Men fear other men most: gender specific brain activations in perceiving threat from dynamic faces and bodies – an fMRI study

Mariska Kret#, Swann Pichon**, Julie Grèzes and Beatrice de Gelder

INTRODUCTION

Facial and bodily expressions are among the most salient affective signals regulating our daily interactions and they have a strong biological basis (de Gelder, 2006, 2010). Therefore it stands to reason that gender figures prominently among factors that determine affective communication. Previous studies have already reported gender differences in how the brain processes facial emotions. But it is not known whether gender differences also influence how emotional expressions of the whole body are processed. It is also unclear whether there is a relation between the gender of the observer and that of the image shown. The goal of this study was to address both issues. We first give a systematic overview of the core areas that underlie the perception of facial and bodily expressions of emotion (Kret et al., 2011) and then outline the implications for gender differences.

The cortical network underlying face perception is well known and includes the fusiform face area (FFA; Kanwisher et al., 1997), the occipital face area (Puce et al., 1996; Gauthier et al., 2000), the superior temporal sulcus (STS) and the amygdala (AMG; Haxby et al., 2000). Recent studies show that the brain areas involved in whole body perception overlap with the face network and confirm the involvement of AMG, fusiform gyrus (FG), and STS in face and body perception (Hadjikhani and de Gelder, 2003; de Gelder et al., 2004; Peelen and Downing, 2007; Meeren et al., 2008; van de Riet et al., 2009; Kret et al., 2011). Two areas in the body perception network have been the targets of categorical selectivity research. The one reported first is an area at the junction of the middle temporal and middle occipital gyrus, labeled the extrastriate body area (EBA; Downing et al., 2001). A later added one is in the FG, at least partly overlapping with FFA (Kanwisher et al., 1997) and termed the fusiform body area (FBA; Peelen and Downing, 2005). Recent evidence suggests that these areas are particularly responsive to bodily expressions of emotion (Grèzes et al., 2007; Peelen et al., 2007; Pichon et al., 2008).

Yet so far the relation between categorization by the visual system and emotion perception is not clear. Furthermore, photographs of bodily expressions also trigger areas involved in action perception (de Gelder et al., 2004). Recent studies with dynamic stimuli have proven useful for better understanding the respective contribution of action and emotion-related components. A study by Grosbras and Paus (2006) showed that video clips of angry hands trigger activations that largely overlap with those reported for facial expressions in the FG. Increased responses in the STS and the temporoparietal junction (TPJ) have been reported for dynamic threatening body expressions (Grèzes et al., 2007; Pichon et al., 2008, 2009). Different studies have demonstrated a role for TPJ in “theory of mind”, the ability to represent and reason about mental states, such as thoughts and beliefs (Saxe and Kanwisher, 2003; Samson et al., 2004). Other functions of this area involve reorienting attention to salient stimuli, sense of agency, and multisensory body-related information processing, as well as in the processing of phenomenological and cognitive aspects of the self (Blanke and Arzy, 2005). Whereas TPJ is implicated in higher level social cognitive processing (for a meta-analysis, see Decety and Lamm, 2007), STS has been frequently highlighted in biological motion studies (Allison et al., 2000) and shows specific activity for goal-directed actions and configural and
kinematic information from body movements (Perrett et al., 1989; Bonda et al., 1996; Grossman and Blake, 2002; Thompson et al., 2005). Observing threatening actions (as compared to neutral or joyful actions) increases activity in regions involved in action preparation: the pre-supplementary motor area (pre-SMA; de Gelder et al., 2004; Grosbras and Paus, 2006; Grèzes et al., 2007) and premotor cortex (PM; Grosbras and Paus, 2006; Grèzes et al., 2007; Pichon et al., 2008, 2009). To our knowledge, it is still unclear whether these above described regions relate to gender differences.

Common sense intuitions view women as more emotional than men. Yet research suggests this presumed difference is based more on an expressive and less on an experiential difference (Kring and Gordon, 1998). For example, Moore (1966) found that males reported more violent scenes than females during binocular rivalry, possibly because of cultural influences that socialize males to act more violently than females. A growing body of research demonstrates gender differences in the neural network involved in processing emotions (Kemp et al., 2004; Hofer et al., 2006; Dickie and Armony, 2008). Two observations are a stronger right hemispheric lateralization but also higher activation levels in males as compared to females (Killgore and Cupp, 2002; Schienle et al., 2005; Fine et al., 2009).

A different issue is whether how the gender of the person we observe influences our percept, depends on our gender. Evidence suggests that pictures of males expressing anger tend to be more effective as conditioned stimuli than pictures of angry females (Öhman and Dimberg, 1978). Previous behavioral studies indicate enhanced physiological arousal in men but not in women during exposure to angry male as opposed to female faces (Mazurski et al., 1996). Fischer et al. (2004) observed that exposure to angry male as opposed to angry female faces activated the visual cortex and the anterior cingulate gyrus significantly more in men than in women. A similar sex-differential brain activation pattern was present during exposure to fearful but not neutral faces. Aleman and Swart (2008) report stronger activation in the STS in men than women in response to faces denoting interpersonal superiority. These studies suggest a defensive response in men during a confrontation with threatening males.

Evolutionary theorists suggest that ancestral males formed status hierarchies, and that dominant males were more likely to attract females. Men’s position within these hierarchies could be challenged, possibly explaining why men use physical aggression more often than females (Eron and Huesmann, 1984). Moreover, there are many studies reporting a relationship between high levels of testosterone and increased readiness to respond vigorously and assertively to provocations and threats (Olweus et al., 1988). A physically strong male expressing threat with his body is likely to represent a large threat and may be more relevant for the observer. It is thus conceivable that the perception of and reactivity to emotional expressions depends on the gender of the observer and observed.

Taken together, there are strong indications that males and females differ in the recruitment of cerebral networks following female and male emotional expressions. We tested this hypothesis here by measuring female and male participants’ hemodynamic brain activity while they watched videos showing threatening (fearful or angry) or neutral facial or bodily expressions of female or male actors. First, we expected male observers to react more strongly to signals of threat than females. Second, since threatening male body expressions are potentially harmful, we expected the male as compared to female videos to trigger more activation in regions involved in processing affective signals (AMG), body-related information (EBA, FG, STS, and TPJ), and motor preparation (pre-SMA and PM; de Gelder et al., 2010).

