subjects were required to internally count the number of these occurrences and report them to the experimenter after each block. Target stimuli were discarded from further analysis. The experiment was programmed and ran using the software package Presentation1.

DATA ACQUISITION
Continuous MEG was recorded using a 275 sensor axial gradiometer system (CTF MEG TM Systems Inc., Port Coquitlam, BC, Canada) placed in a magnetically shielded room. The ongoing MEG signals were low-pass filtered at 300 Hz, digitized at 1200 Hz, and stored for off-line analysis. The subjects’ head position was continuously recorded relative to the gradiometer array using coils positioned at the subject’s nasion and at the left and right ear canals. High-resolution anatomical images (1 mm isometric voxel size) were acquired using a 1.5-T Siemens Magnetom Sonata system (Erlangen, Germany). The same earplugs, using vitamin E instead of the coils, were used for coregistration with the MEG data.

DATA ANALYSIS
Magnetoencephalography data was analyzed using the Matlab-based Fieldtrip toolbox, developed at the Donders Institute for Brain, Cognition and Behavior (Oostenveld et al., 2011). Trials containing movement, muscle, and superconducting quantum interference device (SQUID) jumps were discarded by visual inspection. Independent component analysis (ICA) was used to remove eye and heart artifacts. For the sensor-level analysis, planar gradients of the MEG field distribution were calculated using a nearest-neighbor method comparable with the method described by Bastiaansen and Knosche (2000) and also applied by, e.g., Jokisch and Jensen (2007), Nieuwenhuis et al. (2008), Mazaheri et al. (2009), Haegens et al. (2010), and Haegens et al. (2011). The horizontal and vertical components of the estimated planar gradients approximate the signal measured by planar gradiometers while making the sensor-level data easier to interpret as the maximal activity is typically located above the source (Hamalainen et al., 1993). For source reconstruction, however, we used the original data from the axial sensors.

TIME–FREQUENCY AND ERF ANALYSIS ON THE SENSOR LEVEL
For the time window surrounding the stimulus (~0.4 to 1.5 s), time–frequency representations (TFRs) of power were calculated using a Hanning taper approach applied to short sliding time windows (Percival and Walden, 1993) using an adaptive time window of four cycles length (Δt = 4/f). The data in each time window were multiplied with a Hanning taper. The power values were calculated as the sum of the horizontal and vertical component of the estimated planar gradient after subtracting the mean amplitude from the entire time interval. The planar gradient power estimates were subsequently averaged over trials for the pain and control condition. To investigate the event-related changes in activity we calculated the change of power in response to stimulus presentation relative to the average power during the 200-ms before stimulus-onset.

For the time window surrounding the stimulus (~0.4 to 1.5 s), event-related fields (ERFs) were calculated. The data were then low-pass filtered at 40 Hz using a butterworth filter (order of six), and averaged separately for every condition. Similarly as with the frequency analysis, averaging was done on the planar gradients after which they were summed.

STATISTICAL ANALYSIS ON THE SENSOR LEVEL
To avoid “double dipping” (Kriegeskorte et al., 2009) we restricted our statistical comparison to those sensors where we have previously shown the central mu-rhythm to be maximally modulated in a somatosensory task (Haegens et al., 2011: MLC24, MLC25, MLC31, MLC32, MLP35, MLC42, MLP23, MRC24, MRC25, MRC31, MRC32, MRP35, MRG42, MRG23, see highlighted sensors in Figure 3A); We compared the mean log transformed alpha (7–14 Hz) and beta (15–25 Hz) band power over 1 s post-stimulus period, starting at 400 ms to exclude the contribution of evoked components. Although no differences in the beta-band activity were expected, a clear beta suppression in response to both pain and non-painful stimuli prompted an ad hoc testing for differences between conditions.

