In the blink of an eye: neural responses elicited to viewing the eye blinks of another individual

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Facial movements have the potential to be powerful social signals. Previous studies have shown that eye gaze changes and simple mouth movements can elicit robust neural responses, which can be altered as a function of potential social significance. Eye blinks are frequent events and are usually not deliberately communicative, yet blink rate is known to influence social perception. Here, we studied event-related potentials (ERPs) elicited to observing non-task relevant blinks, eye closure, and eye gaze changes in a centrally presented natural face stimulus. Our first hypothesis (H1) that blinks would produce robust ERPs (N170 and later ERP components) was validated, suggesting that the brain may register and process all types of eye movement for potential social relevance. We also predicted an amplitude gradient for ERPs as a function of gaze change, relative to eye closure and then blinks (H2). H2 was only partly validated: large temporopo-occipital N170s to all eye change conditions were observed and did not significantly differ between blinks and other conditions. However, blinks elicited late ERPs that, although robust, were significantly smaller relative to gaze conditions. Our data indicate that small and task-irrelevant facial movements such as blinks are measurably registered by the observer’s brain. This finding is suggestive of the potential social significance of blinks which, in turn, has implications for the study of social cognition and use of real-life social scenarios.

Keywords: eye movements, faces, social cognition, eye blinks, ERPs

INTRODUCTION

In this digital age, the addition of blinks to the onscreen behavior of an avatar or cartoon character can make it appear “alive” and therefore is an important component of the animation process (Thomas and Johnston, 1995; Heudin, 2004). In real-life, spontaneous blinks in humans typically occur on average 20–50 times a minute (Bentivoglio et al., 1997). The considerable variation in spontaneous blinking occurs because blinking serves a number of very important non-social functions. Blinking protects the corneal surface by allowing it to be remoistened and cleaned, and will vary with individual characteristics of the tear-film or whether an individual wears contact lenses or not (Ousler et al., 2008). Blinks can also function as a defensive reflex action that not only protects the corneal surface from incoming particles and potential physical injury, but is also elicited by startle (Pearce, 2008). Aside from these protective mechanisms for the cornea, changes in blinking behavior have been shown to occur as a function of non-ocular factors. For example, decreases in blink rates have been associated with memory operations, thinking, and attentive behaviors (Holland and Tarlow, 1975; Hirokawa, et al., 2004). It is well-known that in a number of neurological disorders, such as schizophrenia, Tourette’s syndrome, Parkinson’s disease, and progressive supranuclear palsy, patients have been reported to show increased blink rates that are thought to be associated with aberrant changes to dopaminergic systems in the brain (Chan and Chen, 2004; Karson, et al., 1984; Swartztrauber and Fujikawa, 1998; Tulen et al., 1999). Hence, from a social cognitive point of view, blink rates have the potential to give clues about someone else’s inner mental life, or indeed their psychiatric well-being. Therefore, being aware of another’s blinks as well as other non-verbal behaviors might provide information that might be useful during a social interaction.

Thus far, virtually no attention has been paid to the neural consequences of observed blinks. Intuitively, it would seem that we do not notice every time someone blinks, yet we do notice when someone does not blink, or blinks very frequently. This suggests that at some level the brain does monitor the blinks of others (Omori et al., 1997). Results from behavioral studies support the idea that observed blinks are not ignored. Very frequent blinkers are more likely to be rated as nervous or careless, whereas both very infrequent and frequent blinks are rated as “unfriendly” (Omori and Miyata, 2001). In addition, political candidates with low blink rates are rated less favorably than those with normal blink frequencies (Patterson et al., 1992). Reduced blink rate has also been linked with deception, such that criminals who lied under interrogation had lower blink rates than those who told the truth, as verified by forensic and witness evidence (Mann et al., 2002). Another study found that blink rate decreased during lying but increased considerably immediately following the lie (Leal and Vrij, 2008). The relationship of blink rate and lying may be linked to cognitive demand as it has been shown that higher demand results in less frequent blinking (Drew, 1951), followed by an increase (or flutter) when the demand is lifted (Holland and Tarlow, 1972). Overall, these studies suggest that observed blinks may have social importance and can influence higher-order processes such as making judgments about mental states or forming impressions about others.
Changes in gaze are known to elicit reliable neural responses, even when the stimulus has no overt social significance (Puce et al., 2000). The elicited event-related potentials (ERPs) mainly consist of an N170 component that can vary its amplitude as a function of whether gaze is direct or averted (Puce et al., 2000, 2003; Conty et al., 2007), followed by later ERP components that have been shown to be modulated by social attention or social significance (Sabbagh et al., 2004; Carrick et al., 2007; Conty et al., 2007). Previous studies using static faces have shown significantly greater N170 and later ERP amplitudes for stimuli with emotional or social meaning including depictions of intense emotion and personally familiar faces compared with neutral expressions or unfamiliar persons (Caharel et al., 2002; Pizzagalli et al., 2002; Eger et al., 2003).