**MATERIALS AND METHODS**

**PARTICIPANTS**

Twenty-eight participants (14 females, mean age 19.8 years old, range 18–27 years old; 14 males; mean age: 21.6 years old, range 18–32 years old) took part in the experiment. Half of them viewed neutral and angry expressions and the other half neutral and fearful expressions. Participants had no neurological or psychiatric history, were right-handed and had normal or corrected-to-normal vision. All gave informed consent. The study was performed in accordance to the Declaration of Helsinki and was approved by the local medical ethical committee. Two participants were discarded from analysis, due to task miscomprehension and neurological abnormalities.

**MATERIALS**

Video recordings were made of 26 actors expressing six different facial and bodily emotions. For the body video sessions all actors were dressed in black and filmed against a green background. For the facial videos, actors wore a green shirt, similar as the background color. Recordings used a digital video camera under controlled and standardized lighting conditions. To coach the actors to achieve a natural expression, pictures of emotional scenes were, with the help of a projector, shown on the wall in front of them and a short emotion inducing story was read out by the experimenter. The actors were free to act the emotions in a naturalistic way as response on the situation described by the experimenter and were not put under time restrictions. Fearful body movements included stretching the arms as if to protect the upper body while leaning backward. Angry body movements included movements in which the body was slightly bended forward, some actors showed their fists, whereas others stamped their feet and made resolute hand gestures. Additionally, the stimulus set included neutral face and body movements (such as pulling up the nose, coughing, fixing one’s hair, or clothes). Distance to the beamer screen was 600 mm. All video clips were computer-edited using Ulead and After Effects, to a uniform length of 2 s (50 frames). The faces of the body videos were masked with Gaussian masks so that only information of the body was perceived. Based on a separate validation study, 10 actors were included in the current experiment. To check for quantitative differences in movement between the movies, we estimated the amount of movement per video clip by quantifying the variation of light intensity (luminance) between pairs of frames for each pixel (Peelen et al., 2007). For each frame, these absolute differences were averaged across pixels that scored (on a scale reaching a maximum of 255) higher than 10, a value which corresponds to the noise level of the camera. These were then averaged for each movie. Angry and fearful expressions contained equal movement (M = 30.64, SD 11.99 versus M = 25.41, SD 8.71) [t(19) = 0.776, ns] but more movement than neutral expressions (M = 10.17, SD 6.00) [t(19) = 3.78, p < 0.005] and [t(19) = 4.093, p < 0.005]. Threatening male and female video clips did not differ in the amount of movement (M = 31.48, SD 8.89 t(19) = 2.54, p < 0.05].

**RESULTS**

Female and male participants showed more activation in the AMG for threatening compared to neutral expressions. Significant differences were observed for both male and female participants: females [t(13) = 3.69, p < 0.005] and males [t(14) = 2.34, p < 0.05]. No differences were observed in the STS, FG, and TPJ.

**DISCUSSION**

This study demonstrates that threat perception is influenced by the gender of the observer. The stronger activation in the AMG for threatening compared to neutral expressions suggests that threatening male body expressions are perceived as more threatening than threatening female expressions. These results are consistent with previous studies demonstrating gender differences in the recruitment of cerebral networks following female and male emotional expressions. Future studies should investigate the role of cultural influences in the perception of emotional expressions.
The experiment consisted of 176 trials (80 non-scrambled volumes were collected for each participant [total scan time = 10 min per run (two runs with the anatomical scan in between)]. A total of 644 functional volumes were collected for each participant [total scan time = 10 min per run (two runs with the anatomical scan in between)]. The localizer scan parameters were: TR = 2000 ms, TE = 30 ms, FA = 90°, FOV = 224 mm, matrix size = 64 mm × 64 mm). An automatic shimming procedure was performed before each scanning session. A trial started with a fixation cross (500 ms), followed by a video (2000 ms) and a blank screen (2450 ms). An oddball task was used to control for attention and required participants to press a button each time an inverted video appeared so that trials of interest were uncontaminated by motor responses. Stimuli were back-projected onto a screen positioned behind the subject’s head and viewed through a mirror attached to the head coil. Stimuli were centered on the display screen and subtended 11.4° of visual angle vertically for the body stimuli, and 7.9° of visual angle vertically for the face stimuli.

**PROCEDURE**

Participants’ head movements were minimized by an adjustable padded head-holder. Responses were recorded by a keypad, positioned on the right side of the participant’s abdomen. After the two experimental runs, participants were given a functional localizer. Stimulus presentation was controlled by using Presentation software (Neurobehavioral Systems, San Francisco, CA, USA). After the scanning session participants were given a 10-min break. They were then guided to a quiet room where they were seated in front of a computer and categorized the non-scrambled stimuli they had previously seen in the scanner.

**fMRI DATA ACQUISITION**

**Parameters of the functional scans**

Functional images were acquired using a 3.0-T Magnetom scanner (Siemens, Erlangen, Germany). Blood oxygenation level dependent (BOLD) sensitive functional images were acquired using a gradient echo-planar imaging (EPI) sequence (TR = 2000 ms, TE = 30 ms, 32 transversal slices, descending interleaved acquisition, 3.5 mm slice thickness, with no interslice gap, FA = 90°, FOV = 224 mm, matrix size = 64 mm × 64 mm). An automatic shimming procedure was performed before each scanning session. A total of 644 functional volumes were collected for each participant [total scan time = 10 min per run (two runs with the anatomical scan in between)].

