Emotional and movement-related body postures modulate visual processing

Khatereh Borhani,^{1,2} Elisabetta Làdavas,^{1,2} Martin E. Maier,^{2,3} Alessio Avenanti,^{1,2,4} and Caterina Bertini^{1,2}

¹Department of Psychology, University of Bologna, 40126 Bologna, Italy, ²CSRNC, Centre for Studies and Research in Cognitive Neuroscience, Cesena Campus, University of Bologna, 47521 Cesena, Italy, ³Department of Psychology, Catholic University of Eichstätt-Ingolstadt, 85072 Eichstätt, Germany, and ⁴IRCCS Fondazione Santa Lucia, 00142 Rome, Italy

Human body postures convey useful information for understanding others' emotions and intentions. To investigate at which stage of visual processing emotional and movement-related information conveyed by bodies is discriminated, we examined event-related potentials elicited by laterally presented images of bodies with static postures and implied-motion body images with neutral, fearful or happy expressions. At the early stage of visual structural encoding (N190), we found a difference in the sensitivity of the two hemispheres to observed body postures. Specifically, the right hemisphere showed a N190 modulation both for the motion content (i.e. all the observed postures implying body movements elicited greater N190 amplitudes compared with static postures) and for the emotional content (i.e. fearful postures elicited the largest N190 amplitude), while the left hemisphere showed a modulation only for the motion content. In contrast, at a later stage of perceptual representation, reflecting selective attention to salient stimuli, an increased early posterior negativity was observed for fearful stimuli in both hemispheres, suggesting an enhanced processing of motivationally relevant stimuli. The observed modulations, both at the early stage of structural encoding and at the later processing stage, suggest the existence of a specialized perceptual mechanism tuned to emotion- and action-related information conveyed by human body postures.

Keywords: body postures; emotion perception; visual structural encoding; N190; early posterior negativity (EPN)

INTRODUCTION

Human body postures comprise a biologically salient category of stimuli, whose efficient perception is crucial for social interaction. Although in natural environments human bodies and faces are usually integrated into a unified percept, the neural networks underlying the processing of these two categories of stimuli, though closely related, seem to be distinct. In particular, neuroimaging evidence has demonstrated selective responses to human bodies in two focal brain regions: the extrastriate body area (EBA), located in the lateral occipitotemporal cortex (Downing *et al.*, 2001), and the fusiform body area (FBA), in the posterior fusiform gyrus (Peelen and Downing, 2005; Taylor *et al.*, 2007). Interestingly, both EBA and FBA responses generalize to schematic depictions of bodies, suggesting that body representation in these two areas is independent of low-level image features (Downing *et al.*, 2001; Peelen *et al.*, 2006).

As is the case with faces (e.g. Adolphs, 2002), the perceptual processing of bodies seems to represent a specialized mechanism, in which perception is configural (i.e. based on relations among the features of the stimulus), rather than based on the analysis of single body features. This is suggested, for example, by the inversion effect, a phenomenon in which bodies presented upside-down are more difficult to recognize than inverted objects (Reed *et al.*, 2003). At the electrophysiological level, event-related potentials (ERPs) in response to bodies show a prominent negative deflection at occipitotemporal electrodes peaking in a range between 150 and 230 ms after stimulus presentation (Stekelenburg and de Gelder, 2004; Meeren *et al.*, 2005; Van Heijnsbergen *et al.*, 2007; Minnebusch *et al.*, 2010). More specifically,

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Correspondence should be addressed to Caterina Bertini, Centre for Studies and Research in Cognitive Neuroscience, University of Bologna, Viale Europa 980, 47023 Cesena, Italy. E-mail: caterina.bertini@unibo.it

Thierry *et al.* (2006) found a negative component peaking at 190 ms post-stimulus onset (N190), reflecting the structural visual encoding of bodies, which was distinct in terms of latency, amplitude and spatial distribution compared with the typical negative component elicited by the visual encoding of faces (i.e. the N170; Rossion and Jacques, 2008). The neural generators responsible for the negative deflection in response to bodies are thought to be located in a restricted area of the lateral occipitotemporal cortex, corresponding to EBA, as suggested by source localization analysis (Thierry *et al.*, 2006), magnetoencephalographic recordings (Meeren *et al.*, 2013) and electroencephalogram (EEG)-fMRI correlation studies (Taylor *et al.*, 2010).

Studies on the perceptual processing of faces have shown that the component reflecting visual encoding (N170) is modulated by the emotional expressions of faces processed both explicitly (Batty and Taylor, 2003; Stekelenburg and de Gelder, 2004) and implicitly (Pegna et al., 2008, 2011; Cecere et al., 2014), suggesting that relevant emotional signals are able to influence the early stages of structural face encoding. In addition, non-emotional face movements, such as gaze and mouth movements, seem to be encoded at an early stage of visual processing and to modulate the N170 amplitude (Puce et al., 2000; Puce and Perrett, 2003; Rossi et al., 2014). At a later stage of visual processing (typically around 300 ms after stimulus onset), salient emotional faces are known to modulate the amplitude of the early posterior negativity (EPN), which reflects stimulus-driven attentional capture, in which relevant stimuli are selected for further processing (Sato et al., 2001; Schupp et al., 2004a; Frühholz et al., 2011; Calvo and Beltran, 2014).

