The color red attracts attention in an emotional context. An ERP study

Michał Kuniecki1*, Joanna Pilarczyk1 and Szymon Wichary2

1 Psychophysiology Laboratory, Institute of Psychology, Jagiellonian University, Kraków, Poland, 2 Interdisciplinary Center for Applied Cognitive Studies, University of Social Sciences and Humanities, Warsaw, Poland

The color red is known to influence psychological functioning, having both negative (e.g., blood, fire, danger), and positive (e.g., sex, food) connotations. The aim of our study was to assess the attentional capture by red-colored images, and to explore the modulatory role of the emotional valence in this process, as postulated by Elliot and Maier (2012) color-in-context theory. Participants completed a dot-probe task with each cue comprising two images of equal valence and arousal, one containing a prominent red object and the other an object of different coloration. Reaction times were measured, as well as the event-related lateralizations of the EEG. Modulation of the lateralized components revealed that the color red captured and later held the attention in both positive and negative conditions, but not in a neutral condition. An overt motor response to the target stimulus was affected mainly by attention lingering over the visual field where the red cue had been flashed. However, a weak influence of the valence could still be detected in reaction times. Therefore, red seems to guide attention, specifically in emotionally-valenced circumstances, indicating that an emotional context can alter color’s impact both on attention and motor behavior.

Keywords: attention, emotion, visual perception, the color red, event-related potentials

Introduction

Color vision aids the visual processing of natural scenes on several levels, such as scene segmentation (Hansen and Gegenfurtner, 2009), object recognition, and stimulus discrimination (Gegenfurtner and Rieger, 2000). Colors may also guide attention towards important objects, since color is used both in nature and culture as a powerful signal (Hutchings, 1997). Information regarding hue is extracted at early stages of visual processing (Gegenfurtner, 2003), which makes its role in triggering early attention shifts plausible. Vivid colors, especially, or those particularly fine-tuned to the perceptual scope of a visual system, can be used to convey messages of two basic kinds: repelling and attracting (King, 2005). In the animal world stripes of contrasting colors, such as seen on wasps, are often used to repel potential predators by signaling unpleasant consequences of attack, however same contrasting colors on a courting bird are an invitation to approach. Interestingly, attraction and repulsion also comprise the basic pleasant-unpleasant dimension of emotion (Lang et al., 1998a), therefore colors may be relevant to the processing of emotional stimuli. In fact, people tend to assign an emotional meaning to particular hues (Moller et al., 2009) and have consistent preferences regarding colors (Palmer and Schloss, 2010). However, some studies reject a general role of color in the processing of emotional stimuli (Bradley et al., 2001; Junghöfer et al., 2001; Codispoti et al., 2011).
Recently, Elliot and Maier (2012) proposed a theory of colors’ impact on psychological functioning. According to their color-in-context theory, color carries meaning, and this has a direct and automatic influence on cognitive processes, including attention. This influence is consistent with the emotional evaluation of color as either hospitable or hostile. Thus, color may facilitate approach- or avoidance-oriented psychological processes, serving as an automatically and rapidly processed affective prime. Colors’ connotations stem from pairings of a particular color with experiences, objects and messages, which have sources both in biology and culture. However, these connotations are not uniform, and colors’ impact on behavior is context-dependent. The modulatory role of the context proposed by Elliot and Maier’s (2012) theory is plausible even in case of fast processes like attention switching, given the temporal course of the semantic analysis of visual stimuli. People are able to extract gist of a scene, even when it is presented for only a split second (12–26 ms; Thorpe et al., 1996; Bacon-Macé et al., 2005; Rousselet et al., 2005), and outside of the focus of attention (Li et al., 2002; Peelen et al., 2009). More importantly, emotionally-loaded visual stimuli are promptly discriminated, even when their presentation is very brief (Junghöfer et al., 2001; Schupp et al., 2004), and when they are concealed among distracting neutral stimuli (Ohman et al., 2001; Calvo et al., 2006). The aim of the present study was to test Elliot and Maier’s theory by exploring the engagement of attention by color, and the modulatory role of emotional context in this process.

Red is a particularly good example of the aforementioned properties of colors, which prompted several lines of research within the color-in-context theory framework. Firstly, it was established that viewing red immediately before or during a motor response increases the response’s strength and velocity, most probably due to the elicitation of fear (Elliot and Aarts, 2011). However, when participants are exposed to red several seconds before the motor task, it impairs the motor production, because a relatively distal threat cue causes anxiety rather than fear (Payen et al., 2011). Secondly, red seems to possess emotion-eliciting properties. Moller et al. (2009) have shown that people tend to associate red with negative, danger-bearing emotions, since it is the color of fire, blood, anger, and sometimes of poisonous or dangerous animals. Yet, red does not always signal hostility or danger. Among many species (e.g., primates and fish), red is an evolved biological signal of attractiveness (e.g., Milinski and Bakker, 1990; Waitt et al., 2006). In humans, women and men wearing red clothes are regarded by the opposite sex as more desirable (Elliot et al., 2010; Kayser et al., 2010). The other positive connotation of red is in food, namely in ripe fruits. Notably, detection of ripe fruits is a possible reason why trichromacy has evolved in primates (Sumner and Mollon, 2000; Surridge et al., 2003; Osorio and Vorobyev, 2005). Clearly, red has a double meaning: it may signal either threat or opportunity, depending on the context. This duality is reflected in infants’ preference for red in safe conditions, (i.e., red stimulus primed by a smiling face), and avoidance of this color in threatening circumstances (i.e., red stimulus primed by an angry face; Maier et al., 2009).

The emotional connotation of red switches between negative and positive, but in both emotional extremes red signals the presence of a significant stimulus and thus should require an attentional shift towards it. Indeed, Fortier-Gauthier et al. (2013) have recently shown that simple red targets to be detected among gray distractors evoke larger N2pc than green targets, matched in other physical qualities, pointing to possible special status of the color red in visual search. The N2pc component provides a marker of the spatial location of the attentional focus, being more negative at the posterior scalp locations contralateral to the attended stimulus (Luck and Hillyard, 1994). Before, attentional bias towards red was postulated on a less straightforward basis. Hill and Barton (2005) concluded that wearing red clothes in combat sports significantly increased the chance of victory. Hagemann et al. (2008) argued that this tendency can be explained, at least partially, by the referees’ perceptual bias towards red.

Although attentional bias to red is apparent not only in emotional circumstances (Fortier-Gauthier et al., 2013), the primary context of red seems to be an emotionally arousing one, not one that is calm and neutral. Thus, in line with Elliot and Maier’s (2012) theory, red should have different properties in emotional and neutral circumstances. Provided that signaling is one of the most important functions of color, red should affect attention, particularly in emotional conditions. Indeed, red is the most widespread signaling color in the natural world, as it is easily visible both on the blue background of the sky, and on the green of foliage (Humphrey, 1976). It is also commonly used in the urban environment to raise an alert and draw one’s attention, e.g., through the use of road signs, traffic lights, and significant notices.

