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Independent effects of eye gaze and spatial attention on the processing of tactile events: Evidence from event-related potentials

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Abstract

Directing one’s gaze at a body part reduces detection speed and enhances the processing of tactile stimuli presented at the gazed location. Given the close links between spatial attention and the oculomotor system it is possible that these gaze-dependent modulations of touch are mediated by attentional mechanisms. To investigate this possibility, gaze direction and sustained tactile attention were orthogonally manipulated in the present study. Participants covertly attended to one hand to perform a tactile target-nontarget discrimination while they gazed at the same or opposite hand. Spatial attention resulted in enhancements of the somatosensory P100 and Nd components. In contrast, gaze resulted in modulations of the N140 component with more positive ERPs for gazed than non gazed stimuli. This dissociation in the pattern and timing of the effects of gaze and attention on somatosensory processing reveals that gaze and attention have independent effects on touch.

Keywords:
Spatial attention – Eye gaze - Somatosensory processing  - Event-related brain potentials
Highlights

- Is the effect of gaze on touch mediated by attentional mechanisms?
- Previous studies manipulated gaze direction but not spatial attention
- Here, both gaze and spatial attention are simultaneously manipulated
- SEPs modulations by gaze and attention have different patterns and time courses
- The effects of gaze and attention on touch are independent
Introduction

When we feel a touch at a certain location on our body, we tend to direct our eyes to that location to look at the source of stimulation. While this orienting behavior does not alter directly the tactile input, that is the operations of the mechanoreceptors on the stimulated skin, a number of recent studies have now demonstrated that tactile processing is modulated not only by the availability of visual information about the stimulated body part but also by the direction of the eyes.

Viewing the touched body part during a tactile task improved the discrimination of stimuli and lowered the tactile threshold (e.g. Kennett et al., 2001; Press et al., 2004; Tipper et al., 1998; 2001). For instance, responses to tactile targets were faster when presented to the visible hand (displayed on a monitor thorough a video camera) than when they were presented to the non-visible hand, suggesting that vision of the hand facilitated the discrimination of tactile stimuli (Tipper et al., 1998). In addition, psychophysical studies showed improved performance in a two-point discrimination threshold task (2ptD) when participants viewed their stimulated arm, as compared to when their arm was not visible or when a neutral object was presented in the same location (Kennett et al., 2001; Press et al., 2004). Crucially, in these experiments, visual enhancement of touch was observed despite the fact that vision of the tactually stimulated body site was completely non-informative (i.e. viewing the body did not provide any information about the tactile stimulation). Neuroimaging studies have started to unravel the neural mechanisms underlying this facilitatory effect of vision on touch. For instance, TMS and ERP evidence showed that non-informative vision can modulate early somatosensory processing, already within the primary somatosensory cortex, and that the multisensory integration of visual and tactile information is likely to be responsible for the visual enhancement of touch (e.g. Cardini et al., 2011; 2012; Forster and Eimer, 2005; Fiorio and Haggard, 2005; Longo et al., 2011; Taylor-Clarke et al., 2002).

Typically, when we look at a specific body site we tend to direct our eyes towards the relevant body location, to foveate the source of the relevant visual information. Thus, very often the possibility to further process visual information from a specific body site is
mediated by changes in the position of the eyes (gaze direction). Intriguingly, it has been observed that also gaze direction can modulate tactile perception even when no visual information relative to the stimulated body site is available (Honoré et al., 1989; Pierson et al., 1991; Tipper et al., 1998). While the effect of eye gaze on touch has been less investigated, initial evidence suggests that gazing towards a specific body location facilitates the processing of tactile stimuli presented at that location. Behavioural studies in which participants were asked to detect or discriminate tactile stimuli presented to either hands while their gaze was directed to one of the hands, showed faster responses to stimuli presented to the hand gaze was directed, under conditions where the hands were hidden from view (Honoré et al., 1989; Pierson et al., 1991; Tipper et al., 1998).

The studies described above suggest that gazing to the tactilely stimulated body part can improve tactile processing and that this effect is independent from the presence of visual information. While the available evidence indicates that vision and gaze have independent effects on touch (e.g. Forster and Eimer, 2005; Tipper et al., 1998), the functional and neural mechanisms that mediate the effect of gaze on touch remain almost entirely unknown. The proprioceptive orienting of the eyes towards the tactilely stimulated body site has been suggested as one of the possible mechanisms responsible for the observed changes in tactile processing due to the manipulation of gaze direction (Tipper et al., 2001; 1998; Honoré et al., 1898). In addition, it has been proposed that the effect of gaze on touch might be mediated by spatial attention (Forster and Eimer, 2005; Pierson et al., 1991). Given that the mechanisms responsible for the allocation of attention in space are closely linked to the oculomotor system, spatial attention might be automatically directed to the gazed body location. While these two hypotheses are not mutually exclusive, none of them has been directly investigated.

