Neural correlates of correct and errant attentional selection revealed through N2pc and frontal eye field activity.

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The goal of this study was to obtain a better understanding of the physiological basis of errors of visual search. Previous research has shown that search errors occur when visual neurons in the frontal eye field (FEF) treat distractors as if they were targets. We replicated this finding during an inefficient form search and extended it by measuring simultaneously a macaque homologue of an event-related potential indexing the allocation of covert attention known as the m-N2pc. Based on recent work, we expected errors of selection in FEF to propagate to areas of extrastriate cortex responsible for allocating attention and implicated in the generation of the m-N2pc. Consistent with this prediction, we discovered that when FEF neurons selected a distractor instead of the search target, the m-N2pc shifted in the same, incorrect direction prior to the erroneous saccade. This suggests that such errors are due to a systematic misorienting of attention from the initial stages of visual processing. Our analyses also revealed distinct neural correlates of false alarms and guesses. These results demonstrate that errant gaze shifts during visual search arise from errant attentional processing.
Introduction

Comprehensive models of cognition must account for patterns of both correct and errant behavior. This has proven difficult because errors can arise in many ways, both through faulty sensory processing and through hasty response preparation. Still, behavioral measures of errant responding have been of interest for many years (Rabbitt 1966), and leading models of perceptual decision making aim to account for errant behavior along with correct responding (Ratcliff and Rouder 1998). Unfortunately, these efforts are limited by a paucity of evidence identifying the neural correlates of errors. Understanding the neural activity associated with errant performance provides a more complete picture of the neurophysiological basis of behavior and offers important constraints on cognitive models (e.g., Purcell et al., in press).

In this study we obtained new information about how errors of visual search occur by recording the macaque homolog of the human N2pc using electrodes embedded in the macaque skull (Woodman et al. 2007) simultaneously with single units and local field potential (LFP) in macaque frontal eye field (FEF), an area recognized as contributing to attentional selection on the one hand (Cohen et al. 2009a; Sato and Schall 2003; Schall et al. 1995a) and saccade production on the other (Bruce and Goldberg 1985; Hanes and Schall 1996; Ray et al. 2004; Schall 1991).

Previous research has demonstrated that the N2pc is a signature of covert attentional selection (Luck et al. 1993; Luck and Hillyard 1994) and can be used to monitor dynamic shifts of attention (Woodman and Luck 1999). The N2pc was
discovered when human subjects performed visual search for a lateralized target stimulus (Luck et al. 1993). In humans, the N2pc is observed as a greater negativity at electrode sites contralateral to the target stimulus, 175-200 ms after array onset. Importantly, the N2pc appears in tasks designed to prevent eye movements, occurs well before any manual responses are generated, and is even elicited by attended objects that do not require a response of any kind (Woodman and Luck 1999). Furthermore, because the N2pc is a lateralized component, a prerequisite for its emergence is a consistency in the direction of attentional orienting. If attention were directed in a haphazard fashion, no N2pc could be found because of the averaging across locations. Similarly, the N2pc does not emerge when conditions preclude attentional selection, such as when sensory input is data limited (Woodman and Luck 2003). Hence, the N2pc reflects consistent, task related movements of covert attention apart from any overt response. We recently discovered that macaque monkeys exhibit a homologue of the human N2pc (referred to henceforth as the m-N2pc) that demonstrates identical functional characteristics (Woodman et al. 2007).

This study is the first to examine whether the m-N2pc occurs during visual search errors, though FEF activity during such behavior has been well documented (Shen and Paré 2007; Thompson et al. 2005; Trageser et al. 2008). Briefly, FEF neurons incorrectly select distractor items prior to an errant saccade into the receptive field (RF). The equating of target selection in FEF with attention allocation (Armstrong et al. 2009; Kodaka et al. 1997; Sato and Schall 2003) requires that other indices of covert attention, such as the N2pc, mirror the
neural activity known to occur in FEF during visual search. While we have recently demonstrated this for correct trials (Cohen et al. 2009a), it remains to be seen whether the relationship will continue to hold when processing breaks down. The placement of FEF as an important contributor to attentional orienting suggests a similar FEF neuron – mN2pc mirroring on error trials. A demonstration that the m-N2pc tracks errant attentional orienting concordant with errant target selection in FEF would strengthen the hypothesis that FEF processing is closely related to the m-N2pc and thereby plays a role in covert orienting.