We aimed to test alternative accounts of how the color red might influence the processing of emotional visual stimuli. On one side, red might serve as a cue that a given image is important, and thus a shift of attention is desirable. In that case, all red images should capture attention, regardless of their emotional meaning. Alternatively, a possible attentional bias towards red may interact with valence, in which case both positive and negative, but not neutral, red images should attract attention. Finally, red may not influence attention at all, leading to random attending to the red and non-red stimuli.

In order to test the specific influence of red on the capturing of attention, we used a version of the Posner spatial cueing task (Posner et al., 1980), called the “dot-probe”, originally introduced by MacLeod et al. (1986) to study attentional biases towards emotional vs. neutral stimuli. In our modification of this task, a cue comprising a pair of images from the same valence category, one of which always contained a red object while the other did not, were presented simultaneously, one on each side of the screen. They were followed by a target (an asterisk) displayed to the left or right side of the visual field; this target prompted the participant to press the corresponding
button on the response box. Comparing reaction times (RTs) in congruent conditions (the target appearing on the side of the red image) to reaction times in incongruent conditions (the target on the side opposite the red image) should allow us to detect an attentional bias towards red in each valence condition. Additionally, we presented cues formed by pairs of images where neither comprised the color red, which created a control condition dubbed “non-aligned”.

Apart from RTs, we also measured the early attentional bias using the event-related potentials (ERP), which can be extracted from the ongoing EEG recording. Changes in event-related potentials may appear faster than any overt behavioral reaction. ERP components that mark shifting of attention in space initiated by cue presentation, particularly early directing attention negativity (EDAN) and subsequent anterior directing attention negativity (ADAN), were of special interest. Both EDAN and ADAN are called event-related lateralizations, as they assume more negative values contralaterally to the location of attentional focus. EDAN exhibits posterior-occipital scalp distribution in the time window between 200 and 400 ms after cue onset. It is hypothesized to represent either a shift of attention caused by decoding the meaning of the cue (Harter et al., 1989; Hopf and Mangun, 2000; Nobre et al., 2000; Talsma et al., 2005; Praamstra and Kourtis, 2010) or the selection of a relevant cue feature (van Velzen and Eimer, 2003; van der Lubbe et al., 2006; Jongen et al., 2007; Brigiani et al., 2009). Thereby, according to van Velzen and Eimer (2003), EDAN is a reaction to cue analogous to the target-related N2pc. A following component, ADAN, commencing around 300 ms post cue and lasting till 500 ms has frontal distribution (Praamstra et al., 2005; van der Lubbe et al., 2006) and represents a modality-independent attentional control mechanism (Hopf and Mangun, 2000; Nobre et al., 2000; Eimer et al., 2002, 2003; Seiss et al., 2007, 2009). In addition to the lateralized EEG components, we examined the occipito-parietal P1 component with latency around 100 ms, to check for efficacy of valence manipulation. A number of EEG studies utilizing various image presentation schemes suggest that cortical differentiation between affective categories occurs as early as 100 ms after stimulus onset, evidenced by the modulation of the P1 by valence of presented stimuli (Smith et al., 2003; Carretié et al., 2004; Delplanque et al., 2004; Keil et al., 2007; Van Strien et al., 2009; Bublatzky and Schupp, 2012; Feng et al., 2014; for review, see Olofsson et al., 2008).

According to the interaction hypothesis consistent with Elliot and Maier’s (2012) theory, we expected to observe more prominent EDAN and ADAN components contralateral to the visual field where the red image was shown, especially in the emotional condition. We also anticipated to find shorter reaction times and less errors in trials in which the target appeared in the same visual field as the red image (congruent condition), compared with the incongruent and non-aligned conditions. We assumed that also this effect of the color red would depend on the emotional valence of the stimulus pair, with more pronounced facilitation of the behavioral reactions in case of the emotional stimuli compared with the neutral ones.

### Materials and Methods

**Participants**

Twenty-three (5 male and 18 female) students from the Jagiellonian University, Kraków, Poland, with an age range of 18 to 21 years (M = 19.3, SD = 0.7), participated in the study for course credit. All participants were right-hand dominant, had normal or corrected-to-normal vision, and did not report any color vision deficiencies. Participants gave written informed consent. The experiment was conducted according to guidelines of the ethic committee of Institute of Psychology, Jagiellonian University in Kraków.

**Stimuli**

Forty-five pairs of images from the International Affective Picture System (IAPS; Lang et al., 1999) were selected for the experiment. The first subset was composed of thirty pairs: ten negative, ten positive, and ten neutral. One image of each pair contained a predominant red object (“Red” in Table 1), the other lacked any shades of red (“Non-red” in Table 1), while both had equal ratings of valence and arousal and, whenever possible, were similar in terms of content and layout. The remaining fifteen pairs (five for each valence category) served as a control condition for the color manipulation (“Non-aligned” in Table 1) with both images in each pair lacking red coloring, but meeting the other criteria of selection described above. Brightness and contrast, calculated as the mean and SD of the luminance component in CIELAB color space, were equated in each pair of images using Adobe Photoshop. Table 1 presents IAPS ratings of images and their basic physical properties compared across experimental conditions. For the sake of clarity in the study, we made sure that the selected pictures depicted neither faces, due to the unique mechanism of face processing (Allison et al., 1999; Alpers et al., 2011, for review, see Dekowska et al., 2008), nor sexual scenes, whose valence and arousal ratings differ largely between men and women (Lang et al., 1999).

**Procedure**

The stimuli were presented against a gray background on a 22-inch flat TFT monitor with a refresh rate of 100 Hz. The software package DMDX (Forster and Forster, 2003) was used to control the experiment. The response pad was connected directly to the recording equipment via an optical transducer.

The experiment was carried out in a dimly-lit chamber. Participants were seated in a comfortable chair approximately 70 cm away from the screen. After receiving both verbal and written instructions, participants completed the dot-probe task.
In a pair contained a red feature, while sharing other properties. Secondly, to create a control condition in which neither image
attention from being guided simply by feature-driven pattern. This prevented the subjects’ purposes. Firstly, to maintain uncertainty regarding the color
with pairs in which red was featured. This design served two
noting is that all pairs lacking a red object were randomly mixed
spanning from 5.5° to 28.6°, either to the left or the right of the center. Although the pairing of the pictures was permanent, their relative location on the screen was randomized. Worth noting is that all pairs lacking a red object were randomly mixed with pairs in which red was featured. This design served two purposes. Firstly, to maintain uncertainty regarding the color properties of an upcoming cue; this prevented the subjects’ attention from being guided simply by feature-driven pattern. Secondly, to create a control condition in which neither image in a pair contained a red feature, while sharing other properties of experimental pairs, which enabled us to assess the pure
role of color. Finally, the target—an asterisk (*, of size 1.2° × 1.2°) appeared for 150 ms, presented 16.7° to the left or right side of the screen. Laterality of the target location was fully counterbalanced across the experiment. The participants were instructed to make a response (as quickly and as accurately as possible) when the asterisk was flashed on the screen, by pressing the button corresponding to the location of the target with their left or right thumb, accordingly. The participants were informed that before each target a pair of images would be flashed and were told not to pay attention to it. No further information about images was given. The next trial immediately followed the response.