Indirect support for a role of attention in the effect of gaze on touch comes from the only ERP study to date reporting modulations of somatosensory processing by gaze direction (Forster and Eimer, 2005, Exp. 3; but see also Hesse et al., 2004, for null effects of gaze on touch). In this study, participants were instructed to gaze to one hand while responding to infrequent tactile targets randomly presented to either hands (both hands were hidden under a table top). ERPs elicited by tactile stimuli presented to the gazed hand as compared to stimuli presented to the non gazed hand were characterized by
enhanced N140 components followed by enhanced sustained negativities. Importantly, the pattern and time course of this effect of gaze (Forster and Eimer, 2005) are remarkably similar to those reported in previous ERP studies of covert tactile spatial attention (e.g. Eimer and Forster, 2003; Forster and Eimer, 2004; García-Larrea et al., 1995; Michie et al., 1987). When participants are explicitly instructed to covertly attend to one of their hands (and to maintain their gaze onto a ‘neutral’ central location equidistant to both hands), enhanced mid-latency somatosensory ERP components (P100 and/or N140), followed by enhanced Nd components (Eimer and Forster, 2003; Michie, 1984; Josiassen et al., 1982) are typically elicited by attended stimuli.

The similarities between the effects of gaze (Forster and Eimer, 2005) and of spatial attention (e.g. Eimer and Forster, 2003; Forster and Eimer, 2004; García-Larrea et al., 1995; Michie et al., 1987) on somatosensory processing might suggest that manipulating gaze direction activates the same mechanisms that are responsible for the covert orienting of spatial attention in touch. However, existing data cannot provide unequivocal support for the hypothesis of a functional link between gaze and spatial attention. In all previous studies on the effect of gaze on touch (Honoré et al., 1989; Tipper et al., 1998; Forster and Eimer, 2005; Hesse et al., 2004; Pierson et al., 1991) tactile stimuli were equally likely to be presented to either hands, and participants had no incentive to focus their attention on a specific location. In other words, spatial attention was not directly manipulated. Under these experimental conditions it is possible that spatial attention was directed to the gazed location simply because it was not engaged in any other specific spatial task. Thus, current evidence does not allow to disentangle between the effects of gaze and of spatial attention on touch.

To directly investigate whether the effects of gaze and spatial attention on touch are mediated by the same mechanisms or whether these two effects are independent, both gaze direction and spatial attention were simultaneously and independently manipulated in the present ERP study. Participants were instructed to focus and maintain their covert attention on the task-relevant (attended) hand throughout a block of trials to carry out a difficult discrimination between target and non-target tactile stimuli presented to that hand, while ignoring all tactile stimuli presented to the other task-irrelevant (unattended) hand. In addition, they had to direct and maintain their gaze on the task-relevant or on the
task-irrelevant hand in different blocks of trials. Somatosensory ERPs in response to tactile non-target stimuli presented to the left and right hand were compared as a function of attention (stimulus presented to the attended – task-relevant -, A+, vs. unattended – task-irrelevant - hand, A-) and gaze (stimulus presented to the hand gaze was directed, G+, vs. the opposite hand, G-). Because gaze and attention were orthogonally manipulated so that each condition of attention (A+ vs. A-) could be matched to any condition of gaze (G+ vs. G-), it was possible to investigate the electrophysiological correlates of their effects on tactile processing. If the effect of gaze and spatial attention are mediated by the same neural mechanisms, gaze-dependent and attention-dependent ERP modulations should have similar profiles and influence analogous stages of somatosensory processing. In contrast, if the effect of gaze on touch is mediated by mechanisms that are independent of spatial attention, different patterns of SEPs modulations by gaze and by spatial attention are likely to be observed. Note that if previously reported effects of gaze on touch were confounded with (and not mediated by) spatial attention, there is the possibility that no effect of gaze is observed when the direction gaze and spatial attention are orthogonally manipulated.
Material and methods

Participants

Nine paid volunteers (2 males) aged 21-35 (mean age of 26.9 years) participated in the experiment. Two were left handed and they all had normal or corrected-to-normal vision by self-report. All participants gave written informed consent. The study was performed in accordance with the ethical standards laid down in the 1964 Declaration of Helsinki and was approved by the Ethics committee, Department of Psychology, City University London.

Stimuli and apparatus

Participants sat in a dimly lit experimental chamber. Tactile stimuli were presented using a 12V solenoids, driving a metal rod with a blunt conical tip to the top segment of the index fingers, making contact with the fingers whenever a current was passed through the solenoid. Two tactile stimulators were used, each attached with adhesive medical tape to the left and right index finger, placed so that the metal rod made contact with the outer side of the top phalanx.

Tactile stimuli were either continuous (non-target stimuli), consisting of one rod contacting one finger for 200 ms, or contained a 6-ms gap where this contact was interrupted after a duration of 97 ms (gap stimuli). Throughout the experimental blocks, white noise (62 dB SPL) was continuously delivered from a loudspeaker centrally located in front of the participants, to mask any sounds made by tactile stimulators.

Participants were instructed to place their hands palm side down on a table with their left and right index finger positioned 20 cm from the left and the right of the body midline. A black cardboard panel (69 x 41 cm) was placed on the table and was used to prevent the visibility of the hands and lower parts of the arms. Two markers (white circles, 0.2 cm diameter) were located on this panel 40 cm apart (20 cm to the right or left of the panel centre) and were used as fixation points when participants were instructed to gaze to their left or right hand. To ensure the accurate alignment of unseen hands and
gaze direction throughout all experimental blocks, participants had to hold a small stick between their middle and the index finger with each hand. These sticks passed through two small holes practiced on the horizontal panel so that their top segment was visible. The white circles used as fixation points were glued to the top of these sticks. Participants’ body posture was unchanged through the experiment, with the exception of their gaze that was maintained to the left or right fixation point (corresponding with the left and right hand, respectively) in different blocks.