The localizer scan parameters were: TR = 2000 ms, TE = 30 ms, FA = 90°, 28 slices, matrix size = 256 mm × 256 mm, FOV = 256 mm, slice thickness = 2 mm (no gap), number of volumes = 328 (total scan time = 11 min).
Table 1 | Experimental design.

| Non-scrambled | Scrambled | Oddball* |
|---------------|-----------|---------|
| Anger         | Neutral   |         |
| **RUN 1**     |           |         |
| Faces         | Male      | 5 * 2   | 5 * 2   | 5 * 2   | 5 * 2   | 2       | 2       |
|               | Female    | 5 * 2   | 5 * 2   | 5 * 2   | 5 * 2   | 2       | 2       |
| Bodies        | Male      | 5 * 2   | 5 * 2   | 5 * 2   | 5 * 2   | 2       | 2       |
|               | Female    | 5 * 2   | 5 * 2   | 5 * 2   | 5 * 2   | 2       | 2       |
| Total         |           | 40      | 40      | 40      | 40      | 8       | 8       |
| **RUN 2**     |           |         |
| Faces         | Male      | 5 * 2   | 5 * 2   | 5 * 2   | 5 * 2   | 2       | 2       |
|               | Female    | 5 * 2   | 5 * 2   | 5 * 2   | 5 * 2   | 2       | 2       |
| Bodies        | Male      | 5 * 2   | 5 * 2   | 5 * 2   | 5 * 2   | 2       | 2       |
|               | Female    | 5 * 2   | 5 * 2   | 5 * 2   | 5 * 2   | 2       | 2       |
| Total         |           | 40      | 40      | 40      | 40      | 8       | 8       |

Version 1. anger–neutral (N = 14)

Version 2. fear–neutral (N = 14)

The oddball condition consisted of videos that were presented upside down. 5 * 2: five unique identities that were presented twice. All stimuli were presented in random order.

Parameters of the structural scan

A high-resolution T1-weighted anatomical scan was acquired for each participant (TR = 2250 ms, TE = 2.6 ms, FA = 9°, 192 sagittal slices, voxel size 1 × 1 × 1 mm, Inversion Time (TI) = 900 ms, FOV = 256 mm × 256 mm, 192 slices, slice thickness = 1 mm, no gap, total scan time = 8 min).

Statistical parametric mapping

Functional images were processed using SPM2 software (Wellcome Department of Imaging Neuroscience)\(^1\). The first five volumes of each functional run were discarded to allow for T1 equilibration effects. The remaining 639 functional images were reoriented to the anterior/posterior commissures plane, slice time corrected to the middle slice and spatially realigned to the first volume, subsampled at an isotropic voxel size of 2 mm, normalized to the standard MNI space using the EPI reference brain and spatially smoothed with a 6 mm full width at half maximum (FWHM) isotropic Gaussian kernel. Statistical analysis was carried out using the general linear model framework (Friston et al., 1995) implemented in SPM2.

At the first level analysis, eight effects of interest were modeled: face threat (fear or anger, depending on the version of the experiment) female, face neutral female, body threat female, body neutral female, face threat male, face neutral male, body threat male, body neutral male. Null events were modeled implicitly. The BOLD response to the stimulus onset for each event type was convolved with the canonical hemodynamic response function over a duration of 2000 ms. For each subject’s session, six covariates were included in order to capture residual movement-related artifacts (three rigid-body translations and three rotations determined from initial spatial registration), and a single covariate representing the mean (constant) over scans. To remove low frequency drifts from the data, we applied a high-pass filter using a cut-off frequency

\(^1\)www.fil.ion.ucl.ac.uk/spm
of 1/128 Hz. Statistical maps were overlaid on the SPM’s single subject brain compliant with MNI space, i.e., Colin27 (Holmes et al., 1998) in the anatomy toolbox (see Eickhoff et al., 2005 for a description). The atlas of Duvernoy was used for macroscopical labeling (Duvernoy, 1999).

The beta values of the ROIs (see next paragraph) were extracted for the following conditions: face threat male and female, body threat male and female, face neutral male and female, and body neutral male and female. The reason for combining two studies into one analysis was because the overall pattern of responses to these two types of threat stimuli was similar. Since there were almost no differences across the two hemispheric ROIs and our interest does not concern hemispheric lateralization, we pool bilateral ROIs to reduce the total number of areas. Main and interaction effects were tested in SPSS in an ANOVA with three within-subject variables [emotion (threat and neutral), category (face and body), gender of actor (female and male)] and one between subject variable (gender of observer) and were further investigated with Bonferroni-corrected paired comparisons and Bonferroni-corrected two-tailed t-tests.

LOCALIZATION OF FACE- AND BODY-SENSITIVE REGIONS

Face- and body-sensitive voxels in EBA, FG, STS, and AMG were identified using a separate localizer scan session in which participants performed a one backward task on face, body, house, and tool stimuli. The localizer consisted of 20 blocks of 12 trials of faces, bodies (neutral expressions, 10 male, and 10 female actors), objects, and houses (20 tools and 20 houses). Body pictures were selected from our large database of body expressions and we only included the stimuli that were recognized as being absolutely neutral. To read more about the validation procedure, we refer the reader to the article of van de Riet et al. (2009). The tools (for example pincers, a hairdryer etc.) and houses were selected from the Internet. All pictures were equal in size and were presented in grayscale on a gray background. Stimuli were presented in a randomized blocked design and were presented for 800 ms with an ISI of 600 ms.

Preprocessing was similar to the main experiment. At the first level analysis, four effects of interest were modeled: faces, bodies, houses, and tools. For each subject’s session, six covariates were included in order to capture residual movement-related artifacts (three rigid-body translations and three rotations determined from initial spatial registration), and a single covariate representing the mean (constant) over scans. To remove low frequency drifts from the data, we applied a high-pass filter using a cut-off frequency of 1/128 Hz. We smoothed the images of parameter estimates of the contrasts of interest with a 6-mm FWHM isotropic Gaussian kernel. At the group level, the following t-tests were performed: face > house, body > house, and subsequently a conjunction analysis [body > house AND face > house]. The resulting images were thresholded liberal (p < 0.05, uncorrected) to identify the following face- and body-sensitive regions: FG, AMG, STS, EBA (see Table 2).