Sides of the red image and target display were independent. Combined, they created two conditions: congruent with the target spatially aligned with the red cue, and incongruent with the target and cue appearing on the opposite sides of the screen. Naturally, this division was impossible to maintain in the case of pairs lacking a red feature, due to the absence of the color cue. Hence, those trials formed the non-aligned condition.

**EEG Recording**

The EEG was recorded with the ActiveTwo BioSemi system (BioSemi, Amsterdam, The Netherlands) from thirty-two monopolar locations (Fp1/Fp2, AF3/AF4, F3/F4, F7/F8, FC1/FC2, FC5/FC6, C3/C4, T7/T8, CP1/CP2, CP5/CP6, P3/P4, P7/P8, PO3/PO4, O1/O2, Fz, Cz, Pz, Oz). All the electrodes were placed on the scalp using an Electro-Cap according to the 10–20 system, and referenced to the common mode sense (CMS) electrode with an additional drive right leg (DRL) electrode serving as a ground. The ocular activity was recorded with four bipolar electrodes placed at the outer canthi of each eye for horizontal movements, and above and below the middle of the left eye for vertical movements. An additional two electrodes were placed at both mastoids. The EEG signal was recorded at a sampling rate of 512 Hz with a 104 Hz anti-aliasing filter was continuously stored on a computer for offline analysis. Using BrainVision analyzer software (BrainProducts GmbH), the data were offline re-referenced to the average of the mastoids, and filtered with bandpass filter 0.016–30 Hz (12 dB). Subsequently, segments

---

### TABLE 1 | IAPS ratings (valence and arousal) and basic physical features (luminance, contrast, energy in low and high spatial frequencies) of negative, neutral and positive images, separately for images containing color red, non-red images, and images from control non-aligned condition.

|                 | Negative      | Neutral       | Positive      | Statistics |
|-----------------|---------------|---------------|---------------|------------|
|                 | Red           | Non-red       | Non-aligned   | Red        | Non-red | Non-aligned | Red        | Non-red | Non-aligned |
| Valence         | 2.91          | 2.88          | 2.80          | 5.06       | 5.09    | 4.97        | 7.10       | 7.13    | 7.17        |
| Arousal         | 6.03          | 5.80          | 5.92          | 3.73       | 3.58    | 3.37        | 4.99       | 4.90    | 4.96        |
| Contrast        | 0.49          | 0.49          | 0.55          | 0.53       | 0.52    | 0.52        | 0.49       | 0.49    | 0.50        |
| Luminance       | 86            | 87            | 79            | 105        | 106     | 90          | 79         | 78      | 96          |
| HSF (μ-score)   | −0.20         | 0.03          | −0.32         | −0.11      | 0.04    | 0.30        | −0.31      | 0.12    | −0.19       |
| LSF (μ-score)   | −0.13         | −0.06         | 0.13          | 0.35       | 0.44    | −0.10       | −0.26      | −0.38   | 0.03        |

Energy in high spatial frequency (HSF; >16 cycles per image) and low spatial frequency (LSF; <8 cycles per image) were calculated using algorithm provided by Delplanque et al. (2007). Luminance and contrast were calculated as the mean and SD of the luminance component in CIELAB color space. ANOVA analyses were performed with factors of valence (negative, positive, neutral) and image type (red, non-red, non-aligned). In statistics columns F(2,81), values for main effects and F(4,81) values for interactions are reported, with **indicating p < 0.001 and n.s. indicating p > 0.05.
FIGURE 1 | Overview of trial structure in congruent, incongruent and non-aligned conditions. Presented images are similar to IAPS images used in the experiment.

of 600 ms in duration were extracted for each trial, starting 100 ms before the image onset. The first 100 ms period of each segment was defined as a baseline. Data were edited for the artifacts by rejecting trials with zero activity, voltage steps over 50 µV, and voltage differences larger than 100 µV. Segments contaminated by ocular artifacts were excluded from the analysis by rejecting trials with HEOG activity greater than 40 µV. Artifact-free trials in which participants correctly responded to the target stimulus, were averaged separately for valence type (positive, neutral, negative) location of red cue (left and right visual field), and SOA (50 or 100 ms). On average 55.4 trials (SD = 4.9) were used to calculate ERP waveforms, after rejection of 7.5% of trials (Table 2). ANOVA analysis conducted on the number of averaged trials with factors of valence, location of the red cue, and SOA revealed no significant main effects. Only the interaction of valence with location of cue proved to be significant $F_{(2,42)} = 5.3$, $p = 0.009$. Pairwise comparison revealed that this interaction was due to difference in positive condition. It is unlikely that this effect impacted on our ERP results, since there was no corresponding effect in either EDAN ($F_{(2,42)} = 0.84$, $p = 0.44$), or ADAN ($F_{(2,42)} = 0.19$, $p = 0.83$) components.

Lateralized EDAN and ADAN potentials were calculated using standard procedure involving averaging of mean EEG activity recorded contralaterally and ipsilaterally in respect to the location of the red cue (Seiss et al., 2009). Since for every cue in any valence and SOA conditions there was the same number of targets in the left and right visual fields, subtraction and averaging procedure over left and right electrode locations eliminated the overlap of cue and target-related EEG activity, including motor preparation (for discussion of this issue see Luck, 2014; for similar experimental design see Anzorge et al., 2009). Following Praamstra et al. (2005, 2009), we scored EDAN and ADAN as a mean EEG activity over pooled electrode locations. The specific time windows and electrode locations were determined using grand-average waveforms, and scalp topographies. For EDAN, with time window 200–250 ms, we averaged electrode pairs O1/O2, P3/P4, PO3/PO4, and P7/P8,
TABLE 2 | Mean number of artifact-free trials (with SD reported in brackets) used to calculate EDAN and ADAN components for all valence (negative, neutral, positive), SOA (50 ms and 100 ms), and location of red cue (left or right visual field) conditions.

|               | SOA 50 ms |               | SOA 100 ms |
|---------------|-----------|---------------|------------|
|               | Left VF   | Right VF      | Left VF    | Right VF   |
| Negative      | 56.2 (4.2) | 55.4 (6.4)    | 55.3 (6.2) | 55.9 (5.2) |
| Neutral       | 55.1 (4.7) | 55.1 (6.5)    | 55.9 (5.2) | 55.4 (5.3) |
| Positive      | 54.7 (5.1) | 55.5 (6.5)    | 54.5 (5.3) | 56.5 (4.1) |

while for ADAN, with time window 300–350 ms, the following ones: FC1/FC2, AF3/AF4, F3/F4, and F7/F8. Non-aligned trials were not taken into account in this analysis, as they lacked a laterally-located cue which could serve as a reference.