Methods

Behavioral tasks and recording. Two male macaques (macaca radiata) were trained to perform visual search for a form-defined target among similar distractors (Cohen et al. 2009b). The monkeys were also trained to perform a memory-guided saccade task for use in characterizing the response properties of FEF single units (Bruce and Goldberg 1985; Hikosaka and Wurtz 1983). During the memory-guided task, monkeys began a trial by fixating for 750-1000 ms. Then, a circular target was presented for 100 ms at one of 8 iso-eccentric locations. Eccentricity was adjusted based on the response properties of the neuron being recorded and target size was scaled with the cortical magnification factor. Following a delay of 500 – 1000 ms, monkeys made a saccade to the remembered location of the now-absent target. Monkeys were rewarded for fixating the remembered location for 1000 ms and were not rewarded when
saccades were either not produced or made to an incorrect location. In the visual search task, monkeys were shown an iso-eccentric, circular array of rotated T and L shapes (see Figure 1A). On a given session, either a T or L of a specific orientation was defined as the target. Distractor items were drawn from the non-target set. The number of distractors presented on each trial was 1, 3, or 7 (leading to set sizes of 2, 4, and 8) and was determined randomly. Each trial began with monkeys fixating a central point for 750-1000 ms. To earn liquid reward, monkeys had to make one saccade directly to the target location and hold that position for 1000 ms.

*Direction errors* were those trials in which the monkey made a valid eye movement to a screen location containing a distractor when a target had been present. Target-present errors of other types (e.g., lack of any eye movement within 2000 ms of target appearance, failure to maintain fixation in the target window for 1000 ms, eye movements to empty screen locations) were rare and were not analyzed further. Some sessions included catch trials where all display elements were distractor items. The proportion of catch trials was between 10-30% on these days. To earn reward, monkeys had to maintain central fixation for 750 ms. Thus, the second type of error was a *catch error* in which an eye movement of any type was made. In some sessions, the catch trial display immediately disappeared at 750 ms when reward was delivered while in other sessions, the displays remained visible for an additional 500 ms to eliminate any possibility for contamination due to an offset response. This had no effect on behavioral or neural data and patterns were identical across the two schemes.
Our analysis of FEF activity and the m-N2pc contrasted activity on correct trials with direction errors in which gaze shifted opposite to the actual target location (Figure 1A). This selection of error trajectories allowed us to use the neuron-antineuron approach to quantify the target selection process (Britten et al. 1992; Thompson et al. 1996).

During aseptic surgery, monkeys were implanted with a head post, recording chambers, electroencephalogram (EEG) electrodes, and a subconjunctival eye coil under isoflurane anesthesia. Antibiotics and analgesics were administered postoperatively. All surgical and experimental procedures were in accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals and approved by the Vanderbilt Institutional Animal Care and Use Committee.

Neurons and LFPs were recorded from FEF, located in the rostral bank of the arcuate sulcus. FEF was identified by evoking saccades with < 50µA of microstimulation current. Neurons and LFPs were recorded simultaneously from both hemispheres using tungsten microelectrodes (2-4 MΩ, FHC, Inc.) and were referenced to a guide tube in contact with the dura. The EEG and the averaged event-related potentials (ERPs) were recorded from the monkeys using an array of electrodes implanted in the exterior mantle of the skull (Woodman et al. 2007). Spikes were sampled at 40 kHz; LFP and EEG signals were sampled at 1 kHz. LFP and EEG signals were acquired in one of two ways. About half of the data for monkey Q was band-pass filtered between 0.7 and 170 Hz, amplified using a Plexon, Inc. HST/8o50- G20 head-stage. The other half of monkey Q’s data and
nearly all of Monkey S’s data were band-pass filtered between 0.2 and 300 Hz, amplified using a Plexon, Inc. high impedance HST/8050-G1 head-stage. The differences in head-stage may account for the slight difference in visually-evoked LFP polarization between the monkeys, as low impedance head-stage are known to create distortions in LFPs (Nelson et al. 2008).