SOURCE RECONSTRUCTION
Source reconstruction was performed using a frequency-domain beamformer approach (Dynamic Imaging of Coherent Sources) which uses adaptive spatial filters to localize power in the entire brain (Gross et al., 2001; Liljestrom et al., 2005). The brain volume of each individual subject was discretized to a grid with a 0.8-cm resolution. For every grid point a spatial filter was constructed from the cross-spectral density matrix and the lead field. The lead fields were calculated from a subject specific realistic single-shell model of the brain (Nolte, 2003), based on the individual anatomical MRIs. We calculated the cross-spectral density matrix based upon both the post-stimulus (200–1400 ms) as well as pre-stimulus (1400–200 ms pre-stimulus) interval to obtain the most accurate estimation of the alpha source. Furthermore, both conditions were combined for the purpose of calculating the spatial filter, after which the power at each grid point was estimated for both conditions separately in every subject. Sources were estimated using a multitaper approach to accomplish accurate frequency smoothing for the alpha band (10 ± 2 Hz by using three Slepian tapers). Prior to averaging, the source estimates of the individual subjects’ functional data were spatially normalized using SPM2 to the International Consortium for Brain Mapping template (Montreal Neurological Institute, MNI, Montreal, QC, Canada2).

1http://nbs/neurobs.com

2http://www.bic.mni.mcgill.ca/brainweb
RESULTS

Subjects were presented with static images depicting limbs in painful and non-painful situations from a first-person perspective. We investigated the role of sensorimotor rhythms in processing the painful content.

WIDE-SPREAD MODULATION OF ALPHA- AND BETA-BAND ACTIVITY FOLLOWING VISUAL STIMULI

First, we investigated the responses to combined painful and non-painful stimuli. As shown in Figure 2A, these resulted in marked reductions in occipital alpha (mean 49.9% of baseline, SEM 5.3%), extending to central sensors power (mean 74% of baseline, SEM 4.3%). At central sensors separate alpha and beta components could readily be distinguished (Figures 2B,C). This demonstrates widespread modulation in the alpha and beta frequencies including both occipital and central regions.

GREATER SENSORIMOTOR ALPHA SUPPRESSION IN RESPONSE TO PAINFUL PICTURES

We then tested if these oscillatory responses were different in response to painful compared to non-painful stimuli. As predicted, the strongest modulation was observed at central regions over the head (Figure 3A), showing consistency in topography with a previous investigation of attention-related alpha modulation using the same MEG system and comparable spectral analysis (Haegens et al., 2011). A t-test comparing the average (log transformed) power between pain and control pictures on these sensors (Figure 3B) yielded a significant difference (mean difference 2.19%, SEM = 0.63%, p = 0.019, one-sided) demonstrating that the painful stimuli resulted in stronger sensorimotor alpha suppression than pictures showing non-painful situations. Difference in beta power only showed a trend toward increased suppression (mean difference 1.41%, SEM = 1.07%, p = 0.062, one-sided). Source analysis confirmed that the origin of the alpha difference was located along the central sulcus (Figure 3D).

NO DIFFERENCE IN THE ERF BETWEEN PAINFUL AND CONTROL PICTURES

We also calculated the ERFs to investigate if pain observation could be observed in neuronal activity time-locked to the stimulus. Early visual evoked components (<400 ms) were clearly reflected in the ERFs, while during the interval in which we found sustained alpha suppression (>400 ms) the ERF deflection returned toward baseline. A cluster-based randomization test based upon every timepoint found no difference between conditions for neither occipital nor central sensors (depicted in the boxes of Figure 2A), nor for the central sensors selected for our frequency analysis (Figure 3C). Our findings of increased alpha suppression therefore seem unrelated to differences in evoked responses.

NO CORRELATIONS WITH SUBJECTIVE EMPATHY REPORTS

We also tested for correlations between the magnitude of the modulation of the neuronal response by the pain effect and subjective reports of empathetic distress in daily life. Individual scores on the Interpersonal Reactivity Index (Davis, 1983) did not correlate reliably with the magnitude of the alpha modulation (perspective results).