The non-lateralized P1 component in response to cue onset was scored as a mean EEG activity on parietal and occipital locations (PO4, O2, Oz, O1, PO3) in time window 70–100 ms. The P1 was calculated for each valence condition, regardless whether a pair was congruent, incongruent or non-aligned.

**Statistical Analyses**

A valence (positive, neutral, negative) by congruency (congruent, incongruent, non-aligned) repeated measures ANOVA was conducted on reaction times (medians aggregated for participant and condition) and arc-sine transformed mean error rates. Reaction time analysis was performed on all correct responses between 100 and 1000 ms.

The mean amplitude of P1 component was analyzed using repeated measures ANOVA with factors of valence (positive, neutral, negative), electrode (PO4, O2, Oz, O1, PO3) and SOA (50 and 100 ms). With respect to EDAN and ADAN components two complementary analyses were conducted. Firstly, mean raw ERP data over pooled locations in EDAN and ADAN time windows were investigated using repeated measures ANOVA with factors of valence (positive, neutral, negative), red cue visual field (left, right) lateralization of electrode cluster (ipsilateral, contralateral to the red cue) and SOA (50 and 100 ms). Secondly, lateralized EDAN and ADAN components were analyzed using repeated measures ANOVA with factors of valence (positive, neutral, negative), and SOA (50 and 100 ms). In all ANOVA analyses with factors featuring more than two levels, if the sphericity assumption was violated, the Huynh-Feldt correction was applied, and the adjusted *p*-values are reported. All simple effects were investigated using post hoc comparisons with the Bonferroni correction. Data from one participant had to be rejected from the analyses due to very high error rates (29.5% vs. the average of 7.5%), leaving data from 22 participants for all further analyses.

**Results**

**Behavioral Performance Accuracy**

The overall pattern of accuracies and reaction times with respect to congruency and valence is shown in Figure 2. Error rates were affected by congruency (*F*\(_{2,42}\) = 6.9, *p* = 0.002, *η\(_p^2\) = 0.248), such that participants’ responses were most frequently correct in the congruent (94.7%), and least correct in the incongruent condition (92.4%), with intermediate accuracy in the non-aligned condition (93.5%). It seems that, as expected, a red cue facilitated responding on the side where it appeared, causing both a decrease of performance in the incongruent condition, and its increase in the congruent condition, whereas absence of the color cue indeed acted as a neutral baseline. Also, the valence altered response accuracy (*F*\(_{2,42}\) = 7.0, *p* = 0.002, *η\(_p^2\) = 0.251). That is, participants performed slightly more accurately in the positive (93.9%) than in the neutral condition (93.4%), and the lowest accuracy was observed in the negative condition (93.1%). Most importantly, valence and congruency factors produced an interaction effect (*F*\(_{4,84}\) = 3.7, *p* = 0.008, *η\(_p^2\) = 0.149), which was further investigated using simple effects tests. In the negative valence condition, congruency resulted in significantly higher accuracy than incongruency (*p* = 0.006), or the absence of the red cue (*p* = 0.002). The positive primes elicited a slightly distinct pattern of response: participants performed the task more correctly both in the congruent and non-aligned trials, compared with the incongruent trials (*p* = 0.033 and *p* = 0.019, respectively). Finally, the analysis revealed that in the neutral condition, congruency of the prime did not alter accuracy significantly.

**Reaction Times Time-Locked to Target**

Overall, participants were faster to respond when the target was preceded by the red cue (M = 392 ms), than in the incongruent (402 ms) or the non-aligned condition (402 ms),
which was reflected in the effect of congruency ($F_{(2,42)} = 39.7$, $p < 0.001$, $\eta_p^2 = 654$). Although there was no significant main effect of valence, the emotional content of the cue interacted with the congruency factor, affecting the reaction times significantly ($F_{(4,84)} = 3.8$, $p < 0.007$, $\eta_p^2 = 154$). A follow-up analysis revealed that in the congruent condition both negative and positive cues shortened the response time, in comparison to the neutral cue condition, by 4.9 ms ($p = 0.036$) and 7.8 ms ($p = 0.01$), respectively. On the contrary, an incongruent cue or the absence of red cue evoked no differences in reaction times with respect to valence. Summing up, this pattern of results indicates that red is an effective cue on the behavioral level, reducing the latency of reaction to the target, whereas the overall emotional charge of a given pair of images bears only an indirect relationship to RTs, enhancing the effect of color cueing.

**Cue-Locked ERP Components**

The P1 component elicited at occipito-parietal locations proved to be sensitive to the affective value of presented images (Figure 3). Differences in the P1 amplitude evoked by negative ($M = 1.1$, $SD = 3.1$), positive ($M = 0.66$, $SD = 2.9$), and neutral cues ($M = 0.33$, $SD = 2.81$) were significant, as validated by main effect of valence, $F_{(2,42)} = 10.13$, $p < 0.001$, $\eta_p^2 = 0.325$.

![Cue-locked P1 component for negative, neutral and positive condition elicited at occipito-parietal locations.](image.png)
Pairwise comparison revealed that P1 elicited by negative cues differed significantly from P1 elicited by neutral ($p = 0.004$) and positive cues ($p = 0.021$), but P1 to positive cues did not differ from P1 to neutral ones ($p = 0.14$). No other effects or interactions were significant in the P1 amplitude analysis.

Statistical analysis confirmed the presence of EDAN component (Figure 4) by significant effect of lateralization.
(F(1,21) = 6.7, p = 0.017, η² = 0.242) with potentials assuming smaller values at electrodes contralateral to the red cue (M = 3.97, SD = 2.7) than at electrodes ipsilateral to the red cue (M = 4.13, SD = 2.7). Crucially, substantiating the influence of context on attention deployment, the interaction between valence and laterality proved to be significant, F(2,42) = 10.5, p < 0.001, η² = 0.333. Further investigation of this interaction with simple effects revealed that in the negative and positive conditions values at contralateral electrodes were more negative (M = 3.98, SD = 2.75 and M = 3.68, SD = 2.72, respectively) than at ipsilateral electrodes (M = 4.33, SD = 2.86 and M = 4.03, SD = 2.73, respectively), while in the neutral condition an opposite effect emerged with potentials at contralateral electrodes being slightly more positive (M = 4.25, SD = 2.79) than at ipsilateral ones (M = 4.02, SD = 2.63). These simple effects were statistically significant, with p = 0.005 for negative, p = 0.001 for positive, and p = 0.045 for neutral condition. Furthermore, the main effect of SOA proved to be significant (F(1,21) = 8.94, p = 0.007, η² = 0.299), as well as interaction of SOA with red cue visual field (F(1,21) = 4.71, p = 0.042; η² = 0.183). This effect was driven by the overall larger values of mean EEG activity in 50 ms SOA condition as compared with 100 ms SOA condition, especially if the red cue appeared in the left visual field (M = 4.55, SD = 3.77 for 50 ms SOA and M = 3.51, SD = 3.72 for 100 ms SOA condition), as compared with right visual field (M = 4.31, SD = 2.5 for 50 ms SOA and M = 3.82, SD = 2.26 for 100 ms SOA condition).