Here we recorded neural responses elicited to viewing blinks and eye movements of other individuals. Specifically, we studied viewing eye blinks, eye closure, as well as upward and leftward gaze shifts [in previous studies no differences to viewing right and leftward gaze had been observed (Puce et al., 2000)]. As in a naturalistic situation, the blink had an extremely short duration, whereas all other eye movement conditions including eye closure were longer and matched in duration. We formulated two hypotheses relating to the ERP results of the study:

H1. That N170 and later ERP components elicited to blinks would be robust and reliable. If this prediction were to be true, it would suggest that the brain can register and process all types of eye changes for potential social relevance.

H2. That there would be a graded change in N170 and later ERP amplitude as a function of condition in the following order: averted eyes (leftward or upward), eye closure, blinks. Larger and earlier N170s and subsequent ERPs would be seen in the former conditions as they are more likely to signal social or attentional significance than the latter.

We did not expect to observe any differences in P100 amplitudes (or latency) across conditions, based on literature indicating that this ERP component is modulated by stimulus attributes such as contrast, luminance, and spatial frequency, and does not systematically vary as a function of facial attributes or movement (in particular, see a discussion of this issue in Rossion and Caharel, 2011).

MATERIALS AND METHODS

PARTICIPANTS

Data from 14 volunteer participants between 18 and 29 years of age (mean age 21.6 years; six females; all right-handed) were included in this study. Handedness was confirmed using the Edinburgh Handedness Inventory (mean handedness quotient of 88R). All participants had normal or corrected-to-normal hearing and vision and reported no previous history of neuropsychiatric disorders. ERP data from an additional four participants were collected, but were excluded due to excessive and persistent movement or blink artifacts. Informed consent was acquired for the experiment, as approved by the Institutional Review Board of West Virginia University.

STIMULI

Stimuli consisted of gray scale video clips of a full-on face with a neutral expression and eyes looking directly at the observer. The eyes would randomly change to blink, to gaze upward and back, to gaze leftward and back, or make a duration-matched (with gaze conditions) eye closure (Figure 1). The direct gaze face was present for the entire experimental run (see also Puce et al., 2000), such that it appeared the face was looking at the observer and periodically blinking, closing the eyes, or looking up or away, always returning to the baseline direct gaze state. These apparent movements made up the four eye conditions: Blink, Eye Closure, Leftward Gaze, and Upward Gaze. All gaze aversion conditions (Eye Closure, Leftward and Upward gaze) had equal durations (100 ms) and were designed to look natural, whereas the eye blink was very brief (33 ms). The direct gaze face (no blank screen, or scrambled stimulus) remained on the screen between trials, with an interstimulus interval of 2 s between each eye change. There were two experimental runs of 4 min each, one with a male face and the other with a female face, and presentation order was counterbalanced across participants. Both faces were unfamiliar to all participants. To encourage participants to maintain their attention on the stimulus display, an additional target detection condition was included. Participants were required to detect a transparent checkerboard pattern briefly superimposed on the direct gaze face (target stimulus) and make a button press response each time it appeared. There were a total of 40 trials for each eye movement condition, as well as 40 target trials (half of these occurred with the female face, the other half with the male face).