For each isolated neuron, we mapped the RF using activity during the memory-guided task. Cells were classified as visual or visuomovement (pure movement cells were not analyzed). Visual neurons had above-baseline activity in the 100 ms following stimulus onset but no modulation before a saccade while visuomovement neurons exhibited both a visual response and buildup activity before a saccade. LFP and EEG signals were assigned an RF based on hemisphere. That is, signals recorded in the right hemisphere were assigned left hemifield RFs, and vice versa. LFPs and EEGs were included only if they exhibited statistically significant selectivity across hemifields. Using a more restricted RF for the LFP and EEG, based on the concurrently recorded neuron’s RF, did not change the data or conclusions qualitatively. We assessed selectivity by comparing activity on correct trials when a target fell in the RF (RF$_{\text{correct}}^T$) to that when a distractor appeared there (RF$_{\text{correct}}^D$). Running Wilcoxon rank-sum tests were computed ms by ms, and a signal was categorized as selective if it reached significance for 10 consecutive ms at the 0.05 level. We verified this by computing selectivity using receiver-operating curves (ROC). Each point on an ROC curve reflects the probability that RF$_{\text{correct}}^T$ activity is greater than some criterion as a function of the probability that RF$_{\text{correct}}^D$ activity is greater than that
criterion. In the case of single units, the criterion was incremented from 0 spikes/second to the maximum firing rate observed across all trials in steps of 1 spike/sec. For LFP and EEG signals, the criterion incremented from the smallest voltage observed to the largest voltage observed in steps of 10 µV. The area under this curve, at a given time point, can be interpreted as the ability of an ideal observer to determine whether the target was in the RF given only the neural data. If the ROC area reached .70, the signal considered selective. Importantly, the neurons identified as being selective using running Wilcoxon tests were the same as those identified using ROC analyses. This ensured that the neural signals we analyzed were significantly tuned for target identity. We imposed this constraint only on correct trials, as error trials had significantly lower signal-to-noise ratio. This did not present a problem for our analyses, as we were primarily interested in the direction of selectivity. While this undoubtedly included neurons with weak activity during error trials, doing so only made it more difficult for us to reject the null hypothesis.

ERP analyses were conducted with electrode pairs OL/OR and T5/T6 in the macaque analogue of the human 10/20 system (Jasper 1958; Woodman et al. 2007) for monkeys S and Q, respectively. A frontal electrode at position Fz was used as the EEG reference for both monkeys. We present only data from electrode OR and T6 in monkey S and Q, as skull electrode OL in monkey S demonstrated a curiously inverted P1/N1 polarity change after a frontal craniotomy and was unreliable. Importantly, all patterns of m-N2pc data were
verified using the left hemisphere T5 electrode of monkey Q, which were identical to those from the right hemisphere.

Data Processing. Spikes were convolved with a kernel resembling a post-synaptic potential (Thompson et al. 1996). LFPs and ERP data were processed as follows. Trials on which signals saturated the amplifier were excluded from the analyses. Signals were baseline corrected using the average voltage in a 100 ms window ending at the time of array presentation. EEG signals were truncated trial-by-trial 20 ms before the saccade to eliminate any influence of the saccade artifact. This tended to increase noise levels; therefore, ERP waveforms were filtered (50 Hz low-pass, zero-phase shift filter) for display purposes only. All statistics were computed on unfiltered data.

Results
We identified neuron, LFP, and ERP recordings that met the criteria described above. Monkey Q contributed 42 neurons, 31 LFPs, and 14 ERPs. Monkey S contributed 19 neurons, 46 LFPs, and 19 ERPs.

Behavior.
Figure 1A depicts a typical visual search session in which a T (rotated 90 degrees) was the target stimulus. The left panel shows saccade trajectories for the $RF^T_{\text{correct}}$ (when a target item fell in the RF and the monkey responded correctly) and $RF^D_{\text{correct}}$ (when a distractor fell in the RF and the monkey
responded correctly) conditions in a representative session. The right panel shows the opposing trial types (black traces), when a target fell in the RF and a saccade was made away (RF\textsuperscript{error}) as well as when a distractor fell in the RF and a saccade was made into the RF (RF\textsuperscript{error}).

Across the recording sessions, both monkeys demonstrated significantly slower RTs on error trials than on correct trials (Figure 1B; monkey S: \( t(35) = -2.47, p < 0.05 \); monkey Q: \( t(117) = -10.71, p < 0.001 \)), but the difference was larger for monkey Q (\( M_{\text{Correct}} = 318 \text{ ms}; M_{\text{Error}} = 431 \text{ ms} \)) than for monkey S (\( M_{\text{Correct}} = 267 \text{ ms}; M_{\text{Error}} = 279 \text{ ms} \)). Error rates for monkey Q were much lower (12%) than for Monkey S (35%), (\( U = 779, n_1 = 118, n_2 = 36, p < 0.001 \)). This pattern of RTs and accuracy suggest that the two monkeys may have been operating with different speed-accuracy tradeoff criteria (Lohman 1989). Most importantly, both RT and error rate tended to increase with search set size (Figure 1B). A long history of research implicates this ‘set size effect’ as a signature of attention, as it is absent in conditions allowing pre-attentive selection of target stimuli (Treisman and Gelade 1980). A statistically significant increase in RT with set size was observed for monkey Q (correct trials: \( F(1,105) = 902.38, p < .001 \); error trials: \( F(1,104) = 348.40, p < .001 \)) and for monkey S (correct trials: \( F(1,34) = 177.27, p < .001 \); error trials: \( F(1,34) = 146.90, p < .001 \)). The same linear increase with set size was observed for error rate (monkey Q: \( F(1,105) = 117.54, p < .001 \); monkey S: \( F(1,34) = 155.26, p < .001 \)). We have previously documented the relationship between neural activity and the set size effect (Cohen et al. 2009b) and so will not discuss this further here.
**N2pc and FEF signals during correct and error trials**