Although faces represent a primary source of information about others' states (Adolphs, 2002), human bodies can also be a powerful tool for inferring the internal states of others (de Gelder *et al.*, 2010). Indeed, body postures convey information about others' actions and emotions, both of which are useful for interpreting goals, intentions and mental states. Neuroimaging studies have shown that motion and emotion-related information conveyed by bodies activates a broad network of brain regions (Allison *et al.*, 2000; de Gelder, 2006; Peelen and Downing, 2007). On the one hand, the observation of

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human motion increases activation in occipitotemporal areas close to and partly overlapping with EBA (Kourtzi and Kanwisher, 2000; Senior *et al.*, 2000; Peelen and Downing, 2005), the superior temporal sulcus (STS), the parietal cortex (Bonda *et al.*, 1996) and the premotor and motor cortices (Grèzes *et al.*, 2003; Borgomaneri *et al.*, 2014a), which might take part in perceiving and reacting to body postures (Rizzolatti and Craighero, 2004; Urgesi *et al.*, 2014). On the other hand, emotional body postures, compared with neutral body postures, enhance activation not only at the approximate location of EBA, the fusiform gyrus and STS but also in the amygdala (de Gelder *et al.*, 2004; Van de Riet *et al.*, 2009) and other cortical (e.g. orbitofrontal cortex, insula) and subcortical structures (e.g. superior colliculus, pulvinar) known to be involved in emotional processing (Hadjikhani and de Gelder, 2003; Peelen *et al.*, 2007; Grèzes *et al.*, 2007; Pichon *et al.*, 2008).

Although the pattern of neural activation for bodies conveying motion and emotion-related information suggests a similarity between perceptual mechanisms for faces and bodies, it is still unclear whether, like the information conveyed by faces, the information conveyed by body postures is already encoded at the early stage of structural representation and is therefore able to guide visual selective attention to favor the recognition of potentially relevant stimuli. Thus, this study was designed to investigate, using the high temporal resolution of ERPs, whether the structural encoding of bodies, reflected in the N190 component and visual selective attention, measured by the subsequent EPN component, are influenced by motion and emotionrelated information represented in body postures. To this end, an EEG was recorded from healthy participants performing a visual task in which they were shown pictures of bodies. These bodies had static postures (without implied motion or emotional content), impliedmotion postures without emotional content or implied motion postures expressing emotion (fear or happiness). In addition, stimuli were peripherally presented to the left or the right of a central fixation point to investigate whether the two hemispheres differentially contribute to the processing of body postures. In keeping with previous evidence that the right hemisphere plays a prominent role in responding to bodies (Chan et al., 2004; Taylor et al., 2010) and processing emotional information (Gainotti et al., 1993; Làdavas et al., 1993; Adolphs et al., 2000; Borod, 2000), a more detailed perceptual analysis of the different body postures was expected in the right hemisphere, compared with the left. More specifically, a low-level discrimination of motion-related information, reflected by an enhancement of the N190 component in response to postures with implied motion (either neutral or emotional) compared with static postures, was expected in both hemispheres. In contrast, discrimination of emotional content, reflected by an enhanced N190 in response to fearful compared with happy bodies, was only expected in the right hemisphere. Finally, at a later stage of visual processing, the salience of fearful body postures was expected to increase visual selective attention, resulting in an enhanced EPN component. Unlike the emotion-related modulation of the N190, we expected the EPN enhancement for salient fearful postures to occur in both hemispheres, since attention-related emotional modulations are known to occur in a widespread bilateral network of brain regions, including extrastriate occipital cortex, superior and inferior parietal areas and medial prefrontal regions (for a review, Pourtois and Vuilleumier, 2006).

METHODS

Participants

Twenty-two right-handed healthy volunteers (two males; mean age: 21.6 years; range: 20–26 years) took part in the experiment. They all had normal or corrected-to-normal vision. Since alexithymia is a relatively stable personality trait (Nemiah *et al.*, 1976; Taylor *et al.*, 1991),

which is known to affect emotion recognition and processing (Jessimer and Markham, 1997; Parker *et al.*, 2005), all volunteers underwent a screening for alexithymia, using the 20-item Toronto Alexithymia Scale (TAS-20; Taylor *et al.*, 2003). Only volunteers with scores in the normal range (TAS score: >39 and <61) were selected to participate. Participants were informed about the procedure and the purpose of the study and gave written informed consent. The study was designed and performed in accordance with the ethical principles of the Declaration of Helsinki and was approved by the Ethics Committee of the Psychology Department at the University of Bologna.