**Procedure**

Each trial started with a 200 ms stimulus presentation (either target or non-target) followed by a 1000 ms interval used to collect vocal responses. Intertrial interval was varied randomly between 200 and 300 ms.

The experiment consisted of 12 blocks, with 64 trials per block. In each block, a non-target stimulus was presented on 48 trials with equal probability to the task-relevant or to the task-irrelevant hand. A target stimulus was presented in the remaining 16 trials. Of these, twelve were trials where a target stimulus was presented to the task-relevant hand, while four were trials where a target stimulus was presented to the task-irrelevant hand. Participants were instructed to respond vocally (by saying ‘yes’) whenever a target stimulus was presented to the task-relevant hand, they had to ignore target stimuli to the task-irrelevant hand as well as all non-target stimuli. At the beginning of the session, a block of trials was run to familiarize participants with the task and the stimuli.

On six out of twelve blocks they had to gaze to the task-relevant (attended) hand, while on the other half they had to gaze to the opposite task-irrelevant (unattended) hand. The order of task-relevant hand (left vs. right hand) and gaze direction (left vs. right hand) in which these blocks were delivered was counterbalanced across participants. Overall three successive blocks of the same task were run for each combination of task-relevant hand (attend to the left hand vs. attend to the right hand) and gaze-direction (left hand vs. right hand). Prior to the beginning of each block, participants were instructed/reminded about the task-relevant hand (left or right hand) and about the gaze direction (left or right hand) they had to maintain throughout the block.
Participants’ gaze direction was monitored via a video camera throughout the experiment.

**G recording and data analyses**

EEG was recorded from 28 Ag–AgCl electrodes (Fp1, Fp2, F7, F8, F3, F4, Fz, Fc5, Fc6, Fc1, Fc2, Fcz, T7, T8, C3, C4, Cz, Cp5, Cp6, Cp1, Cp2, P7, P8, P3, P4, Pz, O1, O2) relative to a right earlobe reference. Horizontal EOG was recorded unipolarly from the outer canthi of both eyes. Electrode impedance was kept below 5 kΩ, and efforts were made to equalize the impedance of the earlobe electrodes. Amplifier bandpass was 0.1–100 Hz, and digitization rate was 500 Hz. EEG was digitally re-referenced to the average of the left and right earlobes and HEOG was averaged for the left and right eye. Trials with eye blinks (Fp1 or Fp2 exceeding ±60 µV relative to baseline), horizontal eye movements (HEOG exceeding ±30 µV relative to baseline) or other artefacts (a voltage exceeding ±60 µV at any other electrode location relative to baseline) were excluded. On average, these artefact rejection criteria led to the exclusion of 6.5% of trials.

ERPs to non-target stimuli were averaged relative to a 100 ms pre-stimulus baseline for 300ms after stimulus onset, separately for all combinations of attended hand (left versus right), gaze direction (left versus right) and stimulus location (left versus right). ERP mean amplitudes were computed within measurement windows centered on the latency of somatosensory P45 (40-60ms), N80 (70-90ms), P100 (90-120ms) and N140 (130-150 ms) components. To investigate longer-latency effects of gaze and attention, mean amplitudes were also computed between 160-240 ms post-stimulus. Analyses of somatosensory ERPs were conducted separately for lateral anterior (F7/8, F3/4, and FC5/6), lateral central (FC1/2, C3/4, and CP1/2) and lateral posterior (P7/8, P3/4, and CP5/6) sites, contralateral and ipsilateral to the stimulated hand, and for midline sites (Fcz, Cz, and Pz).

To investigate the effects of gaze and attention on somatosensory processing, separate repeated measures analyses of variance were carried out for each time window (40-60ms; 70-90ms; 90-120ms;130-150ms; 160-240ms) and each electrode site (lateral anterior, central and posterior sites, contralateral and ipsilateral to the stimulated hand,
and midline sites) with gaze (stimulus presented to the gazed hand versus the non gazed hand: G+ vs. G-), attention (stimulus presented to the attended versus unattended hand: A+ vs. A-), stimulus location (left versus right), and electrode site (F7/8, F3/4, and FC5/6 for lateral anterior electrodes; FC1/2, C3/4, and CP1/2, for central electrodes; P7/8, P3/4, and CP5/6 for lateral posterior electrodes and Fcz, Cz, and Pz for midline electrodes) as within-subjects factors.

When appropriate, Greenhouse-Geisser adjustments to the degrees of freedom were performed. For all ERP analyses, only significant main effects or interactions are reported.