ROIs were defined using a sphere with a radius of 5 mm centered onto the group peak activation. All chosen areas appeared in the whole brain analysis (faces and bodies versus scrambles) and are well known to process facial and bodily expressions (de Gelder, 2006; de Gelder and Hadjikhani, 2006; Grèzes et al., 2007; Meeren et al., 2008; Pichon et al., 2008, 2009; van de Riet et al., 2009; for a recent review, see de Gelder et al., 2010). We were also interested to investigate gender differences in TPJ (Van Overwalle and Baetens, 2009), which is involved in higher social cognition and pre-SMA and the PM that are involved in the preparation of movement and environmentally triggered actions (Hoshi and Tanji, 2004; Nachev

Table 2 | Coordinates used to create regions of interest.

| Hemisphere | Anatomical region | x    | y    | z    | Reference            | Contrast                                           |
|------------|-------------------|------|------|------|----------------------|----------------------------------------------------|
| R          | Fusiform face/body area | 42   | -46  | -22  | Localizer            | [Body > house AND face > house]                   |
| R          | Amygdala          | -42  | -46  | -22  | »                     |                                                    |
| L          | Superior temporal sulcus | 54   | -52  | 18   | Localizer            | Face > house                                      |
| L          | Extrastriate body area | 52   | -70  | -2   | »                     | Body > house                                      |
| R          | Temporo-parietal junction | 62   | -40  | 26   | 1 + 2 + 3*           |                                                   |
| L          | Premotor cortex  | -60  | -40  | 24   | 2 + 3*               |                                                   |
| R          | Pre-supplementary motor area | 42   | 44   | 2    | »                     |                                                    |
| L          |                   | -46  | 10   | 54   | 2                    |                                                   |
| R          |                   | 8    | 18   | 66   | 1                    |                                                   |
| L          |                   | -8   | 18   | 66   | »                     |                                                   |

*Average coordinate:
1. Grèzes et al. (2007), fear body > neutral body.
2. Pichon et al. (2008), anger body > neutral body.
3. Pichon et al. (2009), anger body AND fear body.
R, right; L, left.
» Coordinate was taken from the other hemisphere.
These three areas could not be located with our localizer. However, since there are arguments against using the same data set of the main experiment for the localization of specific areas (Kriegeskorte et al., 2009), we defined TPJ, PM, and pre-SMA by using coordinates from our former studies (Grèzes et al., 2007; Pichon et al., 2008, 2009). In Kret et al. (2011), we revealed that FG was equally responsive to emotional faces and bodies, and therefore we chose to pool FFA and FBA. Also, our localizer showed a considerable overlap between the two areas and our whole brain analysis on the main experiment revealed that the FG as a whole responded much more to dynamic bodies than to dynamic faces.

The localization of functional regions may differ between participants and there are arguments to define a fROI per participant and not base its location on the group level. However, the fROIs that we chose are very different from one another in terms of the location but also of the activity level and specific function. We opted for the group level for a number of reasons. First, group localization may be preferable for testing the behavior of a functional ROI if the region is small or if the criteria response is weak (for example in case of the AMG; Downing et al., 2001). Second, not all the fROIs were detectable in each participant. Localization at the individual level risks that one fails to identify an ROI in some individuals. Third, participants often show multiple peaks in one area and it is sometimes arbitrary to decide which one to take if the stronger peak lies further away from the group peak than the weaker peak. Fourth, not all our fROIs were easy to localize with our localizer and we therefore for some areas use a coordinate from the literature (see Table 2). Our definition of the ROIs has the advantage that their anatomy is easy to report. In tables, we specify exactly around which peak the sphere was drawn which may be meaningful for meta-analyses.

RESULTS

In order to examine gender differences in face and body responsive areas, we specified the gender of the actors and observers by extracting the beta values of pre-defined ROIs. We checked the patterns of both fear and anger to ensure that both factors contributed the same way to the common effects that are reported below (See Figure 3).

1. Extrastriate body area showed a main effect of emotion \(F(1,24) = 44.597, p < 0.001\) and category \(F(1,24) = 147.764, p < 0.001\) and was more active for threatening versus neutral and for bodily versus facial expressions \((p < 0.001)\). An interaction between actor and emotion was found \((F(1,24) = 4.706, p < 0.05)\). Both male and female actors induced more activity when expressing threatening versus neutral emotions (female actors: \(t(25) = 4.206, p < 0.001\); male actors: \(t(25) = 6.412, p < 0.001\)). Since we expected more brain activity in response to threatening male than threatening female actors, we conducted two planned comparison t-tests. First, we compared the difference between threatening and neutral expressions in male actors versus female actors and found a significant difference \((t(25) = 2.216, p < 0.05, \text{one-tailed})\). Second, to ascertain that the interaction was mainly driven by male threat, we compared threatening male versus threatening female expressions, which yielded a difference \((t(25) = 1.760, p < 0.05, \text{one-tailed})\). EBA showed an interaction between category and emotion \((F(1,24) = 8.775, p < 0.01)\). Faces and bodies induced more activity when expressing a threatening versus neutral emotion (faces: \(t(25) = 3.362, p < 0.05\); bodies: \(t(25) = 6.349, p < 0.001\)) yet this difference was larger in bodies versus faces \((t(25) = 11.501, p < 0.001)\). An interaction between category, emotion, and observer was found \((F(1,24) = 9.499, p < 0.005)\).

Male observers showed more activity for threatening than neutral bodies \((t(11) = 7.481, p < 0.001)\), threatening bodies than faces \((t(11) = 8.662, p < 0.001)\), and neutral bodies than faces \((t(11) = 7.661, p < 0.001)\). Female observers showed more activity for threatening than neutral faces \((t(13) = 3.987, p < 0.05)\), threatening bodies than faces \((t(13) = 7.688, p < 0.001)\), and neutral bodies than faces \((t(13) = 8.353, p < 0.001)\). The interaction was partly driven by female observers’ enhanced activity for threatening versus neutral faces but this difference was not significantly larger in female than male observers \((p = 0.227)\).

Since we expected more brain activity in response to threatening male body expressions in male observers, we conducted two planned comparisons. First, we compared male observers’ response to threatening versus neutral male body expressions \((t(11) = 3.601, p < 0.001, \text{one-tailed})\). Second, we expected this difference to be larger in male than female observers and therefore we compared the difference between threatening minus neutral male body expressions between male and female observers which yielded a difference \((t(24) = 1.716, p < 0.05, \text{one-tailed})\).

In summary, EBA not only processes bodies but is also sensitive to emotion. Moreover, highest activity was observed in male participants while watching threatening male body expressions.