Experimental task

The experimental session was run in a sound-attenuated and dimly lit room. Participants sat in a relaxed position on a comfortable chair in front of a 17" PC monitor (refresh rate 60 Hz) at a distance of \sim 57 cm. Prior to the experiment, a short practice session was administered to familiarize the participants with the task.

The stimuli were presented on a PC running Presentation software (Version 0.60; www.neurobs.com) and consisted of 64 static color pictures of human bodies (two males and two females; $10^{\circ} \times 16^{\circ}$) with the faces blanked out. The images were selected from a validated database (Borgomaneri et al., 2012, 2014a). Half of the stimuli were the original pictures and the other half were mirror-reflected copies. Stimuli represented bodies in different postures, in which implied motion was absent (static body posture) or motion was implied with different emotional expressions and body movement. In particular, the body images included 16 static body postures (static body stimuli; S) in which neither motion nor emotion was implied, 16 neutral body postures in which motion was implied (neutral body stimuli; N), 16 fearful body postures in which motion was implied (fearful body stimuli; F) and 16 happy body postures in which motion was implied (happy body stimuli; H; Figure 1). Two independent psychophysical studies (Borgomaneri et al. 2012, 2014a) provided evidence that N, F and H are subjectively rated as conveying the same amount of implied motion information and as conveying more body motion information than S stimuli. Moreover, H and F were rated as more arousing than N and S. Critically, although H and F were rated as conveying positive and negative emotional valence, respectively, these two classes of stimuli received comparable arousal ratings. The stimuli were displayed against a white background, 11° to the left [left visual field (LVF) presentation] or the right [right visual field (RVF) presentation] of the central fixation point (2°) . Each trial started with a central fixation period (100 ms), followed by the stimulus (500 ms). Participants were asked to keep their gaze fixed on the central fixation and decide whether the presented stimulus was emotional (fearful or happy) or non-emotional (static or neutral) by pressing one of two vertically arranged buttons on the keyboard. The task was selected to balance the number of stimuli assigned to each response while maximizing the number of correct responses to minimize the rate of rejected epochs. Behavioral responses were recorded during an interval of 2400 ms. Half of the subjects pressed the upper button with the middle finger to emotional stimuli and the lower button with the index finger to non-emotional stimuli, while the remaining half performed the task with the opposite button arrangement. Eye movements were monitored throughout the task with electrooculogram (EOG; see below).

Participants performed 12 blocks in an experimental session of \sim 45 min. In half of the blocks, the stimuli were presented in the LVF, while in the remaining half, they were presented in the RVF. Blocks with LVF and RVF presentation were interleaved, and the sequence of the blocks was counterbalanced between participants. In each block, 67 trials were randomly presented (16 trials × 4 body stimuli: static, motion neutral, motion fearful, motion happy=64



Fig. 1 Graphical representation of the trial structure in the behavioral task. The figure depicts example trials with stimuli showing fearful (A), happy (B), neutral (C) and static body postures (D).

trials + 3 practice trials). Each participant completed a total of 768 trials (384 trials in the LVF and 384 in the RVF).

EEG recording

EEG was recorded with Ag/AgCl electrodes (Fast'n Easy-Electrodes, Easycap, Herrsching, Germany) from 27 electrode sites (Fp1, F3, F7, FC1, C3, T7, CP1, P3, P7, O1, PO7, Fz, FCz, Cz, CPz, Pz, Fp2, F4, F8, FC2, C4,T8, CP2, P4, P8, O2, PO8) and the right mastoid. The left mastoid was used as reference electrode. The ground electrode was placed on the right cheek. Impedances were kept below $5 \text{ k}\Omega$. All electrodes were off-line re-referenced to the average of all electrodes. Vertical and horizontal EOG was recorded from above and below the left eye and from the outer canthi of both eyes. EEG and EOG were recorded with a band-pass of 0.01-100 Hz and amplified by a BrainAmp DC amplifier (Brain Products, Gilching, Germany). The amplified signals were digitized at a sampling rate of 500 Hz and offline filtered with a 40-Hz low-pass filter.

ERP data analysis

ERP data were analyzed using custom routines in MATLAB 7.0.4 (The Mathworks, Natic, MA) and EEGLAB 5.03 (Delorme and Makeig, 2004; http://www.sccn.ucsd.edu/eeglab). Segments of 200 ms before and 800 ms after stimulus onset were extracted from the continuous EEG. The baseline window ran from -100 ms to 0 ms relative to stimulus onset. Epochs with incorrect responses were rejected (5.8% per body stimulus type). In addition, epochs contaminated with large