All video stimuli were presented using Presentation software (version 11; Neurobehavioral Systems, CA, USA). Video clips of the faces were dynamic and featured apparent motion of the eyes, which maintained their end positions for 100 ms (Leftward gaze, Upward gaze, and Eye Closure) or 33 ms (Blink) before returning to the baseline direct gaze condition. To maintain consistent stimulus timing, target duration (superimposed checkerboard pattern) was also equal to the duration of the presented eye change stimulus. Although there were only three video frame changes from direct gaze to eye change back to direct gaze in this apparent motion stimulus, as in a previous study (Puce et al., 2000), all participants
described the eye movements as realistic. The stimulus faces were projected on a white screen placed directly in front of the participant at a distance of 195 cm, and subtended a visual angle of 14.3° × 16.7° (horizontal × vertical). The two eyes in each face subtended a visual angle of 3.4° × 0.73°. Stimulus condition order was randomized across trials.

**DATA COLLECTION**

Each participant was tested individually in a quiet, dimly lit room, while seated in an armchair. Visual stimuli were projected onto a white screen in front of the participant rather than viewed from a computer monitor to improve visibility of the facial movements and make the face seem more realistic (projected dimensions were approximately life size). Each participant completed two experiments in one ERP session in randomized order. Participants were given a short rest period (several minutes) between experiments. The second experiment had an unrelated purpose and will not be reported here. Before the experiment, participants were instructed to minimize their own eye and face movements, to keep their gaze centered on the bridge of the nose of the stimulus face, and to press the response button to targets as quickly and accurately as possible.

**ERP RECORDING**

The electroencephalogram (EEG) was recorded from participants using a 128 channel Neuroscan Electrocap with sintered Ag/AgCl electrodes, with a midline frontal ground and dual reference electrodes placed on either side of the nose. Four electro-oculographic (EOG) electrodes captured horizontal and vertical EOG components (Figure A1 in Appendix). EEG and EOG signals were recorded with a gain of 5000, sampled at 250 Hz/channels, and band pass filtered at 0.1–100 Hz. A continuous EEG recording was made during each experimental run and included a digital time stamp to indicate the onset of each stimulus condition.

At the completion of the recording session, electrode locations were digitized with a Polhemus 3 Space Fastrak digitizer (see Appendix).

**ERP ANALYSIS**

Electroencephalogram data were segmented offline into epochs of 1100 ms, which included a 100 ms pre-stimulus baseline. A semi-automated artifact detection routine identified participant eye blinks in segmented EEG epochs, and data were also visually inspected for additional electromyographic artifacts. Artifactual EEG epochs were excluded from subsequent analyses. An average of 10.6 out of 40 trials were excluded per participant per condition. There were no significant differences in artifact removal between conditions (average for viewing Blink 10.6, Eye Closure 10.7, Upward gaze 9.6, Leftward gaze 11.2 trials). Segmented EEG epochs were low pass filtered using a filter cut-off of 40 Hz, then baseline corrected with respect to the pre-stimulus baseline, and averaged as a function of each stimulus condition for each participant. ERPs elicited in response to target stimuli were excluded from further analyses.

The averaged ERPs for each participant were used to create grand average ERPs for each stimulus type across the participant group. Potential latency ranges for a subsequent semi-automated peak identification analysis were defined for each ERP component from the grand averaged ERPs waveforms. Peak latencies and amplitudes for all stimulus types in individual participants and grand averaged ERPs were identified within those pre-selected time intervals. From the average ERPs of each individual participant, ERP component peak latency, and amplitude measures were calculated for each condition.

Topographical voltage maps and Laplacians (second spatial derivative of the voltage map; Puce et al., 2007) of the grand average ERPs at ERP component peaks were created using EMSE software (Source Signal Systems, CA, USA). Based on inspection of topographic voltage maps, clusters of electrodes were selected whose mean ERP amplitude and latency values were calculated for P100, N170, and amplitudes for later ERP components. Data for P100 and N170 components were entered into two-way repeated measures analysis of variance (ANOVA) using SPSS V15 to test for Condition and Hemisphere. Amplitude data for later ERP components were analyzed using a repeated measures ANOVA for Condition.

**RESULTS**

**BEHAVIORAL DATA**

Participants indicated by keyboard press when they detected a transparent black and white checkerboard overlaid on the direct gaze face. The mean reaction time for target detection was 501.0 ms (SD = 138.6 ms), with 95.1% accuracy.