2. Fusiform gyrus showed a main effect of emotion \((F(1,24) = 10.430, p < 0.005)\) and was more active for threatening than neutral expressions \((p < 0.005)\). There was an interaction between category, emotion, and observer \((F(1,24) = 4.695, p < 0.05)\). Male observers showed more activity for threatening than neutral bodies \((t(11) = 5.106, p < 0.001)\).

Since we expected more brain activity in response to threatening male body expressions in male observers, we conducted two planned comparisons. First, we compared male observers’ response to threatening versus neutral male body expressions \((t(11) = 3.054, p < 0.05, \text{one-tailed})\). Second, we expected this difference to be larger in male than female observers and therefore we compared the difference between threatening minus neutral male body expressions between male and female observers which yielded a difference \((t(24) = 1.835, p < 0.05, \text{one-tailed})\).

Fusiform gyrus was responsive to emotional expressions from faces and bodies and most responsive to threatening male bodies in male observers.

3. Superior temporal sulcus showed main effects of emotion \((F(1,24) = 21.191, p < 0.001)\) and category \((F(1,24) = 6.846, p < 0.05)\) and was more active for bodies than faces \((p < 0.05)\) and for emotional than neutral videos \((p < 0.001)\). An interaction between emotion and category was observed \((F(1,24) = 10.799, p < 0.005)\). Whereas STS did not differentially respond to emotional versus neutral faces \((p = 0.371)\),
activity was higher for emotional versus neutral bodies \((t(25) = 4.571, p < 0.005)\). The gender of the actor interacted with emotion \((F(1,24) = 7.632, p < 0.05)\). STS was more active following male threatening than neutral expressions \((t(25) = 4.669, p < 0.001)\) and threatening male versus female stimuli \((t(25) = 2.881, p < 0.05)\). An interaction between category, emotion, and observer was observed \((F(1,24) = 6.498, p < 0.05)\). Male observers showed more activity for threatening versus neutral bodies \((t(11) = 5.536, p < 0.001)\) threatening bodies versus faces \((t(11) = 4.178, p < 0.05)\) and females did not \((p = 0.420; p = 0.882)\). The interaction was partly driven by female observers' enhanced activity for threatening versus neutral faces but this was not significantly different \((p = 0.230)\). The difference between threatening versus neutral faces was not significantly larger in female versus male observers \((p = 0.810)\).

Since we expected more brain activity in response to threatening male body expressions in male observers, we conducted two planned comparisons. First, we compared male observers response to threatening versus neutral male body expressions \((t(11) = 4.251, p < 0.001, \text{one-tailed})\). Second, we expected this difference to be larger in male than female observers and therefore we compared the difference between threatening minus neutral male body expressions between male and female observers which yielded a marginally significant difference \((t(24) = 1.454, p = 0.079, \text{one-tailed})\).

Similar to EBA and FG, STS was responsive to emotional expressions from faces and bodies and was most responsive to threatening male bodies in male observers.

4. Amygdala showed a main effect of category \((F(1,24) = 22.402, p < 0.001)\) and was more active for faces than bodies \((p < 0.001)\). An interaction between category and observer was found \((F(1,24) = 4.325, p < 0.05)\). Male observers only showed more activity for faces than bodies \((\text{male observers: } t(11) = 4.914, p < 0.001 \text{ female observers: } p = 0.294)\). Inspecting the graph (see Figure 2) gives more insight into this effect. The enhanced response in the male observers for facial expressions, was only significant in case of female faces \((\text{female faces versus all bodies: } t(11) = 5.185, p < 0.001)\); male faces \((p = 0.813)\).

In contrast to our expectations, and in contrast to the activity pattern in the regions described above as well as with TPJ, the AMG was not more responsive to emotional than neutral stimuli but showed enhanced activity for faces, in particular in the male observers, but only when they observed female faces. 5. Temporo-parietal junction showed a main effect of category \((F(1,24) = 16.227, p < 0.001)\) and emotion \((F(1,24) = 4.374, p < 0.05)\) and was more active for threatening than neutral expressions \((p < 0.05)\) and for bodies than faces \((p < 0.001)\). There was an interaction between gender of the actor and observer \((F(1,24) = 4.351, p < 0.05)\). Follow-up comparisons did not yield any significant effects.

6. Premotor cortex showed a main effect of category \((F(1,24) = 4.670, p < 0.05)\) and responded more to bodies than to faces \((p < 0.05)\). There was an interaction between gender of the actor and emotion \((F(1,24) = 5.764, p < 0.05)\) but follow-up t-tests did not yield significant differences. Since we expected more brain activity in response to threatening male than threatening female actors, we conducted two planned comparison t-tests. First, we compared the difference between threatening and neutral expressions in male actors versus female actors and found a marginally significant difference \((t(25) = 1.514, p = 0.07, \text{one-tailed})\). Second, to ascertain that the interaction was mainly driven by male threat, we compared threatening male versus threatening female expressions, which yielded a difference \((t(25) = 1.720, p < 0.05, \text{one-tailed})\). Moreover, an interaction between gender of the actor and gender of the observer enhanced response in the male observers for facial expressions, was only valid in case of female faces. The amygdala was not more responsive to emotional than neutral stimuli but showed enhanced activity for faces, in particular in the male observers, especially when they observed female faces.
FIGURE 3 | Threatening facial and bodily expressions as a function of gender. Difference scores between threatening and neutral videos. *Tests are two-tailed, Bonferroni-corrected. Planned comparisons are described in the text and are not indicated in this figure. Regions are followed by the MNI coordinate. EBA, STS, pre-SMA, and PM were active following bodily expressions, especially when threatening, even more so when expressed by a male actor and above all when observed by a male participant. FG was equally responsive to faces and bodies but the interaction between category, emotion, and observer revealed more activation for threatening than neutral male bodies in male participants. AMG was more active for faces than bodies, specifically for male observers watching female faces. TPJ showed an effect of emotion and was more responsive to bodies than faces.
was found \((F(1,24) = 4.670, p < 0.05)\) but follow-up \(t\)-tests did not yield significant differences. Since we expected more brain activity in response to threatening male body expressions in male observers, we conducted two planned comparisons. First, we compared the difference in brain activity between threatening minus neutral male body expressions versus this difference for female actors which yielded a significant difference in the male observers \((t(11) = 1.853, p < 0.05,\) one-tailed). Second, we compared the difference between threatening minus neutral male body expressions in male versus female observers which yielded a difference \((t(24) = 1.827, p < 0.05,\) one-tailed).