artifacts were identified using two methods from the EEGLAB toolbox (Delorme et al., 2007): (i) an epoch was excluded whenever the voltage on an EOG channel exceeded 100 µV to remove epochs with large EOG peaks and (ii) an epoch was excluded whenever the joint probability of a trial exceeded five standard deviations to remove epochs with improbable data (mean excluded epochs: 9.6%). Remaining blinks and EOG horizontal artifacts were corrected using a multiple adaptive regression method (Automatic Artifact Removal Toolbox Version 1.3; http://www.germangh.com/eeglab_plugin_aar/index.html; Gratton et al., 1983), based on the Least Mean Squares algorithm. Finally, epochs were discarded from the analysis when saccadic movements (>30 mV on horizontal EOG channels) were registered in a time window of 500 ms following stimulus onset (1.73%). The remaining epochs (mean: 83 epochs per body stimulus type) were averaged separately for each participant and each body stimulus type. The N190 amplitude was quantified as the mean amplitude in a time window of 160-230 ms post-stimulus presentation (Figure 2). Scalp topographies for the N190 component were calculated as mean amplitude in a time window of 160-230 ms post-stimulus presentation (Figure 3g and h). In addition, the EPN was calculated as the mean amplitude in a time window of 290-390 ms post-stimulus presentation (Figure 2).

Both the N190 and the EPN mean amplitudes were analyzed with a three-way analysis of variance (ANOVA) with electrode (P8, P7), visual field (LVF, RVF) and body stimulus (static: S; neutral: N; fearful: F; happy: H) as within-subjects variables. To compensate for violations of sphericity, Greenhouse–Geisser corrections were applied whenever



Fig. 2 Grand-average ERPs elicited by fearful, happy, neutral and static body postures. ERP waveforms at the representative electrodes P8 (A,B) and P7 (C,D) when stimuli were presented in the LVF (A,C) and in the RVF (B,D).

appropriate (Greenhouse and Geisser, 1959) and corrected *P* values (but uncorrected degrees of freedom) are reported. Post-hoc comparisons were performed using the Newman–Keuls test.

RESULTS ERP results

N190

The mean N190 amplitude averaged for all body stimuli (fearful, happy, neutral and static body postures) reached a maximum negative deflection in a time window of 160–230 ms on electrodes P7 and P8, as shown in the scalp topographies (Figure 3g and h). Electrodes P7 and P8 were therefore chosen as electrodes of interest in the N190 analyses, in line with previous studies (Stekelenburg and de Gelder, 2004; Thierry *et al.*, 2006). Grand average waveforms for the electrodes P7 and P8 are shown in Figure 2.

The ANOVA showed a significant main effect of Body stimulus (*F*(3, 63) = 29.14; *P* < 0.0001; $\eta_p^2 = 0.58$), a significant Electrode × Visual field interaction (*F*(1, 21) = 11.84; *P* = 0.002; $\eta_p^2 = 0.36$) and, more importantly, a significant Electrode × Visual field × Body stimulus interaction (*F*(3, 63) = 7.43; *p* = 0.0003; $\eta_p^2 = 0.26$). This interaction was further explored with two-way ANOVAs with Visual field (LVF, RVF) and Body stimulus (static, neutral, fearful, happy) as within-subjects factors for the two electrodes (P8 and P7) separately, to investigate possible differences between the two hemispheres.

The results of the ANOVA on the N190 amplitude over electrode P8, located in the right hemisphere, revealed a significant main effect of Visual field (F(1, 21) = 6.74; P = 0.016; $\eta_p^2 = 0.24$), with larger amplitudes for stimuli presented in the contralateral LVF ($-2.83 \,\mu$ V), compared with the ipsilateral RVF ($-1.65 \,\mu$ V; P = 0.016). Moreover, the

main effect of Body stimulus was significant (F(3, 63) = 18.42; $P < 0.0001; \quad \eta_p^2 = 0.47$). Post-hoc analyses showed a significantly smaller N190 amplitude in response to static postures $(-1.27 \,\mu\text{V})$ compared with all the motion postures (all $Ps \le 0.001$; H: $-2.10 \,\mu\text{V}$; N: $-2.65 \,\mu\text{V}$; F: $-2.93 \,\mu\text{V}$). In addition, a significant difference was found between the emotional postures, with a significantly larger N190 amplitude for fearful postures ($-2.93 \,\mu\text{V}$; P = 0.003) compared with happy postures $(-2.10 \,\mu\text{V})$. More importantly, the Visual field × Body stimulus interaction was significant (F(3, 63) = 6.99;P = 0.0007; $\eta_p^2 = 0.25$). Post-hoc analyses revealed that, in both the LVF and the RVF, static postures (S-LVF: $-1.60 \,\mu\text{V}$; S-RVF: $-0.93 \,\mu\text{V}$) elicited a significantly smaller N190 compared with all the motion postures (LVF: all $Ps \le 0.0001$; H-LVF: $-2.66 \,\mu\text{V}$; N-LVF: -3.24 µV; F-LVF: -3.80 µV; RVF: all Ps < 0.0006; H-RVF: $-1.54 \,\mu\text{V}$; N-RVF: $-2.06 \,\mu\text{V}$; F-RVF: $-2.07 \,\mu\text{V}$). Also, in both the LVF and the RVF, the N190 amplitude was significantly larger for fearful postures than for happy postures (F-LVF vs H-LVF: P = 0.0001; F-RVF vs H-RVF: P = 0.01). In addition, in the LVF, the N190 amplitude was significantly larger for fearful postures (F-LVF: $-3.80 \,\mu\text{V}$) than for neutral postures (N-LVF: $-3.24 \,\mu\text{V}$; P = 0.001; Figure 3a, b, e and f).