Calculation of lateralized EDAN produced an enhanced parietal negativity on the scalp contralateral to the red cue in both emotional conditions, while in the neutral condition slightly opposite effect emerged (Figure 5). This resulted with EDAN assuming negative values in both positive (M = −0.35, SD = 0.43) and negative valences (M = −0.34, SD = 0.51), as opposed to neutral one (M = 0.23, SD = 0.51). The difference was overall significant (F(2,42) = 10.5, p < 0.001, η² = 0.333), and pairwise comparisons revealed that it was indeed driven by neutral condition differing significantly from both negative (p < 0.004) and positive (p < 0.001) conditions. Neither main effect of SOA nor its interaction with valence reached statistical significance.

Similarly, the ADAN component (Figure 4) was substantiated by significant main effect of lateralization (F(1,21) = 13.4, p = 0.001, η² = 0.39), which confirmed that potentials at contralateral electrodes were more negative (M = −3.73, SD = 4.02) than at ipsilateral electrodes (M = −3.53, SD = 4.11). Again, significant interaction of lateralization with valence (F(2,42) = 5.72, p = 0.006, η² = 0.214) supported the idea

**FIGURE 5** | Cue-locked grand average waveforms for (A) EDAN and (B) ADAN, obtained by subtracting ERPs at ipsilateral from contralateral locations to the red image, separately for negative, neutral, and positive slides. EDAN was calculated from pooled parietal locations (O1/O2, P3/P4, PO3/PO4, P7/P8) and ADAN from pooled frontal locations (FC1/FC2, AF3/AF4, F3/F4, F7/F8). The gray bar indicates time-window used to obtain amplitudes for corresponding effects. Scalp topographies present difference in activity contra-ipsilateral to the red cue for the left hemisphere and difference ipsi-contralateral for the right hemisphere in the time window of EDAN (C) and ADAN (D) components.
of modulatory influence of context on attentional shift. A follow up analysis of simple effects showed that ADAN was more negative at contralateral sites than at ipsilateral sites for negative cues \((M = -3.74, SD = 4.19\) for contralateral and \(M = -3.39, SD = 4.3\) for ipsilateral location) and for positive cues \((M = -3.93, SD = 4.02\) for contralateral and \(M = -3.6, SD = 4.07\) for ipsilateral location), while in case of neutral cues this difference was negligible \((M = -3.53, SD = 4.05\) for contralateral and \(M = -3.6, SD = 4.14\) for ipsilateral locations). Consequently, simple effect of laterality was significant in case of negative \((p = 0.003)\) and positive conditions \((p = 0.004)\) but not in case of neutral condition \((p = 0.46)\). Additionally, main effect of SOA was also significant \((F(1,21) = 59.35, p < 0.001, \eta^2_p = 0.739)\) with smaller mean EEG activity in 50 ms SOA condition \((M = -2.48, SD = 3.87)\) as compared with 100 ms SOA condition \((M = -4.78, SD = 4.34)\).

Lateralized ADAN (Figure 5) exhibited negative deflection in positive \((M = -0.33, SD = 0.47)\) and negative \((M = -0, 35, SD = 0.48)\) conditions as compared to neutral one \((M = 0.06, SD = 0.41)\). This significant effect \((F(2,42) = 5.72, p < 0.006, \eta^2_p = 0.214)\) was driven by the difference between negative and neutral \((p < 0.026)\) and positive and neutral \((p < 0.038)\) conditions. Neither main effect of SOA nor its interaction with valence proved significant.

In summary, the attention-capturing properties of the color red seemed to be modulated by the overall emotional value of stimuli. When presented images were emotionally charged, the EDAN and ADAN components reflected an attention shift towards the red image. On the other hand, in case of the neutral pairs, the data indicated no significant shift of attention.

**Discussion**

In the present study, using a dot-probe task coupled with EEG recording, we demonstrated that the color red captures attention and facilitates congruent motor response, particularly in an emotional context.

Participants were faster and more accurate in their responses when the target followed a red-colored cue. The effect was significantly stronger when the color cue was emotionally loaded, revealing interaction between the physical feature and the emotional valence of the image. This is consistent with propositions put forward by Elliot and Maier (2012) that color influences psychological functioning, and that the actual manifestation of this influence depends on the context. Specifically, it appears that the color red generally facilitates motor output. The magnitude of this effect is potentiated by the emotional context, a phenomenon most likely mediated by the evolutionary meaning of red, which serves as a significant signal, both in the appetitive and the aversive conditions.

The hypothesis postulating an early automatic shift of attention by red emotional images is corroborated by the analysis of the event-related potentials. We found that both the EDAN and ADAN components were larger contralateral to the red cue, but only if the cue was emotional. We also observed significant effects of SOA on raw EEG in both EDAN and ADAN time windows. However, these effects do not interact with laterality, representing EEG activity averaged across contra- and ipsilateral electrode locations. As we are only interested in comparing activity contra- and ipsilateral to the red cue, our main experimental effect is the interaction of laterality with valence. From this perspective, effects of SOA that do not interact with laterality cannot be interpreted in any meaningful way. Moreover, SOA effects disappear completely when analysis is conducted on the subtracted data.

Cueing by color observed only after emotional images suggests that the appearance of an emotional image increased the visual cortex activation linked to the processing of a potentially significant stimulus. This in turn might have led to enhanced cortical susceptibility to the evolutionarily relevant features, such as red coloration. Indeed, the visual cortex activation, evidenced by the magnitude of the P1 component, was modulated by valence of the presented cue within the first 100 ms after image onset. The P1 evoked by both negative and positive cues was numerically larger than evoked by neutral cues. However, it was most eminent in the negative condition and proved to differ significantly from the P1 in positive and neutral conditions, possibly reflecting so-called negativity bias (Smith et al., 2003; Delplanque et al., 2004; Bublatzky and Schupp, 2012; Feng et al., 2014). It has been suggested that the P1 enhancement in emotional context reflects more efficient sensory processing of potentially relevant stimuli (Smith et al., 2003; Carretié et al., 2004; Keil et al., 2007; Bublatzky and Schupp, 2012; Pourtois et al., 2013; Feng et al., 2014). The evidence indicating that emotional stimuli boost the activation of visual cortex has been also obtained in neuroimaging studies (e.g., Lang et al., 1998b; Kuniecki et al., 2003).

Closer examination of color with valence interaction reveals that in contrast to the emotional conditions, in the neutral condition the EDAN component proved to be more pronounced ipsilateral to the red image. This result might appear somewhat surprising, since it suggests that in the case of neutral stimuli, early attention was directed away from the red cue. Observed lack of attention engagement in neutral condition might be explained by the superfluous nature of the color red in a non-emotional context, due to a lack of evolutionary significance in the neutral images. Therefore, in the case of the neutral stimuli, attention was either not as focused as it was in the case of the emotional stimuli, or it was quickly directed away from the red image.