**ERP DATA**

All eye change conditions, including the viewed eye blink, produced a robust ERP complex consisting of a P100 and N170 seen mainly at temporo-occipital electrodes (e.g., centered on equivalent 10–10 system sites P07 and P08). Additionally, large later ERP components, were also observed to all stimulus conditions, and these later ERPs tended to be more extensively distributed across the scalp (see white markers on topographic maps in Figure 3). We discuss each ERP component below.

**EARLY ERP COMPONENTS AT TEMPORO-OCCIPITAL ELECTRODES**

**N170**

As we predicted (H1), the Blink condition produced a large and reliable N170 (see Figure 2A); however, all eye conditions produced early components of seemingly equal amplitude (Figure 2A, right temporo-occipital scalp waveforms for Blink and Leftward gaze are shown as examples) over the temporo-occipital scalp, contrary to our hypothesis (H2). The N170 was greater in spatial extent for the right hemisphere. Our second prediction (H2), that we would observe a gradient in amplitude as function of stimulus condition (Leftward and Upward, Eye Closure, Blink) was not sustained for the N170 (see Figures 2B,C histograms). Statistical testing of N170 amplitude differences between conditions via a two-way repeated measures ANOVA for Condition and Hemisphere, was performed on the N170 peak amplitude from an averaged ERP from a cluster of six temporo-occipital electrodes in each hemisphere (see Figures 2B,C white dots; and see Puce et al., 2003; Carrick et al., 2007). There was no main effect of stimulus Condition (Table 1), however, there was a significant main effect of Hemisphere. Comparisons indicated that the right hemisphere generated larger and slower N170s in all conditions (Figure 2C;
Table 1). Similar to N170 amplitudes, a statistical analysis of N170 latency revealed a main effect of Hemisphere, but no main effect of Condition, and no significant interaction effects (data not shown).

**P100**
A two-way ANOVA revealed no main effect or interaction effect of Condition or Hemisphere for P100 amplitude at temporally-occipital electrodes ([Figure 2B](#)). There was no significant main effect of latency, nor were there any significant interaction effects.

**LATE ERP COMPONENTS**
To test the hypothesis that late ERP components would show an amplitude gradient based on putative social significance, we examined later ERPs at clusters of electrodes showing the greatest amplitudes. A series of positive late ERPs were observed, occurring at around 250, 450, and 600 ms post-stimulus, i.e., P250, P450, P600. To facilitate distinction of spatial distributions of the later ERPs, **Figure 3** displays Upward and Leftward gaze topographic voltage maps with higher voltage thresholds than those for Eye Closure and Blink ([Figures 3A–C](#); for Leftward gaze at same threshold see **Figure A2** in Appendix). In terms of spatial distribution and extent, the scalp distributions of the later ERP components showed some differences between the Blink and the other conditions (e.g., **Figure 3**). We discuss each of these ERP components separately.

**P250**
A clear P250 (mean latency 263 ms), which generally peaked over the vertex ([Figure 3A](#), e.g., Leftward gaze condition), was elicited by all four eye movement conditions, but was more posterior (centered closer to Pz) for the Blink condition relative to the other conditions ([Figure 3A](#), compare to Leftward gaze condition). A one-way ANOVA (Condition) of peak amplitudes from a five electrode cluster centered on Cz showed a main effect of Condition. This effect was driven by the Blink and Eye Closure conditions which both showed significantly smaller amplitudes than either the Upward or Leftward gaze conditions ([Table 1](#)). Note, however, that the Blink and Eye Closure conditions did produce

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no significant differences in the two-way ANOVA for Condition or Hemisphere for either the anterior temporal or posterior temporal electrode cluster.

**P600**

On average, all of the eye movement conditions also produced a broad positivity at around 600 ms (P600, mean latency 623 ms). Again there was a difference in the location of the maximal amplitude for this component, with Leftward gaze showing a more anterior focus compared to Blink (see Figure 3C; no difference in Condition effect for Pz vs. Cz electrodes, data not shown). At a nine electrode cluster centered on parietal midline electrode Pz, there was a main effect of Condition with P600s in response to Leftward gaze having significantly greater amplitudes than the three other conditions (Table 1).