The ANOVA for electrode P7, located in the left hemisphere, revealed a significant main effect of Visual field (F(1, 21) = 11.44; P = 0.002; $\eta_p^2 = 0.35$), with larger N190 amplitudes for stimuli presented in the contralateral RVF ($-2.11 \,\mu$ V) compared with the ipsilateral LVF ($-1.14 \,\mu$ V; P = 0.002). In addition, the main effect of Body stimulus was significant (F(3, 63) = 14.26; P < 0.0001; $\eta_p^2 = 0.4$). Posthoc comparisons revealed a significantly smaller N190 amplitude in response to static postures ($-0.76 \,\mu$ V), compared with all the motion postures (all $Ps \le 0.0001$; H: $-1.71 \,\mu$ V; N: $-1.95 \,\mu$ V; F: $-2.08 \,\mu$ V;



Fig. 3 Mean N190 amplitude elicited by fearful, happy, neutral and static body postures from electrode P8 in the right hemisphere (A, B) and electrode P7 in the left hemisphere (C, D) when stimuli were presented in the LVF (A, C) and in the RVF (B, D). Scalp topographies of the difference in mean N190 amplitude between fearful and other body stimuli (happy, neutral and static) when stimuli were presented in the LVF (E) and in the RVF (F) in a time window of 160–230 ms. (G) and (H) represent scalp topographies of the mean N190 amplitude averaged for all body stimuli (fearful, happy, neutral and static body postures) in a time window of 160–230 ms when stimuli were presented in the LVF and RVF, respectively. Error bars represent standard error of the mean (SEM). LF, left fearful body posture; LH, left happy body posture; LF, left neutral body posture; LS, left static body posture; RF, right fearful body posture; RH, right happy body posture; RN, right neutral body posture; RS, right static body posture.

Figure 3c and d). However, in contrast to the results from electrode P8, the N190 amplitude recorded from electrode P7 did not significantly differ between fearful and happy body postures (P = 0.24). No other comparisons were significant (all Ps > 0.57).

EPN

The subsequent EPN amplitudes were measured at the same electrode locations as the N190, in a time window of 290–390 ms post-stimulus onset (Figure 2).

The ANOVA showed a significant main effect of Body stimulus (*F*(3, 63) = 14.87; *P*<0.0001; $\eta_p^2 = 0.41$) and, more interestingly, a significant Electrode × Visual field × Body stimulus interaction (*F*(3, 63) = 9.07; *P* = 0.0002; $\eta_p^2 = 0.3$). This interaction was further explored with two-way ANOVAs with Visual field (LVF, RVF) and Body stimulus (static, neutral, fearful, happy) as within-subject factors for the two electrodes (P8 and P7) separately, to investigate possible differences between the two hemispheres.

The ANOVA for electrode P8, in the right hemisphere, revealed a significant main effect of Body stimulus (F(3, 63) = 7.71; P < 0.0002; $\eta_p^2 = 0.27$), showing a significant more negative amplitude in response to fearful postures (0.97μ V), compared with the remaining postures (all $Ps \le 0.006$; H: 1.64μ V; N: 1.55μ V; S: 1.95μ V). The interaction Visual field × Body stimulus was also significant (F(3, 63) = 5.63; P = 0.003; $\eta_p^2 = 0.21$). Post-hoc analyses revealed that, both in the

LVF and the RVF, fearful body postures (F-LVF: 0.51μ V; F-RVF: 1.43μ V) elicited the most negative amplitude compared with the remaining postures (LVF: all *Ps* \leq 0.0001; H-LVF: 1.42μ V; N-LVF: 1.27μ V; S-LVF: 1.98μ V; RVF: all *Ps* \leq 0.03; H-RVF: 1.86μ V; N-RVF: 1.83μ V; S-RVF: 1.92μ V; see Figure 4a, b, e and f). In addition, in the LVF, happy (1.42μ V) and neutral (1.27μ V) body postures showed a significantly more negative amplitude than compared with static body postures (1.98μ V; all *Ps* \leq 0.02; Figure 4a, b, g and h).