Even though there was no significant interaction between category, observer and emotion, the activity pattern looks very similar to EBA, FG, and STS and again, responds mostly to male threatening body expressions in male observers.

7. Pre-supplementary motor area showed a main effect of gender \((F(1,24) = 9.215, p < 0.01)\) and responded more to male than female expressions \((p < 0.01)\). There was an interaction between gender of the actor and emotion \((F(1,24) = 6.438, p < 0.05)\). Pre-SMA was more responsive to threatening expressions from males than females \((t(25) = 3.555, p < 0.05)\). Moreover, an interaction between gender of the actor and gender of the observer was found \((F(1,24) = 6.157, p < 0.05)\). Although female observers were equally responsive as male observers, they showed more activity for male versus female actors, irrespective of emotion \((t(25) = 3.982, p < 0.05)\). There was an interaction between category \(\times\) emotion \(\times\) gender of the observer \((F(1,24) = 6.239, p < 0.05)\). In contrast to females, male observers were more responding to threatening than neutral bodies (female observer: \(p = 0.256\) male observer: \(t(11) = 4.170, p < 0.05\)).

Further analysis revealed that this effect in male observers was derived from threatening male versus neutral male bodies \(t(11) = 3.130, p < 0.005,\) one-tailed). We expected this difference to be larger in male than female observers and consequently compared the difference between threatening minus neutral male body expressions between male and female observers which yielded a significant difference \((t(24) = 1.944, p < 0.05,\) one-tailed).

Pre-supplementary motor area responded mostly to male threatening body expressions in male observers.

**DISCUSSION**

Previous studies showed gender differences in how the brain processes facial emotions. The present study has three innovative aspects. First, we used facial expressions but also whole body images. Second, all stimuli consisted of video clips. Third, we investigated the role of the gender of the observer as well as that of the stimulus displays.

Overall, we found a higher BOLD response in STS, EBA, and pre-SMA when participants observed male versus female actors expressing threat. But interestingly, in these regions, as well as in FG, there was an interaction between category, emotion, and observer. There was more activation for male threatening versus neutral body stimuli in the male participants. Threatening bodies and not faces triggered highest activity in STS, especially in the male participants.

Whereas male observers responded more to threatening body expressions than females did, the opposite tendency was observed in the EBA for female observers. Females were not more responsive to faces than males, but the difference in brain activity following a threatening versus neutral face was significant in this region in female observers only. However, this difference score between threatening minus neutral faces was not significantly larger in female than in male observers. So, the three-way interaction between category, emotion, and observer in EBA was mainly driven by male observers’ response to threatening body expressions. Ishizu et al. (2009) found that males showed a greater response in EBA than females when imagining hand movement. Our male subjects possibly imagined themselves reacting to the threatening male actor more than females did. Alternatively, there was an automatic, increased response to threatening body expressions (Tamietto and de Gelder, 2010). This latter explanation is plausible because the male observers additionally showed a clear motor preparation response in the PM and the pre-SMA toward threatening male body expressions.

Our results are similar to previous studies that show male observers to be more reactive to threatening signals than female observers (Aleman and Swart, 2008; Fine et al., 2009). Hess et al. (2004) finds that facial cues linked to perceived dominance (square jaw, heavy eyebrows, high forehead) are more typical for men, who are generally perceived as more dominant than women. So far, nothing is known about bodily cues and perceived dominance in humans but the physical differences between men and women may be important for interpreting our results. If there is a significant difference in power or status between men and women, then threat from an anger expression can elicit different responses depending on the status or power of the angry person.

The AMG was more active for facial than bodily expressions, independent of emotion. This effect was specifically for male observers watching female faces. This is consistent with earlier findings. Fischer et al. (2004) reported an increased response in the left AMG and adjacent anterior temporal regions in men, but not in women, during exposure to faces of the opposite versus the same sex. Moreover, AMG activity in male observers was increased for viewing female faces with relatively large pupils indicating an index of interest (Demos et al., 2008). Possibly, female faces provide more information to relevant males than male faces, whereas the distinction at the level of the face between male and female faces is less important for female observers. Other studies have reported that the AMG is face but not emotion specific (Van der Gaag et al., 2007; see Kret et al., 2011 for further discussion). But the striking fact here is that all the other areas that reflect sensitivity of the male observers are emotion sensitive, in contrast with the AMG results. This disjunction between AMG face-gender and STS, EBA, PM, pre-SMA gender–emotion sensitivity indicates that the AMG indeed plays a different role than being at the service of emotion encoding and fits with the notion that it encodes salience and modulates recognition and social judgment (Tsuchiya et al., 2009).

In our study males showed a clear motor preparation response to threatening male body language, and females did not. In males, the fight-or-flight response is characterized by the release of vasopressin. The effects of vasopressin are enhanced by the presence of
testosterone and influence the defense behavior of male animals (Taylor et al., 2002). Testosterone level is a good predictor of the presence of aggressive behavior and dominance (van Honk and Schutter, 2007) and AMG activity to angry but also to fearful faces in men (Derrnt et al., 2009). In contrast, oxytocin has caused relaxation and sedation as well as reduced fearfulness and reduced sensitivity to pain (Uvnas-Moberg, 1997). Testosterone inhibits the release of oxytocin as shown in Jezova et al. (1996; for a discussion, see Taylor et al., 2000). In addition to the increased quantity of oxytocin released in females as compared to males, McCarthy (1995) has found that estrogen enhances the effects of oxytocin. Therefore, oxytocin may be vital in the reduction of the fight-or-flight response in females. Although we cannot report any measure that could support such an interpretation, it is well known that the endocrine system plays an important role in modulating behavior. Comparing the levels of these hormones with specific neuronal responses may give more insight in these gender effects.