Participants were instructed to vocally respond to attended targets only. Thus, they had to discriminate between targets and nontargets presented to the attended hand, while ignoring all stimuli to the unattended hand. Error rates were computed and reported separately for the different types of trials (false alarms to unattended targets, false alarms to nontargets and failure to respond attended targets). In addition, accuracy rates were calculated across all types of trials as a function of gaze (stimulus presented to the gazed, G+, vs. non gazed, G-, hand) and compared with paired t-tests. The latency of vocal responses was measured with a voice key relative to the gap onset of the target stimuli (97 ms after stimulus onset), as target/non-target discriminations were only possible after this interval. Only correct vocal responses between 200 and 900 ms post stimulus were used to compute the mean RTs for the response analysis. To investigate the effect of gaze on participants’ response speed, mean RTs were calculated separately for gazed (G+) and non gazed (G-) attended targets and were compared with paired t-tests.

Results

Behavioural results

False alarms to nontargets occurred on less than 0.2% of these trials (0.06% on G+ and 0.51% on G- trials). No false alarm occurred when targets were presented to the unattended hand. Participants failed to respond to attended targets on 2.62% of these trials. When overall accuracy rates (calculated across all trials) were compared as a
function of gaze, higher accuracy in the G+ as compared to the G- condition was observed, 99.5% and 99.1%, respectively, t(8)=2.7, p=.026. The analysis of correct vocal RTs to attended targets presented to the gazed and non gazed hand did not show any statistically reliable difference between G+ and G- conditions, 475 and 483 ms, respectively; t(8)=-1.5, p=.16).

EEG results

Figures 1 and 2 show SEPs elicited by non-target stimuli in the 300 ms interval after stimulus onset. SEPs are displayed separately for midline sites Fcz, Cz and Pz (centre) and for lateral sites F3/4, C3/4 and P3/4 ipsilateral (right) and contralateral (left) to the stimulated hand. Figure 1 shows the effect of Gaze on tactile processing, obtained by comparing ERP waveforms elicited by non-target stimuli delivered to the hand to which gaze was directed (G+, solid line) and to the other non gazed hand (G-, dashed line), collapsed across currently attended and unattended stimuli (A+ and A-, respectively). The effect of sustained spatial attention on touch is represented in Figure 2 where ERP waveforms are shown separately for tactile non-target stimuli to the attended hand (A+, solid line) and to the unattended hand (A-, dashed line), collapsed across the current direction of gaze (G+ and G- trials). The corresponding difference waveforms for the effects of gaze and attention can be observed in Figure 3 (solid and dashed lines, respectively), while Figure 4 shows separately the four different experimental conditions (A+G+; A-G-; A+G- and A-G+) as observed at central electrodes C3/4 contralateral to the stimulated hand.

While both gaze and attention appeared to modulate the processing of tactile events, their effects seem to be systematically different and to modulate different time windows in which distinct somatosensory ERP components were elicited. The manipulation of gaze resulted in selective modulations of ERPs elicited in the N140 time window. As shown in Figure 1, reduced N140 component were observed for gazed stimuli (G+ trials) as compared to non gazed stimuli (G-), that is, ERPs elicited by stimuli to the gazed hand (G+) were more positive than those to the non gazed hand (G-). In contrast, sustained spatial attention resulted in enhanced P100 components for attended stimuli (A+) as
compared to unattended ones (A-) (Figure 2). While no effect of spatial attention was visible in the N140 time range, enhanced negativities for tactile stimuli to the attended as compared to the unattended hand were visible in the time range of the processing negativity (Nd, Figure 2).

**Figure 1.** Grand-averaged somatosensory ERPs and horizontal EOG waveforms elicited by non-target tactile stimuli delivered to the gazed (G+, solid lines) or non gazed hand (G-, dashed line) in the 300 ms following stimulus onset (relative to a 100 ms pre-stimulus baseline) at fronto-central (FC3/4), central (C3/4) and parietal (P3/4) electrodes contralateral (c) and ipsilateral (i) to the stimulated hand as well as at midline electrodes (Fcz, Cz, Pz).
**Figure 2.** Grand-averaged somatosensory ERPs elicited by non-target tactile stimuli delivered to the attended (A+, solid lines) or unattended hand (A-, dashed line) in the 300 ms following stimulus onset (relative to a 100 ms pre-stimulus baseline) at fronto-central (FC3/4), central (C3/4) and parietal (P3/4) electrodes contralateral (c) and ipsilateral (i) to the stimulated hand as well as at midline electrodes (Fcz, Cz, Pz).
Figure 3. Difference waveforms obtained at contralateral (c), midline and ipsilateral (i) electrodes by subtracting ERPs elicited by tactile non-target stimuli presented to the non gazed hand (G-) from those elicited by stimuli to the gazed hand (G+) (solid lines), and by subtracting ERPs elicited by tactile non-target stimuli presented to the unattended hand (A-) from those elicited by stimuli to the attended hand (A+) (dashed lines).
No significant effect of gaze emerged in the P45 (40-60 ms post stimulus), N80 (70-90 ms), or P100 (90-120 ms) time ranges. In contrast, an enhanced positivity for G+ as compared to G- trials was reliably present in the N140 time window between 130 and 150 ms post-stimulus. In this time window, significant main effects of gaze were found at anterior and central electrodes ipsilateral and contralateral to the stimulated hand as well as at contralateral posterior and midline electrodes (all F(1, 8)>6.04; all p<.04). Gaze x Electrode Site interactions were present at anterior electrodes both ipsilateral and contralateral to the stimulated hand (both F(2, 16)>5.67; both p<.022), and follow-up analyses showed a significant effect of gaze on N140 amplitudes at all sites (all F(1, 8)>5.3; all p<.05) with the exception of contralateral F7/8 where this effect approached significance (F(1, 8)=4.3; p=.72). In the subsequent time window (160-240 ms post stimulus) no main effects of gaze were observed at any of the electrode sites. Table 1 summarizes the mean amplitude values and mean standard errors of tactile ERPs elicited by gazed (G+) and non gazed (G-) stimuli. These values are reported for the specific time windows and electrodes sites where statistically reliable main effects of gaze were observed.