The ANOVA for electrode P7, in the left hemisphere, revealed a significant main effect of Body stimulus (F(3, 63) = 7.47; P = 0.0002; $\eta_p^2 = 0.26$). Post-hoc comparisons revealed that negative amplitude was significantly greater in response to fearful postures (1.66μ V), compared with the remaining postures (all $Ps \le 0.006$; H: 2.34μ V; N: 2.51μ V; S: 2.75μ V; Figure 4c–f). No other comparisons were significant (all Ps > 0.48).

Behavioral results

Reaction times (RTs), accuracy scores and inverse efficiency scores (IES = reaction times/accuracy) were analyzed with separate ANOVAs with Visual field (LVF, RVF) and Body stimulus (static, neutral, fearful, happy) as within-subjects variables. The analysis on RTs revealed a significant main effect of Body stimulus (*F*(3, 63) = 31.81; *P* < 0.0001; $\eta_p^2 = 0.6$), showing faster RTs for static body postures (654 ms) compared with fearful (756 ms), happy (764 ms) and



Fig. 4 Mean EPN amplitude elicited by fearful, happy, neutral and static body postures from electrode P8 in the right hemisphere (A, B) and electrode P7 in the left hemisphere (C, D) when stimuli were presented in the LVF (A, C) and in the RVF (B, D). Scalp topographies of the difference in mean EPN amplitude between fear and other body stimuli (happy, neutral and static) when stimuli were presented in the LVF (E) and in the RVF (*F*) in a time window of 290–390 ms. Scalp topographies of the difference in mean EPN amplitude between static and other body stimuli (happy, neutral) when stimuli were presented in the LVF (G) and in the RVF (*H*) in a time window of 290–390 ms. Error bars represent standard error of the mean (SEM). LF, left fearful body posture; LH, left happy body posture; LF, left neutral body posture; LS, left static body posture; RF, right fearful body posture; RH, right happy body posture; RN, right neutral body posture; RS, right static body posture.

neutral postures (797 ms; all *Ps* < 0.0001). In addition, RTs for neutral body postures were significantly slower than for fearful (*P*=0.02) and happy body postures (*P*=0.03). The ANOVA performed on the accuracy scores revealed a significant main effect of Body stimulus (*F*(3, 63) = 8.42; *P* < 0.0001; η_p^2 = 0.29), showing that participants were slightly more accurate in responding to static body postures (98%), compared with fearful (93%; *P*=0.004), happy (93%; *P*=0.007) and neutral postures (90%; *P*=0.0001). Finally, the ANOVA on inverse efficiency scores revealed a significant main effect of Body stimulus (*F*(3, 63) = 21.62; *P* < 0.0001; η_p^2 = 0.5), showing significantly lower scores (reflecting better performance) for static body postures (661 ms; all *Ps* ≤ 0.0001), compared with fearful (811 ms), happy (822 ms) and neutral postures (891 ms). In addition, IES for the neutral body postures was significantly higher than for the remaining postures (all *Ps* < 0.02).

DISCUSSION

Seeing images of bodies elicits a robust negative deflection peaking at 190 ms post-stimulus onset (N190) reflecting the early structural encoding of these stimuli (Thierry *et al.*, 2006) and a subsequent relative negativity (EPN) indexing attentional engagement to salient stimuli (Schupp *et al.*, 2006; Olofsson *et al.*, 2008). This study revealed that information concerning both the presence of motion and the

emotions expressed by different body postures are able to modulate the early stage of the visual encoding of bodies and the attentional engagement process as reflected by changes in the amplitudes of N190 and EPN, respectively.

In particular, laterally presented pictures of bodies in different postures strongly modulated the N190 component. Interestingly, this component showed differential sensitivity to the observed body postures in the two cerebral hemispheres. On the one hand, the right hemisphere showed a modulation of the N190 both for the motion content (i.e. all the postures implying motion elicited larger N190 amplitudes compared with static, no-motion body postures) and for the emotional content (i.e. fearful postures elicited the largest N190 amplitude). On the other hand, the left hemisphere showed a modulation of the N190 only for the motion content, with no modulation for the emotional content. These findings suggest partially distinct roles of the two cerebral hemispheres in the visual encoding of emotional and motion information from bodies. In addition, at a later stage of perceptual representation reflecting selective attention to salient stimuli, an enlarged EPN was observed for fearful stimuli in both hemispheres, reflecting an enhanced processing of motivationally relevant stimuli (Schupp et al., 2006; Olofsson et al., 2008).

Electrophysiological studies suggest that, akin to the N170 for faces, the N190 component represents the process of extracting abstract and relevant properties of the human body form for categorization (Thierry *et al.*, 2006) and is considered the earliest component indexing structural features of human bodies (Taylor *et al.*, 2010). Our study expands these ideas by demonstrating that the stage of structural encoding reflected by the N190 entails not only the categorization of the visual stimulus as a body but also an analysis of motion-related and emotional features of the body posture. In other words, the visual encoding stage involves not only a perceptual representation of the form, configuration and spatial relations between the different body parts (Taylor *et al.*, 2007, 2010), but it also reflects a discrimination between body postures conveying information about the presence of actions and emotions.