REFERENCES
Aleman, A., and Swart, M. (2008). Sex differences in neural activation to facial expressions denoting contempt and disgust. PLoS ONE 3, e622. doi: 10.1371/journal.pone.0003622
Allison, T., Puce, A., and McCarthy, G. (2000). Social perception from visual cues: role of the STS region. Trends Cogn. Sci. 4, 267–278.
Blanke, O., and Arzy, S. (2005). The out-of-body experience: disturbed self-processing at the temporoparietal junction. Neuroscientist 11, 16–24.
Bonda, E., Petrides, M., Ostry, D., and Evans, A. (1996). Specific involvement of human parietal systems and the amygdala in the perception of biological motion. J. Neurosci. 16, 3737–3744.
Bosson, J. K., Vandello, J. A., Burnaford, R. M., Weaver, J. R., and Arzu Wasti, S. (2009). Precarious manhood and displays of physical aggression. Pers. Soc. Psychol. Bull. 35, 623–634.
de Gelder, B. (2006). Towards the neurobiology of emotional body language. Nat. Rev. Neurosci. 7, 242–249.
de Gelder, B., and Hadjikhani, N. (2006). Non-conscious recognition of emotional body language. Neuroreport 17, 583–586.
de Gelder, B., Snyder, J., Greve, D., Gerard, G., and Hadjikhani, N. (2004). Fear fosters flight: a mechanism for fear contagion when perceiving emotion expressed by a whole body. Proc. Natl. Acad. Sci. U.S.A. 101, 16701–16706.
de Gelder, B., Van den Stock, J., Meeren, H. K. M., Sinke, C. B. A., Kret, M. E., and Tamietto, M. (2010). Standing up for the body. Recent progress in uncovering the networks involved in processing bodies and bodily expressions. Neuropsy. Biobehav. Rev. 34, 513–527.
Decety, J., and Lamm, C. (2007). The role of the right temporoparietal junction in social interaction: how low-level computational processes contribute to meta-cognition. Neuroscientist 13, 580–593.
Demos, K. E., Kelley, W. M., Ryan, S. L., Davis, F. C., and Whalen, P. J. (2008). Human amygdala sensitivity to the pupil size of others. Cereb. Cortex 18, 2729–2734.
Derrnt, B., Windischberger, C., Robinson, S., Kryspin-Exner, I., Gur, R. C., and Moser, E. (2009). Amygdala activity to fear and anger in healthy young males is associated with testosterone. Psychoneuroendocrinology 34, 687–695.
Dickie, E. W., and Armory, J. L. (2008). Amygdala responses to unattended fearful faces: interaction between sex and trait anxiety. Psychiatry Res. 162, 51–57.
Downing, P. E., Jiang, Y., Shuman, M., and Kanwisher, N. (2001). A cortical area selective for visual processing of the human body. Science 293, 2470–2473.
Duvernoy, H. M. (1999). The Human Brain: Surface, Three-Dimensional Sectional Anatomy with MRI and Blood Supply. Wien, NY: Springer Verlag.
Eickhoff, S. B., Stephan, K. E., Mohlbier, H., Grefkes, C., Fink, G. R., Amunts, K., and Zilles, K. (2005). A new SPM toolbox for combining probabilistic cytoarchitectonic maps and functional imaging data. NeuroImage 25, 1325–1335.
Eron, L. D., and Huesmann, L. R. (1984). “The control of aggressive behavior by changes in attitudes, values, and the conditions of learning,” in Advances in the Study of Aggression: Vol. I, eds R. T. Blanchard and C. Blanchard (Orlando, FL: Academic Press), 139–171.
Fine, J. G., Semrud-Clikeman, M., and Zhu, D. C. (2009). Gender differences in BOLD activation to face photographs and video vignettes. Behav. Brain Res. 201, 137–146.
Fischer, H., Sandblom, J., Herlitza, A., Fransson, P., Wright, C. I., and Backman, L. (2004). Sex-differential brain activation during exposure to female and male faces. Neuroreport 15, 233–238.
Friston, K., Holmes, A. P., Worsley, K., Poline, J., Frith, C., and Frackowiak, R. (1995). Statistical parametric maps in functional imaging: a general linear approach. Hum. Brain Mapp. 2, 189–210.
Gauthier, I., Skudlarski, P., Gore, J. C., and Anderson, A. (2000). Expertise for cars and birds recruits brain areas involved in face recognition. Nat. Neurosci. 3, 191–197.
Grèzes, J., Pichon, S., and de Gelder, B. (2007). Perceiving fear in dynamic body expressions. NeuroImage 35, 959–967.
Grohsbras, M. H., and Paus, T. (2006). Brain networks involved in viewing angry hands or faces. Cereb. Cortex 16, 1087–1096.
Grossman, E., and Blake, R. (2002). Brain areas active during visual perception of biological motion. Neuroysr. 35, 1167–1175.
Hadjikhani, N., and de Gelder, B. (2003). Seeing fearful body expressions activates the fusiform cortex and amygdala. Curr. Biol. 13, 2201–2205.
Haxby, J. V., Hoffman, E. A., and Gobbini, M. I. (2000). The distributed human neural system for face perception. Trends Cogn. Sci. 4, 223–233.
Hess, U., Adams, R. B. Jr., and Kleck, R. E. (2004). Facial appearance, gender, and emotion expression. Emotion 4, 378–388.
Hofer, A., Siedentopf, C. M., Ischebeck, A., Rettenbacher, M. A., Verius, M., and Felber, S. (2006). Gender differences in regional cerebral activity during the perception of emotion: a functional MRI study. Neuroimage 32, 854–862.
Holmes, C. J., Hoge, R., Collins, L., Woods, R., Toga, A. W., and Evans, A. C. (1998). Enhancement of MR images using registration for signal averaging. J. Comput. Assist. Tomogr. 22, 324–333.
Hoshi, E., and Tanji, J. (2004). Functional specialization in dorsal and ventral premotor areas. Prog. Brain Res. 143, 507–511.
Ishizu, T., Noguchi, A., Ino, Y., Ayabe, T., and Kojima, S. (2009). Motor activity and imagery modulate the body-selective region in the occipital-temporal area: a near-infrared spectroscopy study. Neurosci. Lett. 465, 85–89.
Jezova, D., Jurankova, E., Mosnarova, A., Kriska, M., and Skultetyova, I. (1996). Neuroendocrine response during stress with relation to gender differences. Acta Neurobiol. Exp. 56, 779–785.
Kanwisher, N., McDermott, J., and Chun, M. M. (1997). The fusiform face area: a module in human extrastriate cortex specialized for face perception. J. Neurosci. 17, 4302–4311.
Kemp, A. H., Silberstein, R. B., Armstrong, S. M., and Nathan, P. J. (2004). Gender differences in the cortical electrophysiological processing of visual emotional stimuli. NeuroImage 21, 632–646.
Killgore, W. D., and Capp, D. W. (2002). Mood and sex of participant in perception of happy faces. Percept. Mot. Skills 95, 279–288.
Kret, M. E., Pichon S., Grèzes, J., and de Gelder, B. (2011). Similarities and