<table>
<thead>
<tr>
<th>N140</th>
<th>G+</th>
<th>Central</th>
<th>Posterior</th>
<th>Anterior</th>
<th>Central</th>
<th>Posterior</th>
<th>Midline</th>
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<tbody>
<tr>
<td></td>
<td>-1.35 (0.42)</td>
<td>-0.059 (0.5)</td>
<td>0.1 (0.5)</td>
<td>0.18 (0.53)</td>
<td>0.7 (0.59)</td>
<td>-</td>
<td>0.56 (0.63)</td>
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<tr>
<td>130-150 ms</td>
<td>G-</td>
<td>-1.9 (0.46)</td>
<td>-0.9 (0.48)</td>
<td>-0.67 (0.43)</td>
<td>-0.49 (0.53)</td>
<td>-0.02 (0.55)</td>
<td>-</td>
</tr>
</tbody>
</table>

Table 1. Mean amplitude values (µV) and mean standard errors of ERPs elicited by gazed (G+) and non gazed (G-) tactile stimuli for the time windows and electrode sites in which significant main effects of gaze emerged.

Reliable effects of attention were first observed in the P100 time range (90-120 ms post-stimulus), with enhanced P100 amplitudes for A+ relative to A- stimuli, as indicated by main effects of attention at central electrodes contralateral and ipsilateral to the stimulated hand as well as at contralateral anterior and midline electrodes (all F(1,8)>6.3;
all p<.036). In contrast, no significant modulation of the N140 component was observed between 130 and 150 ms post stimulus. Finally, in the 160-240 ms time window, enhanced negativities for A+ as compared to A- trials were observed at central and posterior electrodes contralateral and ipsilateral to the stimulated hand as well as midline sites (all F(1,8)>7.08; all p<.03). Attention x Electrode Site interactions were observed in the Nd time range for posterior electrodes both contralateral and ipsilateral to the stimulated hand (both F(2, 16)>4.6; both p<.026) as well as for midline sites (F(2, 16)=11.6; p=.005), and follow-up analyses revealed significant main effects of attention at all sites (all F(1, 8)>6.7; all p<.032), except for ipsilateral P7/8 were this effect did not reach statistical significance (F(1, 8)=3; p=.12). Table 2 summarizes the mean amplitude values and mean standard errors of tactile ERPs elicited by attended (A+) and unattended (A-) stimuli. These values are reported for the specific time windows and electrodes sites where statistically reliable main effects of attention were observed.

<table>
<thead>
<tr>
<th></th>
<th>Attention</th>
<th>Contralateral electrodes</th>
<th>Ipsilateral electrodes</th>
<th>Midline</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>A+</td>
<td>Anterior</td>
<td>Central</td>
<td>Posterior</td>
</tr>
<tr>
<td>P100 (90-120ms)</td>
<td></td>
<td>0.59 (0.38)</td>
<td>1.6 (0.47)</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>A-</td>
<td>0.1 (0.45)</td>
<td>1 (0.54)</td>
<td>-</td>
</tr>
<tr>
<td>Nd (160-240ms)</td>
<td>A+</td>
<td>-</td>
<td>0.38 (0.68)</td>
<td>0.1 (0.62)</td>
</tr>
<tr>
<td></td>
<td>A-</td>
<td>-</td>
<td>2.35 (0.58)</td>
<td>1.4 (0.51)</td>
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</table>

Table 2. Mean amplitude values (µV) and mean standard errors of ERPs elicited by attended (A+) and unattended (A-) tactile stimuli for the time windows and electrode sites where significant main effects of attention were found.

No Gaze x Attention interaction emerged in any of the time window at any of the electrode sites. In the N140 time window, a significant Gaze x Attention x Electrode Site interaction (F(2, 16)=3.89; p=.046) was observed for anterior electrodes ipsilateral to the attended hand. To further explore this interaction, follow up analyses were conducted separately for single anterior electrode site, but no significant Gaze x Attention
interactions were present at any of the electrode sites. Figure 4 shows separately ERPs elicited by the four different experimental conditions. The main difference between ERP waveforms is primarily due to attention in the P100 and Nd time windows (A+ and A-, solid vs. dashed lines, respectively), while it is mainly driven by gaze in the N140 interval (G+ and G-, black vs. grey lines).