Reaction times support this hypothesis, as the difference in RTs between congruent and incongruent trials was significantly larger in the negative and the positive, compared with the neutral condition. The occurrence of a red image caused the shift of attention chiefly in the emotional conditions, which must have affected motor preparation accordingly. Indeed, shifts of the exogenous attention are in most cases linked to motor preparation (Sheliga et al., 1997). Uncoupling of those two processes is possible, though it requires very specific experimental manipulation (Praamstra and Oostenveld, 2003; Praamstra et al., 2005, 2009; Cespón et al., 2012). Elliot and Aarts (2011) obtained evidence that the occurrence of the color red, as compared with blue and gray, enhanced motor...
processes, such as the time and strength of a reaction. Facilitated motor responses corresponding to attentional preference for red over green targets were observed also in Fortier-Gauthier et al. (2013) study. The authors concluded that this property of red, observed also in our study, should be taken into account while choosing stimuli for an experiment with reaction times’ measurement.

Although the EDAN stands for early directing attention negativity it is by no means “early” in terms of the chronometry of visual processing. Foote and Simpson (2002) have shown that evoked potentials occurring around 200 ms after a stimulus onset do not represent the first feedforward sweep of visual information reaching the visual cortex. Instead, they arise as a result of multiple iterations of interactive processing between several cortical and possibly subcortical areas. Accordingly, the observed modulation of the P1 by valence substantiates the idea that differences in the following EDAN component represented the combined influence of emotional valence and coloration, because the visual analysis of both of these features must have been considerably advanced at this stage.

Indeed, the influence of both semantic and physical features on the EDAN component was observed in earlier studies. Presenting an asymmetric cue, such as an arrow, can affect EDAN due to difference in physical features rather than due to the meaning of the cue. van Velzen and Eimer (2003) showed that the EDAN component is not dependent on the direction the cue is pointing to, but instead on the spatial asymmetry of the cue itself. Later it was confirmed that EDAN is related to the asymmetry inherently present in arrow cues (Jongen et al., 2007; Brignani et al., 2009). Additionally, pairs of symmetric cues do not evoke EDAN, even if they are valid, as shown by Brignani et al. (2009) who cued target location by identical textures. On the other hand, Ranzini et al. (2009) found that in certain conditions EDAN can be elicited by completely symmetrical cues, namely by centrally presented numbers. They explain this result in terms of “spatial-numerical association of response codes” (SNARC) theory, stipulating overlearned automatic association between numerals and location in space (Dehaene et al., 1993), i.e., large numbers are mentally associated with the right part of the space, whereas small numbers are associated with the left part (Hubbard et al., 2005). The numerals were effective cues, shifting attention and influencing EDAN, therefore their meaning must have been decoded before the target commenced. In our experiment, the cue images were identical in terms of emotional valence and basic visual properties. The asymmetry was created by the presence of the red hue. Interestingly, the color asymmetry on its own did not influence EDAN in neutral condition, but only did so in emotional condition. Therefore, it seems that in our case the effect of physical asymmetry was modulated by rapid decoding of the images’ semantic meaning. It is plausible that this interaction stems from overlearned association between red hue and emotional stimuli resembling in that aspect the SNARC effect.

Additionally, because cues provided by the IAPS slides were irrelevant to the task, we assume that the enhanced EDAN component reflected a perceptual and automatic bottom-up process, rather than a more elaborate, decision-based top-down process. It appears that even if the explicit instruction requires the participant to ignore the color cue, the early detection system working in the background is unable to completely disregard the color as a valuable hint. This supports Elliot and Maier’s idea that color influences psychological functioning automatically. Indeed, Theeuwes (1992), and Kim and Cave (1999), have shown that an irrelevant-to-the-task but salient color singleton captures spatial attention. This effect persists even if participants are informed in advance that the colored singleton should be ignored (Theeuwes et al., 2006). Importantly, these effects do not depend on any special quality of the color, but rather on its perceptual distance to its surroundings. This might raise doubts if in our case the observed effect is also color-unspecific, and relies chiefly on the relative consistency of the red hues dominating over the background. Although it is not feasible to entirely refute this possibility given our experimental design, the significant interaction of EDAN and ADAN with the valence of images provides strong support for the idea that the color red was not determined to be attention-worthy throughout all stimulus presentations, but only in those with emotional content. Thus, the effect we report here cannot be exhaustively explained by a simple color pop-out. We are not able to conclude if this effect is exclusive for red hue, as we do not contrast effects of red and a single different color. It is possible that other hue would also produce similar results, even regardless of its emotional connotations. However, the unique influence of red on facilitation of motor responses and attentional preference was observed in studies directly comparing effects of different hues (Elliot and Aarts, 2011; Payen et al., 2011; Fortier-Gauthier et al., 2013).

In our experiment the target immediately followed the cue, which might lead to the conclusion that effects attributed to shifts of attention are in fact related solely to motor preparation. Our results however do not provide support for such interpretation. Firstly, the occipito-parietal topography of EDAN and frontal topography of ADAN components are in agreement with literature (Hopf and Mangun, 2000; Nobre et al., 2000; Praamstra et al., 2005; Praamstra and Kourtis, 2010; Van der Lubbe and Utzerath, 2013). Importantly, neither of these components exhibited central-frontal distribution characteristic to motor activity. Secondly, calculation of EDAN and ADAN involved averaging over the same number of left and right target occurrences, therefore cancelling out any attention-unrelated (i.e., motor) lateralizations (Luck, 2014). Double subtraction procedure used for the calculation of EDAN and ADAN, as well as N2pc, compensates also for the overlap of EEG activity evoked by cue and target, leaving only cue-locked activity (Ansorge et al., 2009). It is important to note, however, that ADAN can be evoked not only by attentional shift, but also by motor preparation (Praamstra et al., 2005, 2009; Eimer and van Velzen, 2006; van der Lubbe et al., 2006; Gherri et al., 2007). Motor-related ADAN is bound to the presentation of the cue predicting movement direction or hand selection. The more unambiguous the cue in respect to the expected motor activity, the larger the ADAN component (Praamstra et al., 2009). In our case the cues were irrelevant to the location of the forthcoming target and hence were not valid.
is therefore unlikely that the cue-locked differences observed in the ADAN time window are entirely related to motor processes.

Lastly, it is worth noting that relatively few studies have explicitly tested how emotional processing impacts visual perception. For example, Brosch et al. (2011) explored the relative influence of task-irrelevant emotional cues (angry faces), and exogenous cues (a bright border placed around an image) on early attention deployment indexed by the N2pc component. They established that while the exogenous, physical cue influenced the N2pc, the valence of the cue did not. While this result is inconsistent with the majority of findings exploring the engagement of attention by emotional faces (clearly showing modulation of the N2pc by emotional cues; Eimer and Kiss, 2007; Fenker et al., 2010; Feldmann-Wüstefeld et al., 2011; Ikeda et al., 2013), it provides evidence that—under specific experimental conditions—the exogenous cueing in the emotional dot-probe paradigm is capable of capturing attention. Our results are analogous, insofar as red coloration of the image is regarded as a kind of exogenous cue. However, the present study suggests that the emotionality of the cue plays a role too, by modulating the attention capture through the evolutionarily relevant physical feature.