**DURATION OF SIGNIFICANTLY DIFFERENT NEURAL ACTIVITY AS A FUNCTION OF TIME**

Finally, we examined the data of individual participants for potential differences in the duration of significant activation across the Blink and Leftward gaze conditions. We did this over multiple significant peaks at this latency (Figure 3, charts on right). An ANOVA for P250 latency indicated that there were no significant effects across condition.

**P450**

All four eye movement conditions produced a significant P450 (mean latency 459 ms). P450 was seen over the anterior temporal scalp and tended to be right lateralized for most conditions, including the Blink. The Leftward gaze aversion showed a more posterior temporal distribution than the Blink. We investigated this distributional difference further by plotting topographic Laplacian maps at the peak of P450 activity. For all conditions, there appeared to be a prominent sink and source of activity on the midline and the right posterior temporal scalp, respectively (Figure 4, bottom row). The Laplacian map for the Blink condition showed these same loci and an additional sink over the occipital scalp. This may explain the apparent hemispheric asymmetry in the topographic voltage maps for this condition. In the left hemisphere a sink of activity was also observed in the temporal scalp with the Eye Closure, Upward, and Left conditions, and in the fronto-temporal scalp for the Blink condition.

Given the observed distribution differences in P450 across conditions, we performed statistical analyses for two electrode clusters centered on both these amplitude maxima (anterior temporal and posterior temporal). A two-way ANOVA for Condition and Hemisphere of the posterior temporal electrode amplitudes (white circles in Figure 3B), revealed a significant effect of Condition with Leftward gaze having a significantly greater amplitude than the three other conditions (Table 1). There was no main effect of Hemisphere and no interaction effect. On the other hand, the anterior temporal electrodes (gray stars in Figure 3B), showed main effects of Condition and Hemisphere (Table 1). The Condition effect was driven by larger amplitudes for the gaze conditions. The Hemisphere effect was driven by significantly greater amplitudes in the right vs. left hemisphere for Blink, Eye Closure, and Upward gaze conditions (all comparisons included in Table 1). For P450 latency, no significant differences in the two-way ANOVA for Condition or Hemisphere for either the anterior temporal or posterior temporal electrode cluster.
for Leftward gaze, the condition with the most potential social and attentional meaning, had significantly greater amplitudes than all the other conditions.

**DISCUSSION**

N170 IS COMPARABLE FOR ALL TESTED EYE MOVEMENT CONDITIONS

Our second hypothesis that the Blink condition would elicit reduced N170 amplitudes, compared to the other conditions, was not sustained; robust N170s were elicited by all conditions including Blink. Similarly, no significant differences were seen in N170 latency as a function of Condition. We had previously described significant differences in N170 amplitudes and latencies as a function of eye gaze direction – with larger N170 observed to gaze aversions relative to when eyes look directly at the observer (Puce et al., 2000, 2003). Type of gaze aversion (left, right, upward) does not appear to influence N170 amplitude, irrespective of whether an explicit social context or social judgment is required or not. As expected our tested gaze aversions (Leftward, Upward) also showed no significant differences in N170 amplitude or latency, however, in this study we used transient aversions of around 100 ms in an attempt to move toward more ecologically valid stimulus displays. Here we also tested Eye Closure, which might be regarded as a form of eye closure stimulus, using a similar stimulus duration of 100 ms, in addition to a Blink – a quicker type of eye closure stimulus. In these latter two stimulus conditions, N170 amplitudes were as large as for the gaze aversions, indicating that the brain registers and monitors all types of eye changes. The N170 has been previously suggested to potentially be sensitive to changes in the social attention of another individual (Itier and Batty, 2009), but the results in this study would

- **FIGURE 5 | Return to Baseline (RTB) times in individual participants.** Blink (top row) and Leftward gaze (bottom row) condition data are shown for each participant as bars 1–14. The RTB times indicate a permanent RTB time in ms. The gray bar for each participant extends for the duration of time in which significant peaks (vs. baseline) are observed (timeline on y-axis). The criterion for a waveform to be significantly different from zero was five consecutive time points (20 ms) with a t-value greater than 1.68 (based on a df = 40, p < 0.05). All participants had significant early peaks at temporo-occipital electrodes (up to ~200 ms, see gray shaded region for PO8), but not all participants had later peaks, or peaks at more anterior electrodes, especially for the Blink condition. The final bar indicates a group average RTB with SE. RTB values are shown for electrodes analogous to 10–10 system sites PO8, Cz, Pz, and Fpz.