Figure 2 shows the average m-N2pc simultaneously recorded with a single FEF neuron and associated LFP in a representative session (same session shown in Figure 1A). On correct trials (black lines), all three signals selected the targets (thick lines) rather than distractors (thin lines) through characteristic modulation (Cohen et al. 2009a). The m-N2pc component exhibited a larger positivity for targets than for distractors in the RF, as previously shown (Cohen et al. 2009a; Woodman et al. 2007). Neurons expressed target selection by firing at a higher rate when target stimuli fell in the RF ($RF^T_{\text{correct}}$) relative to when distractors appeared there ($RF^D_{\text{correct}}$), whereas LFPs demonstrated a more negative potential for target stimuli. To compare this pattern to that recorded on error trials, we measured the three types of signal (m-N2pc, spikes, and LFPs) when distractor items fell in the RF but an erroneous saccade was made towards it ($RF^D_{\text{error}}$) and trials in which a target stimulus fell in the RF but a saccade was made away ($RF^T_{\text{error}}$). Consistent with previous analyses of unit activity during visual search for color singletons or color-shape conjunctions (Thompson et al. 2005), neurons demonstrated reversed selectivity on error trials. That is, when an errant saccade was to be made into the RF, neurons responded as if the distractor was actually a target.

We now report two new observations. First, the reversed pattern of polarization was observed in the FEF LFP on error trials relative to correct trials. Second, when monkeys made search errors, the m-N2pc also treated the
misidentified distractor as a target and the missed target as a distractor. That is,
the selectivity is inverted for error trials relative to correct trials, and this inversion
holds across all three kinds of signals. This has not been reported before in
either the literature on human N2pc or on monkey neurophysiology.

As shown in Figure 3, this pattern was replicated across the population in
both monkeys. To evaluate these effects statistically, we compared the average
firing rate or voltage within the period 150-250 ms after array appearance for RF^T
versus RF^D trials. Statistics are combined across monkeys because the effects
were indistinguishable. We carried out a mixed-model ANOVA with within-
session factors of response type (correct versus incorrect) and stimulus array
(RF^T versus RF^D) and the between-session factor of signal (neuron versus LFP
versus m-N2pc) because each session did not necessarily contribute to each of
the signals. As illustrated in Figure 3, the signals observed in RF^T trials and RF^D
trials were significantly different, and the pattern of selectivity reversed between
correct and error trials as evidenced by an interaction of response type X stimuli
array for m-N2pc (F(1,32) = 72.2, p < 0.001), neurons (F(1,60) = 74.0, p < 0.001),
and LFP (F(1,76) = 26.2, p < 0.001). Surprisingly, single unit and LFP activity
were sensitive to the presence of a target item in the RF even when an errant
saccade was made away from that location. For neurons, RF^T_{error} was significantly
greater than RF^D_{correct}, t(60) = -4.4, p < .001; for LFP, RF^T_{error} was significantly less
than RF^D_{correct}, t(76) = 2.9, p < .01.

Figure 4 illustrates the quantitative pattern of selectivity on correct and
error trials across the population. We subtracted the signal on RF^D trials from
that on RF\textsuperscript{T} trials when the response was correct (abscissa) and when it was in error (ordinate) in the period 150-250 ms following array appearance. Each data point represents the direction of selectivity for one signal from one session.

Signals that reversed polarity between correct and error trials will fall in the lower-right quadrant (m-N2pc and single units) and the upper-left quadrant (LFP). LFP data are opposite that of single units and m-N2pc because it demonstrates selection as a relative negativity rather than positivity. Signals that did not reverse between correct and error trials fall in the upper-right or lower-left quadrants. This illustrates the consistency of the reversal between correct and error trials across our sample. These findings show that tight linkage between the focus of selection by FEF activity and that of the m-N2pc is observed even when cognitive processing breaks down and the visual search task is performed incorrectly.