Phelps et al. (2006), in a purely behavioral study using facial expressions as emotional cues, showed that emotion interacts with early visual processing by lowering the contrast required to identify target stimuli. Working in the same vein, Bocanegra and Zeelenberg (2009) obtained evidence that emotion indeed improves discrimination, but only of stimuli with low spatial frequency. Conversely, detection of high spatial frequency stimuli was hindered by the prior exposition of emotional faces. This finding is corroborated by evoked potential evidence showing that low, but not high, spatial frequencies are crucial for early discrimination of emotional from neutral stimuli (Alorda et al., 2007; Carretié et al., 2007). These results suggest that the emotional context selectively facilitates perception of specifically emotion-related aspects of a visual stimulus. Since the color red possesses a signaling value characteristic for the emotional content (e.g., Humphrey, 1976; Elliot and Pazda, 2012), our findings provide further evidence which strengthens this hypothesis.

In summary, our results fit into the color in context theory advocated by Elliot and Maier (2012), as the impact of the color red on motor behavior and attentional shifts is modulated by the emotional context of the presented stimuli. At the relatively early stage of visual processing, only the emotional red stimuli attract attention, while the neutral red images are ignored. At the behavioral level, the color red speeds response in all conditions, however the magnitude of this gain is context-dependent. Furthermore, this process seems to be purely automatic and bottom-up in nature. Our results suggest also that the color of stimuli should be controlled in studies concerning reaction times and attention, especially if the presented stimuli have emotional significance.

Acknowledgments

This work was supported by the Polish National Science Centre [grant number N N106 288939 and 2012/07/E/HS6/01046]. Images used in Figure 1 are shared on Creative Commons Licence by John Sullivan, Mike Pingleton, Tad Arensmeier and Piet Spaans.

References

Allison, T., Puce, A., Spencer, D. D., and McCarthy, G. (1999). Electrophysiological studies of human face perception. I: potentials generated in occipitotemporal cortex by face and non-face stimuli. Cereb. Cortex 9, 415–430. doi: 10.1093/cercor/9.5.415

Alorda, C., Serrano-Pedraza, L., Campos-Bueno, J. J., Sierra-Vázquez, V., and Montoya, P. (2007). Low spatial frequency filtering modulates early brain processing of affective complex pictures. Neuropsychologia 45, 3223–3233. doi: 10.1016/j.neuropsychologia.2007.06.017

Alpers, G. W., Adolph, D., and Pauli, P. (2011). Emotional scenes and facial expressions elicit different psychophysiological responses. Int. J. Psychophysiol. 80, 173–181. doi: 10.1016/j.ijpsycho.2011.01.010

Ansorge, U., Kiss, M., and Eimer, M. (2009). Goal-driven attentional capture by invisible colors: evidence from event-related potentials. Psychon Bull. Rev. 16, 648–653. doi: 10.3758/pbr.16.4.648

Bacon-Macé, N., Macé, M. J., Fabre-Thorpe, M., and Thorpe, S. J. (2005). The time course of visual processing: backward masking and natural scene categorisation. Vision Res. 45, 1459–1469. doi: 10.1016/j.visres.2005.01.004

Bocanegra, B. R., and Zeelenberg, R. (2009). Emotion improves and impairs early vision. Psychol. Sci. 20, 707–713. doi: 10.1111/j.1467-9280.2009.02354.x

Bradley, M. M., Codispoti, M., Cuthbert, B. N., and Lang, P. J. (2001). Emotion and motivation I: defensive and appetitive reactions in picture perception. Emotion 1, 276–298. doi: 10.1016/S1528-3542(13)2.376

Brignani, D., Guzzon, D., Marzi, C. A., and Minnissi, C. (2009). Attentional orienting induced by arrows and eye-gaze compared with an endogenous cue. Neuropsychologia 47, 370–381. doi: 10.1016/j.neuropsychologia.2008.09.011

Brosch, T., Pourtois, G., Sander, D., and Vuilleumier, P. (2011). Additive effects of emotional, endogenous and exogenous attention: behavioral and electrophysiological evidence. Neuropsychologia 49, 1779–1787. doi: 10.1016/j.neuropsychologia.2011.02.056

Bublatzky, F., and Schupp, H. T. (2012). Pictures cueing threat: brain dynamics in viewing explicitly instructed danger cues. Soc. Cogn. Affect. Neurosci. 7, 611–622. doi: 10.1093SCAN/SNR032

Calvo, M. G., Avero, P., and Lundqvist, D. (2006). Facilitated detection of angry faces: initial orienting and processing efficiency. Cogn. Emot. 20, 785–811. doi: 10.1080/02699930500465224

Calvo, M. G., Nummenmaa, L., and Hyöna, N. (2008). Emotional scenes in peripheral vision: selective orienting and gist processing, but not content identification. Emotion 8, 68–80. doi: 10.1016/S1528-3542.8.1.68

Carretié, L., Hinojosa, J. A., López-Martin, S., and Tapia, M. (2007). An electrophysiological study on the interaction between emotional content and spatial frequency of visual stimuli. Neuropsychologia 45, 1187–1195. doi: 10.1016/j.neuropsychologia.2006.10.013

Carretié, L., Hinojosa, J. A., Martín-Loeches, M., Mercado, F., and Tapia, M. (2004). Automatic attention to emotional stimuli: neural correlates. Hum. Brain Mapp. 22, 290–299. doi: 10.1002/hbm.20037

Cespón, J., Galdo-Álvarez, S., and Díaz, F. (2012). The Simon effect modulates N2cc and LRP but not the N2pc component. Int. J. Psychophysiol. 84, 120–129. doi: 10.1016/j.ijpsycho.2012.01.019

Codispoti, M., DeVezac, A., and Ferrari, V. (2011). The influence of color on emotional perception of natural scenes. Psychophysiology 49, 11–16. doi: 10.1111/j.1469-8986.2011.01284.x
Maier, M. A., Barchfeld, P., Elliott, A. J., and Pekrun, R. (2009). Context specificity of implicit preferences: the case of human preference for red. Emotion 9, 734–738. doi: 10.1037/a0016818

Milinski, M., and Bakker, T. C. M. (1990). Female sticklebacks use male coloration in mate choice and hence avoid parasitized males. Nature 344, 330–333. doi: 10.1038/344330a0

Möller, A. C., Elliott, A. J., and Maier, M. A. (2009). Basic hue-meaning associations. Emotion 9, 898–902. doi: 10.1037/a0017811