Figure 4. Grand-averaged somatosensory ERPs elicited by non-target tactile stimuli delivered to the attended and gazed (A+G+), attended and non gazed (A+G-), unattended and non gazed (A-G-) or unattended and gazed (A-G+) hand in the 300 ms following stimulus onset (relative to a 100 ms pre-stimulus baseline) at C3/4 electrodes contralateral to the stimulated hand. ERPs elicited by attended and unattended stimuli are represented by solid and dashed lines, respectively. ERPs elicited by gazed and non gazed stimuli are represented by black and grey lines, respectively.
General discussion

Directing gaze to a body site improves the discrimination of tactile stimuli presented to that location, as demonstrated by initial behavioural and electrophysiological evidence (Forster and Eimer, 2005; Honoré et al., 1989; Pierson et al., 1991; Tipper et al., 1998). However, the neural mechanisms underpinning the effect of gaze on touch remain almost completely unexplored. Here, we directly investigated the relationship between the effects of gaze and spatial attention on tactile processing, to uncover whether these effects are mediated by shared neural mechanisms or whether, in contrast, they represent the outcome of distinct processes. To this aim we orthogonally manipulated the directions of gaze and spatial attention, by instructing participants to covertly attend one of their hands, while directing their gaze to the same hand or to the opposite unattended hand (in different blocks of trials). Because participants’ hands were covered during the experiment, attention and gaze were directed towards the location occupied by the hands. Thus, the effects of gaze and attention on touch were investigated in the absence of visual information about the tactually stimulated body part.

The analysis of ERPs elicited by tactile stimuli as a function of whether they were presented to the attended (A+) or unattended hand (A-) demonstrated reliable effects of spatial attention. These are shown in Figure 2 where ERPs elicited by attended (A+, solid line) and unattended (A-, dashed line) stimuli are represented regardless of gaze (i.e. collapsed across G+ and G-). In line with previous studies on sustained attention in touch (c.f. Desmet and Robertson, 1977; Eimer and Forster, 2003; García-Larrea et al., 1995; Michie et al., 1987; Zopf et al., 2004), the spatial selection of tactile information affected not only mid-latency components, but also post-perceptual processes related to the identification and categorization of tactile stimuli (see Sambo and Forster, 2011, for review). More specifically, we observed enhanced P100 components (90 – 120 ms post-stimulus) for ERPs elicited by attended as compared to unattended stimuli. These attentional effects were maximal over central electrodes ipsilateral to the stimulated hand, but were also reliably present over contralateral central and anterior electrodes as well as at midline electrodes, as shown by the scalp distribution visible in Figure 5 (right column, top panel). While no effects of attention were observed in the subsequent N140 time
window (130-150 ms post-stimulus onset), later attentional ERP modulations were present between 160 and 240 ms post- stimulus overlapping with the descending flank of the N140 component and with the subsequent processing negativity. These enhanced negativities for stimuli presented to the attended as compared to the unattended hand were present over central and posterior electrodes and maximal over midline sites (Figure 4, right column, bottom panel). While a number of different factors contribute to determine the specific time course of the attentional modulations of tactile ERPs (such as the specific type of attention task, e.g. Forster and Eimer, 2003; the availability of visual information relative to the stimulated body site, e.g. Sambo et al., 2009; the difficulty of the discrimination task (e.g. Michie et al.,1987) and the stimulated body site (e.g. Gillmeister et al., 2010), the observation that tactile spatial attention resulted in modulations of both perceptual (here reflected by P100 enhancements) and post-perceptual (as reflected by Nd modulations) stages of somatosensory processing is in line with previous ERP studies (e.g. Sambo and Forster, 2011) and demonstrates that participants engaged with the tactile attention task and covertly attended the task-relevant hand\(^1\).

Crucially, ERPs elicited by tactile stimuli presented to the hands were systematically modulated not only by the direction of covert attention but also by the direction of gaze. A reliable difference between ERPs elicited by gazed (G+) and non gazed (G-) tactile stimuli was observed between 130 and 150 ms post-stimulus, overlapping with the N140 somatosensory ERP component. More specifically, reduced N140 components were found for G+ as compared to G- stimuli, that is, ERPs elicited by stimuli to the gazed hand (G+) were more positive than those to the non gazed hand (G-). The effect of gaze on touch is shown in Figure 1 where somatosensory ERPs elicited by non-target stimuli are represented as a function of whether they were presented to the gazed hand (G+, solid line) or to the opposite non gazed hand (G-, dashed line), regardless of spatial attention (i.e. collapsed across A+ and A-). The scalp distribution of the effect of gaze is visible in Figure 5 (left column, middle panel) and shows that the enhanced positivity for G+ as compared to G- in the N140 time window was not restricted to central electrodes but was also present over anterior, posterior and midline sites. The effect of gaze on touch was not only present in the ERP data, but also in the
behavioural data. While no modulation of response speed was observed, higher accuracy levels were found on trials in which the tactile stimulus was presented to the gazed hand as compared to trials in which it was presented to the non-gazed hand.

![Effect of Gaze and Attention](image)

- **P100** (50-120 ms)
- **N140** (130-150 ms)
- **N200** (160-240 ms)
The aim of the present study was to disentangle the effect of gaze direction on touch from that of spatial attention. First, we asked whether an effect of gaze on touch can be observed when both spatial attention and gaze direction are independently manipulated. Results were clear-cut. Under these experimental conditions, gaze direction still resulted in systematic modulations of somatosensory processing. This finding confirms the presence of an effect of gaze on touch, expanding results of earlier investigations (Forster and Eimer, 2005; Honoré et al., 1989; Pierson et al., 1991; Tipper et al., 1998). Second, we asked whether shared mechanisms are responsible for both the effects of gaze and spatial attention on touch. To address this question we directly compared gaze-dependent and attention-dependent modulations of tactile processing. Both the pattern and the specific time course of these SEPs modulations indicate that the mechanisms underlying the effect of gaze are independent of those responsible for spatial attention.