- **FIGURE 5**
suggest that N170 might more likely reflect a detector of change in the eyes, or perhaps changes in the face in general (see also Puce et al., 2000).

There was an effect of Hemisphere for the N170, with the right hemisphere showing significantly greater amplitude than the left hemisphere in all four conditions. In our own previous studies, N170s elicited to viewing lateral gaze shifts to both the left and right were of equal magnitude, with a tendency for larger N170s to be seen in the right hemisphere regardless of the gaze shift direction (Puce et al., 2000). The right hemisphere preference for face related stimuli, be they the onset of static faces or dynamic faces in ongoing facial displays, has been observed in studies by other laboratories (Rossion et al., 1999; Itier and Batty, 2009). The existing literature, which is devoted largely to examining N170s elicited by static face onset, shows mixed findings with respect to facial changes related to affect or social significance. For example, some studies showed larger N170 amplitudes to static fearful vs. neutral face onset (Pizzagalli et al., 2002; Eger et al., 2003), while others showed no differences (Eimer and Holmes, 2002).

We also examined the behavior of the P100, an ERP component which is thought to reflect low-level stimulus attributes in faces as well as other stimulus categories (see Rossion and Caharel, 2011). We did not observe any differences in P100 attributes as a function of stimulus Condition. Our data indicate that differentiation of neural activity as a function of taking the eyes away from another only occurs at latencies beyond 250 ms (see below). It is our belief that these earlier ERPs signal the change in the facial display (P100, N170), and potentially socially salient stimuli relative to the observer (N170). This latter signal may encourage further attention to a stimulus for potential social or attentional relevance.

**LATE ERP SIGNALS**

Average amplitudes of late ERP components were greater for Leftward and Upward gaze aversions compared to Blinks and Eye Closure, although clear and discernable late peaks were observed even in these latter two conditions. This was true for P250, P450, and P600, supporting our hypothesis that late potential amplitudes may reflect attentional and/or social significance, as seen in a previous study, where social context was explicitly manipulated (Carrick et al., 2007). Traditionally, later ERPs have been associated with higher-order cognitive and affective processes (Sutton et al., 1967; Kutas et al., 1977; Kutas and Hillyard, 1980; Hillyard and Kutas, 1983; Rosler et al., 1986), and more recently have been described in social cognition and theory of mind tasks (Liu et al., 2004; Sabbagh et al., 2004; Puce et al., 2007). Therefore the later responses observed in this study may indicate that simple eye movements, including blinks, have some social importance to us.

Event-related potential studies using static face onsets suggest that P250 may be related to consolidating configurational information of faces (Halit et al., 2000; Milivojevic et al., 2003), while ERPs beyond 350 ms may be related to assessment of stimuli, social cognition, memory, and attentional processes (Cacioppo et al., 1993; Sabbagh et al., 2004). In addition, later ERP amplitudes are known to increase with attentional or emotional salience (Talsma and Woldorff, 2005; Schupp et al., 2007). A recent study from our lab using dynamic facial displays showed differences in later ERPs to averted gaze when embedded in multiface displays where an explicit judgment of social significance is required (Carrick et al., 2007). Even in the present study in which participants did not make explicit social judgments, these changes in later ERPs were robust and reliable. Perhaps this is because, in everyday social interactions, seeing an upward or lateral gaze shift is usually a strong indicator that the attention of the observed person is being directed away from the observer, conceivably toward important external people or stimuli, or as a result of social discomfort (Kleinke, 1986).

It is possible that as we observe the eye movements of another, including blinks, we may activate an embodied response within the oculomotor system. Oscillatory changes for self-blinks in occipital regions at 200 ms and in orbital regions at 400 ms have been documented with MEG (Bardouille et al., 2006). Furthermore, fMRI studies of voluntary self-blinks show activation in visual and orbital frontal cortices (Tsubota et al., 1999; Yoon et al., 2005). For self-blinks, however, these changes might also reflect a transient interruption of input to visual cortex. Activation in different cortical structures may help explain the distributional differences observed in later ERP components across conditions. Examples are the more posteriorly distributed P250 and P600, and more anterior and right lateralized P450 for the Eye Closure vs. Leftward and Upward gaze change conditions. Future studies using neural source modeling might be able to shed some light on these differences.