It has been argued that EBA (i.e. the putative neural generator of the N190; Thierry *et al.*, 2006; Taylor *et al.*, 2010) has a pivotal role in creating a cognitively unelaborated but perceptually detailed visual representation of the human body (Peelen and Downing, 2007; Downing and Peelen, 2011), which is forwarded to higher cortical areas for further analysis. On the other hand, EBA is thought to be modulated by top-down signals from multiple neural systems, including those involved in processing emotion and action information (Downing and Peelen, 2011). Thus, the finding that the N190 is sensitive to information about motion and emotions conveyed by human body postures suggests that emotion- and action-related signals are rapidly extracted from visual stimuli and can exert a fast top-down modulation of the neural processing reflecting structural encoding of bodies in occipitotemporal areas, i.e. the N190.

The smaller N190 amplitudes for static bodies than for bodies with implied motion suggest that both hemispheres operate a perceptual distinction between bodies with static postures and bodies performing actions. Because of the highly adaptive value of motion perception, observers typically extract motion-related information from static images where motion is implied (Freyd, 1983; Verfaillie and Daems, 2002). Occipitotemporal visual areas have been suggested to encode dynamic visual information from static displays of "moving" objects (e.g. human area MT, Kourtzi and Kanwisher, 2000; STS, when implied motion information is extracted from pictures of biological entities, Puce and Perrett, 2003; Perrett et al. 2009) and to respond to static images of human body postures implying an action (Peigneux et al., 2000; Kourtzi et al., 2008). Thus, the static snapshots of moving bodies used here were not only a necessary methodological substitute for real motion that was required to reliably record ERPs but also a sufficient substitute for understanding how the human visual system represents human body movements.

Notably, action observation is also known to activate a wide frontoparietal network of sensorimotor regions involved in action planning and execution. Indeed, observing images of humans during ongoing motor acts is known to enhance the excitability of the motor system (Urgesi et al., 2010; Borgomaneri et al., 2012; Avenanti et al., 2013a,b), where the perceived action is dynamically simulated (Gallese et al., 2004; Nishitani et al., 2004; Keysers and Gazzola, 2009; Gallese and Sinigaglia, 2011). Such motor simulation appears to emerge very early in time (<100 ms after stimulus onset in some cases, e.g. van Schie et al., 2008; Lepage et al., 2010; Ubaldi et al., 2015; Rizzolatti et al., 2014) and is thought to facilitate visual perception through feedback connections from motor to visual areas (Wilson and Knoblich, 2005; Kilner et al., 2007; Schippers and Keysers, 2011; Avenanti et al., 2013a; Tidoni et al., 2013). Thus, the observed enhancement of structural encoding for postures implying motion and action compared with static postures seems to indicate increased perceptual representation of the bodies, possibly triggered by fast action simulation processes in interconnected frontoparietal areas.

On the other hand, a finer perceptual distinction, discriminating not only the presence of action but also the emotional content of that

action, is evident only in the right hemisphere, where the N190 was differentially modulated by fearful and happy body postures, with fearful postures eliciting the largest N190 amplitude. This emotional modulation of structural encoding might reflect an adaptive mechanism, in which the perceptual representation of body stimuli signaling potential threats is enhanced by top-down modulations. In line with this, neuroimaging studies have shown that fearful bodies increase the BOLD signal in the temporo-occipital areas from which the N190 originates and in nearby visual areas (Hadjikhani and de Gelder, 2003; Peelen et al., 2007; Grèzes et al., 2007; Pichon et al., 2008; Van de Riet et al., 2009). Importantly, fearful bodies are known to enhance activation in the amygdala (Hadjikhani and de Gelder, 2003; de Gelder et al., 2004; Van de Riet et al., 2009), the key subcortical structure for signaling fear and potential threat (Adolphs, 2013; LeDoux, 2014). Notably, the magnitude of amygdala activation predicts activity in EBA and FBA during perception of emotional bodies (Peelen et al., 2007). Therefore, the enhanced N190 over the right occipitotemporal electrodes might reflect a rapid and distant functional influence of the amygdala on interconnected visual cortices, useful for processing threat signals efficiently and implementing fast motor reactions (Vuilleumier et al., 2004; Borgomaneri et al., 2014b). Similarly, somatosensory and motor regions, crucial to the processing of threatrelated expressions (Adolphs et al., 2000; Pourtois et al., 2004; Banissy et al., 2010; Borgomaneri et al., 2014a), might also participate in this top-down influence. Indeed, somato-motor regions are connected to occipitotemporal areas via the parietal cortex (Keysers et al., 2010; Rizzolatti et al., 2014) and exert a critical influence on visual recognition of emotional expressions quite early in time (i.e. 100-170 ms after stimulus onset; Pitcher et al., 2008; Borgomaneri et al., 2014a), which may be compatible with the observed N190 modulation.