Nobre, A. C., Sebestyen, G. N., and Miniussi, C. (2000). The dynamics of shifting visuospatial attention revealed by event-related potentials. Neuropsychologia 38, 964–974. doi: 10.1016/s0028-3932(00)00115-4

Öhman, A., Lundqvist, D., and Esteves, F. (2001). The face in the crowd revisited: a threat advantage with schematic stimuli. J. Pers. Soc. Psychol. 80, 381–396. doi: 10.1037/0022-3514.80.3.381

Olofsson, J. K., Nordin, S., Sequeira, H., and Polich, J. (2008). Affective picture processing: an integrative review of ERP findings. Biol. Psychol. 77, 247–265. doi: 10.1016/j.biopsycho.2007.11.006

Osorio, D., and Veronoy, M. (2005). Photoreceptor spectral sensitivities in terrestrial animals: adaptations for luminance and colour vision. Proc. Biol. Sci. 272, 1745–1752. doi: 10.1098/rspb.2005.3156

Palmer, S. E., and Schloss, K. B. (2010). An ecological valence theory of human color preference. Proc. Natl. Acad. Sci. U S A 107, 8877–8882. doi: 10.1073/pnas.0906172107

Payen, V., Elliott, A. J., Coombes, S. A., Chalabaev, A., Brisswalter, J., and Cury, F. (2011). Viewing red prior to a strength test inhibits motor output. Neurosci. Lett. 495, 44–48. doi: 10.1016/j.neulet.2011.03.032

Peelen, M. V., Fei-Fei, L., and Kastner, S. (2009). Neural mechanisms of rapid scene categorization in human visual cortex. Nature 460, 94–97. doi: 10.1038/nnature08103

 Phelps, E. A., Ling, S., and Carrasco, M. (2006). Emotion facilitates perception and potentiates the perceptual benefits of attention. Psychol. Sci. 17, 292–299. doi: 10.1111/j.1467-9280.2006.00810.x

Posner, M. I., Snyder, C. R., and Davidson, B. J. (1980). Attention and the detection of signals. J. Exp. Psychol.: Hum. Percept. Perform. 6, 280–291. doi: 10.1037/0096-3445.7.2.160

Pourtois, G., Schettino, A., and Vuilleumier, P. (2013). Brain mechanisms for emotional influences on perception and attention: what is magic and what is not. Biol. Psychol. 92, 492–512. doi: 10.1016/j.biopsycho.2012.02.007

Praamstra, P., Rousset, G., and Humphreys, G. W. (2005). Fronto-parietal control of spatial attention and motor intention in human EEG. J. Neurophysiol. 94, 764–774. doi: 10.1152/jn.00152.2004

Praamstra, P., and Kourtis, D. (2010). An early parietal ERP component of the frontoparietal system: EDAN not = N2pc. Brain Res. Cogn. Brain Res. 41, 330–333. doi: 10.1016/j.cogbrres.2009.02.008

Ranzini, M., Dehaene, S., Piazza, M., and Hubbard, E. M. (2009). Neural mechanisms of attentional shifts due to irrelevant spatial and numerical cues. Neuropsychologia 47, 2615–2624. doi: 10.1016/j.neuropsychologia.2009.05.011

Rousselet, G., Joubert, O., and Fabre-Thorpe, M. (2005). How long to get to the “git” of real-world natural scenes? Vis. Cogn. 12, 852–877. doi: 10.1080/1350628044400553

Schupp, H. T., Junghofer, M., Weike, A. I., and Hamm, A. O. (2004). The selective processing of briefly presented affective images: an ERP analysis. Psychophysiology 41, 441–449. doi: 10.1111/j.1469-8986.2004.00174.x

Seiss, E., Driver, J., and Eimer, M. (2009). Effects of attentional filtering demands on preparatory ERP's elicited in a spatial cueing task. Clin. Neurophysiol. 120, 1087–1095. doi: 10.1016/j.clinph.2009.03.016

Shepherd, M., and Müller, H. J. (1989). Movement versus focusing of visual attention. Percept. Psychophys. 46, 146–154. doi: 10.3758/bf03204974

Smith, N. K., Cacioppo, J. T., Larsen, J. T., and Chartrand, T. L. (2003). May I have your attention, please: electrocortical responses to positive and negative stimuli. Neuropsychologia 41, 171–183. doi: 10.1016/s0028-3932(02)00147-1

Suriidge, A. K., Osorio, D., and Mundy, N. I. (2003). Evolution and selection of trichromatic vision in primates. Trends Ecol. Evol. 18, 198–205. doi: 10.1016/s1469-8986(03)00012-0

Talsma, D., Slagter, H. A., Nieuwenhuis, S., Hage, J., and Kok, A. (2005). The orienting of visuospatial attention: an event-related brain potential study. Brain Res. Cogn. Brain Res. 25, 117–129. doi: 10.1016/j.cogbrainres.2005.04.013

Theeuwes, J. (1992). Perceptual selectivity for color and form. Percept. Psychophys. 51, 599–606. doi: 10.3758/bf03211658

Theeuwes, J., Reimann, B., and Mortier, K. (2006). Visual search for featural singleton: no top-down modulation, only bottom-up priming. Vis. Cogn. 14, 466–489. doi: 10.1080/13506280500195110

Thorpe, S., Fize, D., and Marlot, C. (1996). Speed of processing in the human visual system. Nature 381, 520–522. doi: 10.1038/381520a0

van der Lubbe, R. H. J., Neegers, S. F. W., Verleger, R., and Kenemans, J. L. (2006). Spatiotemporal overlap between brain activation related to saccade preparation and attentional orienting. Brain Res. 1072, 133–152. doi: 10.1016/j.brainres.2005.11.087

van der Lubbe, R., and Utzerath, C. (2013). Lateralized power spectra of the EEG as an index of visuospatial attention. Adv. Cogn. Psychol. 9, 184–201. doi: 10.2478/v10153-008-0144-7

van Strien, J. W., Langeslag, S. J., Strekalova, N. J., Gootejes, L., and Franken, I. H. (2009). Valence interacts with the early ERP old/new effect and arousal with the sustained ERP old/new effect for affective pictures. Brain Res. 1251, 223–235. doi: 10.1016/j.brainres.2008.11.027

van Velzen, J., and Eimer, M. (2003). Early posterior ERP components do not reflect the control of attentional shifts toward expected peripheral events. Psychophysiology 40, 827–831. doi: 10.1111/j.1469-8986.2000.00083

Waitt, C., Gerald, M. S., Little, A. C., and Kraselbaur, E. (2006). Selective attention toward female secondary sexual color in male rhesus macaques. Am. J. Primatol. 68, 738–744. doi: 10.1002/ajp.20264

Yiend, J. (2010). The effects of emotion on attention: a review of attentional processing of emotional information. Cogn. Emot. 24, 3–47. doi: 10.1080/02699930903205698

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.

Copyright © 2015 Kuniecki, Pilarczyk and Wichary. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution and reproduction in other forums is permitted, provided the original author(s) or licensor are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.