FIGURE 2 | Oscillatory responses to visual stimuli. (A) The topographic representation of alpha (7–14 Hz) suppression in response to visual stimuli (0.5–1.35 s). (B) Time–frequency representations of central and (C) occipital sensors, respectively, as identified by the boxes in (A). Time–frequency window of interest (0.4–1.35 s; 7–14 Hz) outlined in black.
demands (Klimesch, 1999; Neuper and Pfurtscheller, 2001; Schack and Klimesch, 2002; Jensen and Mazaheri, 2010). Increased alpha suppression in the observation of pain is also in accordance with previous reports of oscillatory involvement in the subjective perception of (first-person) pain intensity (Babiloni et al., 2006) and in the anticipation of pain (Babiloni et al., 2008) and fits well within the general understanding that sensorimotor alpha oscillations provide a dynamically modulated cortical context for somatosensory processing (Lopes da Silva, 1991; Pfurtscheller and Lopes da Silva, 1999; Haegens et al., 2010, 2011). What sensorimotor alpha suppression might represent in terms of underlying empathetic mechanisms remains under debate, however. While activation of sensorimotor regions in fMRI studies can be interpreted in terms of perception–action coupling (Keysers et al., 2010), or of unspecific co-activation (Lamm et al., 2011), our results taken together with the emerging understanding of the functional relevance of alpha oscillations (Lopes da Silva, 1991; Klimesch, 1999; Jensen and Mazaheri, 2010), however strongly suggest a disinhibition of the sensorimotor cortices in response to the observation of pain in others. It is in no way suggested, however, that sensorimotor alpha suppression is by itself sufficient for empathy. As has been argued convincingly at length elsewhere (de Vignemont and Singer, 2006; Keysers et al., 2010; taking: $p = 0.494$, empathetic concern: $p = 0.862$; fantasy: $p = 0.433$; personal distress: $p = 0.248$; total: $p = 0.522$), nor with the magnitude of the beta modulation (perspective taking: $p = 0.667$, empathetic concern: $p = 0.910$; fantasy: $p = 0.829$; personal distress: $p = 0.486$; total: $p = 0.960$).

**Discussion**

We used MEG to investigate neural oscillations in vicarious pain perception. We found wide-spread alpha- and beta-band depression in response to visual stimuli, predominantly at posterior sensors. In contrast to these wide-spread visual responses, observing pain depressed alpha power selectively more at central sensors. By applying source modeling we identified the sources of this difference along the central sulcus, implicating sensorimotor regions in the observation of pain. Strikingly, central and posterior ERPs did not show differences between conditions, suggesting a unique role for induced activity in the brain’s response to observing other people in pain. These results provide support for the involvement of sensorimotor oscillations in empathetic responses.

Alpha oscillations might be providing a graded level of excitability and inhibition in task relevant and irrelevant regions, streamlining information flow dependent on moment-by-moment task demands (Klimesch, 1999; Neuper and Pfurtscheller, 2001; Schack and Klimesch, 2002; Jensen and Mazaheri, 2010). Increased alpha suppression in the observation of pain is also in accordance with previous reports of oscillatory involvement in the subjective perception of (first-person) pain intensity (Babiloni et al., 2006) and in the anticipation of pain (Babiloni et al., 2008) and fits well within the general understanding that sensorimotor alpha oscillations provide a dynamically modulated cortical context for somatosensory processing (Lopes da Silva, 1991; Pfurtscheller and Lopes da Silva, 1999; Haegens et al., 2010, 2011). What sensorimotor alpha suppression might represent in terms of underlying empathetic mechanisms remains under debate, however. While activation of sensorimotor regions in fMRI studies can be interpreted in terms of perception–action coupling (Keysers et al., 2010), or of unspecific co-activation (Lamm et al., 2011), our results taken together with the emerging understanding of the functional relevance of alpha oscillations (Lopes da Silva, 1991; Klimesch, 1999; Jensen and Mazaheri, 2010), however strongly suggest a disinhibition of the sensorimotor cortices in response to the observation of pain in others. It is in no way suggested, however, that sensorimotor alpha suppression is by itself sufficient for empathy. As has been argued convincingly at length elsewhere (de Vignemont and Singer, 2006; Keysers et al., 2010;
Lamm et al., 2011), empathy is a multidimensional response to a variety of situations, real and imagined. Not one neural mechanism will be able to explain its full expression. Noteworthy are the results by Betti et al. (2009) who found increased gamma band coherence between primary sensory and motor regions when subjects watched painful movies. Their results further emphasize the importance of understanding empathy through dynamic oscillatory interactions of neuronal assemblies distributed within and across different specialized brain regions. Neither is it claimed here that sensorimotor alpha oscillations are specifically involved in social processes. We understand alpha oscillations more generally as a (sensory) mechanism for gating-through-inhibition, which, in this particular case, is involved in the task of disinhibiting sensory cortices in response to the observation of pain in others.