Although previous electrophysiological findings showed a modulation of fearful body expressions at the stage of the P1 component (i.e. before structural encoding of the stimulus has taken place; Mereen *et al.*, 2005; Van Heijnsbergen *et al.*, 2007), the potentials peaking in the range of the N1 seem to offer more reliable measures of both faceand body-selective perceptual mechanisms. Indeed, earlier potentials such as the P1 could be modulated to a greater degree by low-level features of the stimuli, as they are highly sensitive to physical properties of visual stimuli (Halgren *et al.*, 2000; Rossion and Jacques, 2008).

The observed emotional modulation of the N190 exclusively over the right hemisphere is in keeping with the idea of a possible right hemisphere advantage in processing emotions (Gainotti et al., 1993; Làdavas et al., 1993; Adolphs et al., 2000; Borod, 2000). Alternatively, the more detailed modulation of structural encoding processes observed in the right hemisphere could be due to a higher sensitivity to human bodies, as suggested by preferential activation in response to body stimuli in the right EBA (Downing et al., 2001; Chan et al., 2004; Saxe et al., 2006) and in a broad network of right cortical areas (Caspers et al., 2010). In keeping with the idea of a right hemisphere advantage in processing emotional body postures, recent transcranial magnetic stimulation studies have shown that motor excitability over the right (but not the left) hemisphere is sensitive to the emotional content of the observed body posture at a latency compatible with the initial part of the N190 component (Borgomaneri et al., 2014a). This suggests a strict functional coupling between visual and motor representations during the processing of emotional body postures, which might favor perception of and adaptive motor responses to threatening stimuli.

Interestingly, at a later stage of visual processing (i.e. 300 ms poststimulus onset), the EPN component was enhanced for fearful stimuli in both hemispheres. The EPN is a relative negativity for emotional stimuli (Schupp *et al.*, 2006). This emotional modulation reflects

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attentional capture driven by salient emotional stimuli and might reflect the degree of attention needed to recognize relevant signals (Olofsson et al., 2008). Previous studies have shown increases in the amplitude of the EPN in response to both emotional scenes (Schupp et al., 2003, 2004b; Thom et al., 2014) and emotional faces (Sato et al., 2001; Schupp et al., 2004a; Frühholz et al., 2011; Calvo and Beltran, 2014). Similar to the findings of present study, the EPN is also enhanced during observation of hand gestures, with a greater effect for negatively valenced gestures (Flaisch et al., 2009, 2011). This suggests that viewing isolated body parts with emotional relevance also modulates this component. The present results add to the previous studies by showing strong EPN sensitivity to whole body expressions of fear, supporting the idea that fearful body postures represent a highly salient category of stimuli, able to engage selective visual attention to favor explicit recognition of potentially threatening signals (de Gelder et al., 2004, 2010; Kret et al, 2011; Borgomaneri et al., 2014a). Notably, our data suggest that attentional processes are enhanced by fearful postures in both hemispheres, indicating that, at later stages of visual processing, both the right and the left hemispheres concur to engage attentional resources to aid recognition of salient emotional stimuli. However, it is interesting to note that the right hemisphere also maintains a higher capacity to discriminate between the different body postures at this later stage, as suggested by an increased negativity for happy and neutral body postures compared with static body postures.

Interestingly, the emotional modulations observed both at the early stage of structural encoding and at the later attentional engagement stage might be a by-product of the interaction between movement and emotion-related information conveyed by emotional body postures. Indeed, bodies express emotions through movements, therefore providing concurrent motion-related information. Further studies are needed to disentangle the contributions of emotion and movementrelated information by investigating ERP modulations in response to emotional body postures with a minimal amount of motion content (e.g. sad body postures). Overall, these results suggest that information pertaining to motion and emotion in human bodies is already differentially processed at the early stage of visual structural encoding (N190), in which a detailed representation of the form and configuration of the body is created.

At this early stage, the right hemisphere seems prominent in processing the emotional content of body postures, as shown by the effects of laterally presented body postures on structural encoding. At a later stage of visual processing (EPN), the relevant and salient information represented by fearful body postures recruits visual attention networks in both hemispheres, which might facilitate recognition of potentially dangerous stimuli. Finally, the modulations observed in the visual processing of different body postures, both at the visual encoding and attentional engagement stages, are reminiscent of modulations seen in visual face processing (Batty and Taylor, 2003; Stekelenburg and de Gelder, 2004; Schupp et al., 2004a; Frühholz et al., 2011; Calvo and Beltran, 2014), suggesting that face and body processing might involve distinct but similar perceptual mechanisms. This highly efficient and specialized structural encoding, and the subsequent attentional engagement for salient stimuli, may represent an adaptive mechanism for social communication that facilitates inferences about the goals, intentions and emotions of others.

Conflict of Interest

None declared.

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