Typically, attentional modulations of sensory-specific somatosensory ERP components are characterized by enhanced amplitudes for ERPs elicited by stimuli presented to the attended versus the unattended hand (e.g. Desmet and Robertson, 1977; Eimer and Forster, 2003; García-Larrea et al., 1995; Michie et al., 1987). In the present study, we observed reliable effects of gaze on somatosensory processing in the N140 time window. If spatial attention was at least in part responsible for the effect of gaze on touch observed in the N140 time window, increased N140 components for ERPs elicited by stimuli presented to the gazed hand should have been observed. In contrast, gaze-dependent modulations of somatosensory processing resulted in reduced N140
components for gazed stimuli (that is ERPs were more positive for G+ than for G-
between 130 and 150 ms post stimulus). This finding provides the first direct indication
that gazing to the tactually stimulated body site activates neural mechanisms which are
different from those of spatial attention.

This conclusion is further supported by the specific time course of the effects of
gaze and attention. While both gaze and spatial attention had reliable effects on touch,
they influenced different stages of somatosensory processing. Results demonstrated a
clear dissociation between the timing and therefore the associated locus of the effects of
gaze and attention on touch. Sustained spatial attention resulted in enhanced positivities
starting around 90 ms after stimulus onset. In contrast, the earliest effects of gaze were
only observed after 130 ms post-stimulus. Thus, the effects of attention and gaze
overlapped with different somatosensory ERP components, with attention affecting the
P100 and gaze modulating the following N140 component. Both the P100 and the N140
somatosensory ERP components are considered mid-latency somatosensory ERP
components representing modality-specific stages of tactile processing, but they are
characterized by distinct neural generators. While the P100 component originates
bilaterally from SII (Hari et al., 1984; Forster and Eimer, 2003; Frot and Maguiere,
1999), multiple neural generators are likely to be responsible for the N140 component
including SII and bilateral frontal areas (Allison et al. 1992; Hari et al., 1984; 1993;
Kakigi et al., 2000; Mima et al., 1998). The dissociation between gaze-dependent and
attention-dependent modulations of somatosensory processing is not only related to the
onset of these effects, but also to their time course. Results revealed that the effects of
gaze on touch were short-lived and exclusively present in the N140 time-range, while the
effects of attention were also observed during later stages of processing (between 160 and
240 ms post-stimulus). Taken together, these results suggest that the changes provoked
by gaze and attention occur at different stages of somatosensory processing providing
additional evidence that the effects of gaze and of spatial attention are mediated by
different mechanisms.

Recently, a positive correlation has been demonstrated between the RT cuing effect
measured in an endogenous tactile attention tasks and the attentional modulations of the
N140 and Nd ERP components (Jones and Forster, 2014; see also Talsma et al., 2007, for
a direct relationship between visual N1 component amplitude and response speed. The results of the present experiment appear to confirm a relationship between the ERP modulations in the N140 time window and the behavioural performance. While it is reasonable to expect a strong relationship between the Nd attentional modulations and the behavioural performance, the present task, which was devised to provide a strong incentive for participants to fully focus attention on the task-relevant hand and to maximize the number of trials for the ERP analysis, does not allow such comparison (due to the fact that only attended targets required a response). For this reason, one question which remains open is whether the dissociation observed in the ERP data in the present study would be reflected by independent effects of gaze and attention on performance. Future behavioural studies should directly assess the behavioural effects of gaze and attention on touch when these factors are orthogonally manipulated. If gaze and attention impact different stages of processing, as suggested by the present ERP data, they should have additive effects on the behavioural data (i.e., main effects for both variables and no interaction), following the additive-factors logic (e.g. Sternberg, 1969).

Only one previous ERP study to date has described the direct impact of gaze on somatosensory processing (Forster and Eimer, 2005). Gazing to one hand resulted in enhanced negativities in the N140 time range for G+ as compared to G- in that study, while the opposite pattern of results was observed in the present study with reduced negativities for G+ as compared to G-. Furthermore, enhanced negativities for G+ compared to G- were observed in the 190-250 ms interval by Forster and Eimer (2005) while they were completely absent in the present study. One possible explanation for these discrepancies is that spatial attention was at least in part responsible for the effects of gaze observed in in Forster and Eimer study (2005). In their third experiment stimuli were randomly and unpredictably presented to either hand, therefore participants had no strategic reason to attend to a specific body location. While one might expect that under these conditions spatial attention was equally spread across the two hands locations, it is possible that more attentional resources were allocated to the gazed hand, resulting in attentional enhancements of gazed stimuli. This observation may also have relevant implications for the behavioural studies which investigated the effect of gaze on touch (Honoré et al., 1989; Pierson et al., 1991; Tipper et al., 1998). Because in these studies
gaze direction but not spatial attention were manipulated, there is the possibility that spatial attention was at least in part responsible for the observed effects of gaze. Overall, the present study provides the first direct ERP evidence that the effects of gaze and spatial attention on touch are mediated by distinct neural mechanisms.