Although we were able to localize the source of the empathetic alpha modulation along the central sulcus, we were not able to distinguish between primary somatosensory from primary motor regions. This might have been a problem of lack of signal strength, compared to those studies that have been able to localized alpha modulation to postcentral sulcus in response to actual tactile stimulation. For instance, in Caetano et al. (2007) observation of movement resulted in only 42% of the beta suppression that was found in response to actual movement. As we localized the difference between the pain and no-pain condition, a difference of only a couple of percent, this problem might have been exacerbated even further.

The fact that this study is limited to the observation of limbs might limit a generalization to the observation of other body parts that afford less action related responses. In fact, as mentioned before, besides the fact that sensorimotor alpha is clearly implicated in somatosensory attention and perception, it has also been a hallmark of action–perception. In our study the amount of action suggested in the images was, however, kept similar between conditions. For instance, where in the control condition one situation depicts the sawing through of a wooden log, the painful counterpart differed only by the fact that the second hand was in a painful position underneath the saw. Any difference between the observation of painful and control images therefore cannot be explained by a difference in action–perception between these conditions. This fact, together with the fact that alpha oscillations are commonly found to have a postcentral source (Salmelin and Hari, 1994; Cheyne et al., 2003; Jurkiewicz et al., 2006) makes an interpretation of our findings in terms of motor processes less likely.

Although subjects were instructed to remain relaxed and not move their limbs, and no responses were required, this cannot fully rule out “covert movement” or muscle tension in response to the painful images. Although some (e.g., Yang et al., 2009, but not Perry et al., 2010) record electromyography (EMG), a null-finding comparing average EMG across conditions will not be able to fully rule out differences below the sensitivity of the measurement, and any selection of muscles would have its limitations. More importantly, we did not find significant differences in the beta (≈20 Hz) band where effects of movement would be most pronounced (Salmelin et al., 1995; Hari and Salmelin, 1997; Cheyne et al., 2003; Jurkiewicz et al., 2006; Parkes et al., 2006).

We did not find any correlation with self-report questionnaires and contribute to the existing confusion in the literature. In Fan and Han (2008), self-reported unpleasantness of the observed pain stimuli correlated with early evoked potentials to painful observation, but did so only in an evaluation task and not in the counting task, which would be more similar to our passive paradigm. Mu et al. (2008) reported negative correlations for both perceived pain and self-unpleasantness, also in a similar evaluation task. Conversely, Cheng et al. (2008) found only positive correlations of perspective taking with sensory-evoked alpha rebound, and no correlation with empathy quotient questionnaire (EQ: Baron-Cohen and Wheelwright, 2004, the emotional contagion scale, ECS: Doherty, 1997) or any subscales of the interpersonal reaction index (IRI: Davis, 1983). Finally, Yang et al. (2009) found that the effect of pain on the alpha rebound correlated with the personal distress subscale of the IRI, but only in females. Taken together, it seems that the most consistent findings have been found correlating the subjective quality of individual stimuli trials with oscillatory activity, while correlating individual traits have been less conclusive.

Concluding, observing limbs in pain suppressed ongoing alpha oscillations more than observing control images. This effect was localized along the central sulcus, implicating somatosensory alpha oscillations in the observation of pain in others. These findings support an interpretation in terms of gating-through-inhibition where observation of painful images disinhibits the somatosensory cortex through alpha suppression.

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