**FUTURE DIRECTIONS**

Given that we appear to be aware of the blink rates of others (Omori and Miyata, 2001; Mann et al., 2002) and that social perception is known to be influenced by blink rates of the observed person, a study in which another’s blinking behavior was explicitly associated with particular social attributes might be a way to systematically investigate how late ERPs are modulated by these social attributes. Correlating the degree of higher-order blink processing with blink rate (normal, fast, or slow) and personality assessments would be useful in understanding how non-verbal cues can influence our judgments. A tantalizing possibility for why blinks result in higher-order cognitive processing is that we have learned associations between blink rate and level of interest, arousal level, stress, and/or other factors integral to social interactions. These cognitive and affective states can in turn affect dopaminergic systems which have a direct relationship with blink rate (Brozoski et al., 1979; Carson, 1983; Coull et al., 1995; Coull, 1998; Hirokawa et al., 2000; Dreisbach et al., 2005).

Reflexive attentional shifts that result from observed non-predictive (and sometimes anti-predictive) eye gaze changes (Friesen et al., 2004; Schuller and Rossion, 2005) might potentially drive the observed differences in later ERPs in our study. For example, in the current study, if objects or another face or person were located above or to the left of the face, attention may have shifted to these objects/people causing differences in attentional focus, ERPs, and distraction level (Fichtenholtz et al., 2007). Adding affective expressions to the gaze changes would likely create even more complex attentional effects, which might be influenced by such factors if the depicted emotion encourages approach or withdrawal on the part of the observer (Fichtenholtz et al., 2007, 2009).
CAVEATS
Our task used an apparent motion task with two-dimensional real human faces and near-realistic timing. However, there can be differences in neural responses to viewing our stimuli vs. observing live interactions (Ponkanen et al., 2008). The higher-order response to blinks observed in our study could conceivably occur as a result of the artificial and isolated nature of our computer-presented stimuli. It is possible that in this relatively impoverished laboratory environment, less important stimuli may be processed to a greater extent than they would normally be in a natural environment. In addition, multiple repetitions of the limited stimulus set may have led to habituation, which could have in turn have led to reduced higher-order processing of all eye movements. However, we believe this to be unlikely, as significant differences between conditions were seen in later ERPs and not the early components.

Our sample size (14 subjects) may have resulted in the absence of significant differences in the N170 amplitudes across conditions, although in our previous studies using this sample size, significant differences in N170 amplitudes were observed when social attention was manipulated.

Lastly, there was a stimulus duration difference between the Blink condition and the other conditions. However, the Eye Closure condition was matched in duration with the Leftward and Upward gaze conditions. Eye Closure did not show significant differences from Blink, and, like the Blink condition, was often significantly smaller than the Leftward gaze condition. However, a shorter gaze change condition may be necessary to fully address this duration difference.

CONCLUSION
Behavioral studies indicate that we form social perceptions about others from observing their behaviors, which would include monitoring blinking behavior. Our ERP data show that blinks are registered similarly to gaze aversions, the latter of which are known to be important for gauging the social attention of another. Importantly, later ERP components were sensitive to the potential social/attentional significance of the stimuli and displayed a social relevance related response gradient. Our work demonstrates that blinks in and of themselves may prove to be interesting and important cognitive events.

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APPENDIX

EEG RECORDING
Overall, 128 channel EEG and EOG signals were amplified by 5000 (band pass = 0.1–100 Hz) during the two experimental runs and were continuously recorded and stored on hard disk for offline analysis. Eye movements were recorded with additional electrodes that were placed at the outer canthus of each eye to record the horizontal electro-oculogram (hEOG), and above and below the left eye to record the vertical electro-oculogram (vEOG). Electrode impedances were kept generally below 15 kΩ.