This conclusion has relevant implications for the understanding of the mechanisms of spatial attention and its links with the oculomotor system. Previous studies have demonstrated that the attentional and oculomotor processes are closely, or even mandatorily, linked during the dynamic programming of a saccadic eye movement. For instance, planning a saccadic eye movement towards a spatial location elicits supramodal shifts of attention able to enhance the processing of stimuli presented close to the target location not only in the visual modality (e.g., Deubel and Schneider 1996; Eimer et al., 2006; 2007; Hoffman and Subramaniam 1995; Kowler et al., 1995), but also in the tactile modality (e.g., Gherri and Eimer, 2008; Gherri and Forster, 2012a; 2012b; Juravle and Deuble, 2009; Rorden et al., 2002). In contrast, when the oculomotor system is preset to maintain fixation as opposed to plan a saccade, the attentional and oculomotor processes can be dissociated, as suggested by recent studies on fixational eye movements (Tse et al., 2002; 2004; Horowitz et al., 2007). In line with these observations our results suggest that maintaining gaze on a specific body location does not necessarily results in attentional modulations of tactile stimuli presented at the gazed location. Different operations of the oculomotor system are likely to be coupled in a different way to spatial attention mechanisms. While directing the eyes (or planning to direct the eyes) toward a relevant location elicit a shift of attention toward that location, maintaining the eyes on the same location (fixation) can be de-coupled from attentional processing when gaze and spatial attention are independently manipulated, as observed in the present study.

The finding that modulations of somatosensory processing by gaze are independent of spatial attention raises the question of the mechanisms underlying this effect of gaze. Earlier studies suggested that the proprioceptive orienting of the eyes towards the tactually stimulated body site might be responsible for the observed changes in tactile processing (Honoré et al., 1989; Tipper et al., 2001; 1998). Consistent with the idea of a modulatory effect of eye position on tactile processing, recent evidence showed that gaze direction is used to encode the location of tactile stimuli. When gaze is directed to an
eccentric position, the perceived location of a tactile stimulus is systematically shifted in the direction of gaze as demonstrated by systematic errors related to gaze-direction when participants are asked to localize tactile stimuli (e.g. Harrar and Harris, 2009; Harrar et al., 2013; Pritchett and Harris, 2011). Thus, coding of tactile space requires the integration of tactile and eye position information. Importantly, such integration may only occur within higher level brain areas. Tactile stimuli are initially encoded according to a somatotopic representation of the body which is independent of body posture and of eye position. Only during later stages of processing, tactile information is recoded from somatotopic onto an external representation of space which takes into account the position of the body and of the eyes and is based on the integration of tactile with proprioceptive and visual information about the body (e.g. Azañón and Soto-Faraco, 2008; Longo et al., 2010; Röder et al., 2004). It has been suggested that the posterior parietal cortex might play a pivotal role in the remapping of touch from somatotopic into external coordinates (e.g. Azañón et al., 2010). In line with this observation, and consistent with electrophysiological studies of postural remapping of touch in external space (Heed and Röder, 2010; Rigato et al., 2013), we observed a reliable effects of gaze on touch only after 130 ms post-stimulus onset, suggesting a neural activation of areas in and beyond SII. Given that gaze direction can be used as one of the reference points against which external space is coded (e.g. Harrar and Harris, 2009; Harrar et al., 2013; Pritchett and Harris, 2011), proprioceptive signals of the position of the eyes might influence tactile processing via back projections from multimodal brain areas starting from 130 ms after stimulus presentation. While this hypothesis is speculative at present and should be further investigated in future studies, postural cues of eye position might be responsible for the effect of gaze on touch observed in the present study.

In summary, the present study has provided the first electrophysiological evidence that the effect of gaze on tactile processing is independent of spatial attention. Gazing to the tactually stimulated hand (in the absence of visual information about the hand) affects the processing of tactile events from about 130 ms post stimulus, modulating primarily the N140 somatosensory ERP component, with more positive ERPs for gazed stimuli as compared to non gazed ones. We suggest that this effect of gaze on touch might reflect the update of spatial information relative to the location of tactile stimuli according to the
current position of the eyes. This might involve the integration of tactile, proprioceptive and visual information in higher-order multimodal areas and might be mediated by back-projection from parietal to somatosensory areas, in line with the observation that the effect of gaze modulates stages of tactile processing that are beyond S1.
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Footnotes:

1 Recent evidence suggests that the position of the eyes in the orbit (central vs. eccentric eye position) can modulate attentional processing of tactile events with reduced effects of attention in the Nd time range when the eyes are directed towards an eccentric position (Gherri and Forster, 2014). Thus it is possible that the eccentric eye position held by participants in order to gaze to one of their hands in the present study reduced the late attentional modulations of somatosensory processing as compared to a standard condition in which the eyes were focused on the centre. However, the fact that we observed attentional modulations of both the P100 and Nd components in the present study suggest the presence of reliable effect of attention despite the eccentric eye position held in the present experiment.