CONCLUSION
Increased activation in FG, STS, EBA, PM, and pre-SMA was observed for threatening versus neutral male body stimuli in male participants. AMG was more active for facial than bodily expressions, independent of emotion, yet specifically for male observers watching female faces. Human emotion perception depends to an important extent on whether the stimulus is a face or a body and whether observers and observed are male or female. These factors that have not been at the forefront of emotion research are nevertheless important for human emotion theories.

ACKNOWLEDGMENTS
This study was partly supported by Human Frontiers Science Program RGP54/2004, NWO Nederlandse Organisatie voor Wetenschappelijk Onderzoek 400.04081 to Beatrice de Gelder, and EU FP6-NEST-COBOL043403 and FP7 TANGO to Beatrice de Gelder. We thank the two reviewers for their helpful and insightful comments and suggestions.

Frontiers in Psychology | Emotion Science
January 2011 | Volume 2 | Article 3 | 10
differences in perceiving threat from dynamic faces and bodies. An fMRI study. NeuroImage 54, 1755–1762.
Kriegeskorte, N., Simmons, W. K., Bellgowan, P. S. F., and Baker, C. I. (2009). Circular analysis in systems neuroscience: the dangers of double dipping. Nat. Neurosci. 12, 533–540.
Kring, A. M., and Gordon, A. H. (1998). Kriegeskorte, N. Simmons, W. K., Nachev, P., Hamalainen, M. S., and de Gelder, B. (2008). Functional role of the supple-
mentary motor areas of visual and motor areas by still and dynamic body expressions of anger. Soc. Neurosci. 3, 199–212.
Pichon, S., de Gelder, B., and Grèzes, J. (2009). Two different faces of threat. Comparing the neural systems for recognizing fear and anger in dynamic body expressions. NeuroImage, 47, 1873–1883.
Puce, A., Allison, T., Asgari, M., Gore, J. C., and McCarthy, G. (1996). Differential sensitivity of human visual cortex to faces, letterstrings, and textures: a functional magnetic resonance imaging study. J. Neurosci. 16, 5205–5215.
Samson, D., Apperly, I. A., Chiavarino, C., and Humphreys, G. W. (2004). Left temporoparietal junction is necessary for representing someone else’s belief. Nat. Neurosci. 7, 499–500.
Saxe, R., and Kanwisher, N. (2003). People thinking about thinking people. The role of the temporoparietal junction in “theory of mind.” NeuroImage 19, 274–285.
Schielen, A., Schafer, A., Stark, R., Walter, B., and Vaitl, D. (2005). Gender differences in the processing of disgust- and fear-inducing pictures: an fMRI study. Neuroreport 16, 277–280.
Tamietto, M., and de Gelder, B. (2010). Neural bases of the non-conscious perception of emotional signals. Nat. Rev. Neurosci. 11, 697–709.
Taylor, S. E., Klein, L. C., Lewis, B. P., Gruenewald, T. L., Gurung, R. A. R., and Updegraff, J. A. (2000). Biobehavioral responses to stress in females: tend-and-befriend, not fightor-flight. Psychol. Rev. 107, 441–442.
Taylor, S. E., Klein, L. C., Lewis, B. P., Gruenewald, T. L., Gurung, R. A. R., and Updegraff, J. A. (2002). Sex differences in biobehavioral responses to threat: reply to Geary and Flinn (2001). Psychol. Rev. 109, 751–753.
Thompson, J. C., Clarke, M., Stewart, T., Puce, A. (2005). Configural processing of biological motion in human superior temporal sulcus. J. Neurosci. 25, 9059–9066.
Tsuchiya, N., Moradi, F., Felsen, C., Yamazaki, M., and Adolphs, R. (2009). Intact rapid detection of fearful faces in the absence of the amygdala. Nat. Neurosci. 12, 1224–1225.
Uvnas-Moberg, K. (1997). Oxytocin linked antistress effects: the relaxation and growth response. Acta Physiol. Scand. 640(Suppl.), 38–42.
van de Riet, W. A. C., Grèzes, J., and de Gelder, B. (2009). Specific and common brain regions involved in the perception of faces and bodies and the representation of their emotional expressions. Soc. Neurosci. 4, 101–120.
Van der Gaag, C., Minderaa, R. B., and Keysers, C. (2007). Facial expressions: what the mir ror neuron system can and cannot tell us. Soc. Neurosci. 2, 179–222.
van Honk, J., and Schutter, D. J. L. G. (2007). Testosterone reduces conscious detection of signals serving social correction implications for antisocial behavior. Psychol. Sci. 18, 663–667.
Van Overwalle, F., and Baetens, K. I. (2009). Understanding others’ actions and goals by mirror and mentalizing systems: a meta-analysis. NeuroImage 48, 564–584.

Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Received: 15 September 2010; accepted: 03 January 2011; published online: 26 January 2011.

Citation: Kret ME, Pichon S, Grèzes J and de Gelder B (2011) Men fear other men most: gender specific brain activations in perceiving threat from dynamic faces and bodies – an fMRI study. Front. Psychology 2.3. doi: 10.3389/fpsyg.2011.00003

This article was submitted to Frontiers in Emotion Science, a specialty of Frontiers in Psychology. Copyright © 2011 Kret, Pichon, Grèzes and de Gelder. This is an open-access article subject to an exclusive license agreement between the authors and Frontiers Media SA, which permits unrestricted use, distribution, and reproduction in any medium, provided the original authors and source are credited.