ELECTRODE DIGITIZATION
Once the experiment was completed, electrode locations on the Neuroscan electrocap were digitized with a Polhemus 3 Space Fastrak digitizer. These coordinates were stored for each participant, and a grand average of the electrode locations was made by averaging the coordinates for all participants who were to be included in the experiment.

ERP ANALYSIS
The continuous EEG/EOG recording was first epoched into segments of 1100 ms, beginning 100 ms prior to stimulus onset. Artifact-contaminated EEG epochs were automatically rejected using an amplitude criterion of > ±75 μV (to eliminate EOG and large EOG or electromyographic artifacts caused by any eye and head muscle movements, respectively). The experimenters also visually inspected the accepted epochs, eliminating additional epochs with other artifacts or noise as necessary. All accepted epochs for each stimulus type were averaged as a function of stimulus condition for each individual subject. Target stimuli were not averaged. The averaged ERPs for each participant were individually digitally filtered (low pass, 30 Hz; zero phase shift; cut-off 48 dB/oct). All filtered stimulus type averages for each individual were then detrended across the entire epoch.

Filtered and detrended average ERPs for each participant were used to create grand average ERPs for each stimulus type. A semi-automated method was used to identify peak latencies and amplitudes, for all stimulus types for each individual and the grand average, within pre-determined intervals (outlined below). Topographical voltage maps of these grand average ERPs were created using EMSE software (Source Signal Systems, CA) for all conditions, although only maps for the Blink and Leftward conditions are shown in the figures since they typically differed the most from each other. Grand average ERPs and topographic voltage maps were visually examined at times corresponding with ERP peaks (P100: 108–168 ms; N170: 172–264 ms; P250: 200–300 ms; P450: 348–560 ms; P600: 540–720 ms).

At temporal–occipital electrodes, a two-way repeated measures ANOVA was conducted to examine amplitude and latency differences between Condition (Blink, Eye Closure, Upward gaze, Leftward gaze) and Hemisphere (Left vs. Right), using SPSS V15 software. Temporo-occipital electrodes used for statistical analyses were selected on the basis of inspection of topographical voltage maps, and were similar to those used in previous studies (Puce et al., 2000, 2003). Similarly, for other later ERPs, electrodes were selected based on location of the greatest amplitudes for each component. If lateralized, we would perform a two-way ANOVA, testing main effects for Condition and Hemisphere. For midline electrodes, a one-way ANOVA excluding hemisphere was performed. Significance level was set at p < 0.05.

FIGURE A1 | Average activity in electro-oculogram (EOG) electrodes.
Grand averages (n = 14) for each of the four EOG electrodes are shown relative to eye position. Note that there is persisting positive-going activity in the upper vertical electrode and no (negative-going) activity in the lower EOG electrode. The activity shows a similar time course, but smaller amplitude, in the lateral EOG electrodes. When slow activity is observed in frontal electrodes, there is always the concern that this may have resulted from eye movement or drift. In three of the four EOG electrodes there was a large positivity, with a gradient of activity. It was largest in the upper vertical, and was present albeit to a lesser extent in the bilateral horizontal eye channels. The time course of the positivity was identical to that observed in the frontal EEG channels. Importantly however, the lower vertical EEG channel, located below the right eye, did not show a reflected negative activity as it would for any vertical eye drifts, suggesting that observed activity on the other EOG electrodes represented EOG activity.

FIGURE A2 | Leftward gaze condition at the same threshold as Blink condition from Figure 3.
Leftward gaze results in much greater amplitude than the Blink condition. White circles identify posterior temporal, while stars denote anterior temporal electrode clusters whose ERP data were used in statistical analyses.
For each participant, waveforms from Blink and Leftward gaze were analyzed to test if each time point was significantly different from zero (Paired $t$-test vs. zero, Neuroscan Scan 4.3). For this, we included SD values based on the waveform variation on an epoch by epoch basis for each time point, with $N =$ number of epochs included after artifact removal (in our sample, 22–39 epochs after artifact rejection). We used these $t$-test waveforms to calculate return to baseline (RTB) values used in Figure 5. The RTB value is the point in time after which the waveform is not significantly different from zero. The criterion for a waveform to be significantly different from zero was five consecutive timepoints (20 ms) with a $t$-value greater than 1.68 (based on a df = 40, $p < 0.05$).