**Correlation of N2pc and FEF LFP**

Previously we showed that the amplitude of LFP polarization in FEF is correlated trial-by-trial with the amplitude of the m-N2pc on correct trials (Cohen et al. 2009a). We determined whether this relationship is present even when visual search was incorrectly performed by computing the trial-by-trial correlation between the integral of the FEF LFP and the integral of the m-N2pc from 100 ms after the array appeared until the saccade was initiated. Similar to our previous report, LFP and m-N2pc voltages were correlated for RF\textsuperscript{T}\textsubscript{correct} trials (median $r = 0.29$, $p < 0.05$) and RF\textsuperscript{D}\textsubscript{correct} trials (median $r = 0.33$, $p < 0.05$). Interestingly, this
correlation across brain areas and signal levels was also observed before errors, whether the error was made towards the RF ($RF_{\text{error}}^D$; median $r = 0.25$, $p < 0.05$) or away from it ($RF_{\text{error}}^T$; median $r = 0.30$, $p < 0.05$).

### Catch Trials

We have established that each of the neurophysiological signals we recorded selected a distractor stimulus prior to an errant response. We next investigated whether a relationship existed between neural activity and the type of error made. For this, we turned to sessions that included catch trials when no target appeared. To earn reward, monkeys were required to withhold making any response for at least 750 ms. We distinguished trials in which monkeys made no eye movements during the trial (correct catch) from trials in which the monkey made a saccade to a distractor before the 750 ms deadline (false alarms). We also analyzed trials in which monkeys maintained fixation through the 750 ms deadline (correct catch) but then made a saccade to a distractor item after receiving reinforcement (late catch response). Late catch responses were relatively common, amounting to 56% of all responses on catch trials (across the population, 18,557 / 33,123). Because we did not train the monkeys extensively on catch trials, correct catch trials were less common (17%, 5,694 / 33,123). Monkeys committed catch trial false alarms approximately 27% of the time (8872 / 33123).

Among these various catch trial outcomes we identified neurons, LFPs, and ERPs according to the same criteria described previously. These yielded 17
ERPs, 18 neurons and 13 LFPs. We then verified that saccade trajectories and landing points were similar between false alarms and late catch responses (Figure 5A, see also Figure 6B). Eye movements on correct catch trials were limited only to slow, drifting eye movements typically observed during the inter-trial interval. We contrasted neural activity preceding eye movements into the RF on catch trial false alarms and late catch responses as well as on target present trials when correct ($RF_{\text{correct}}^T$) and errant ($RF_{\text{correct}}^D$) saccades were produced into the receptive field. For single units in FEF (Figure 5B), the activity on false alarm trials was not statistically different from that on $RF_{\text{correct}}^T$ trials ($t(16) = 1.2$, $ns$). This finding replicates a previous report (Thompson et al. 2005). We now report an original observation; spike rate before late catch responses was significantly less than that on catch trial false alarm trials ($t(16) = -4.20$, $p < 0.001$). The spike rate was, though, elevated relative to that measured when a saccade was correctly made away from the receptive field ($RF_{\text{correct}}^D$) ($t(15) = 4.99$, $p < 0.001$). Activity on $RF_{\text{correct}}^D$ trials was not statistically distinguishable from that on correct catch trials ($t(16) = 0.90$, $ns$). To summarize, when monkeys committed a false alarm, the FEF visual neurons responded the same as they did during correct trials. This mimics the pattern of activity observed when erroneous saccades were made to distractors when the target was present. However, when the monkey made a self-generated saccade to a distractor location after receiving reward, neural activity was significantly attenuated. This occurred despite the fact that saccade metrics were closely matched between the conditions.
We determined whether this attenuation persisted through the initiation of the saccade by limiting our analysis to response-aligned visuomovement neurons, which demonstrate a rapid increase in firing rate leading up to a saccadic response. Because each of these categories of saccades have effectively identical trajectories and landing points, one might expect that presaccadic activity is generated whether the saccade was correct, a false alarm, or a late catch response. Surprisingly, the firing rate of the neurons was at the baseline level immediately before late saccades after successful catch trials in contrast to the pronounced level of activity before any other kind of saccade into the RF (Figure 5C). We verified that these neurons did in fact exhibit response-related activity during the memory-guided task. One representative neuron (Figure 6A) clearly demonstrated delay period activity following array onset, and a later buildup of activity around the time of saccade. This neuron was sensitive to target stimuli in the lower left quadrant. Saccades directed to target stimuli in the cell’s RF, and to distractor stimuli during catch trials were quite similar (Figure 6B). Most importantly, this neuron that responded before memory-guided saccades was virtually silent (perhaps even suppressed) before late catch saccades (Figure 6C).

Consistent with the hypothesis that the FEF LFP is a manifestation of local cortical processing during visual search (Cohen et al. 2009a; Monosov et al. 2008), LFP polarization on catch trials was similar to what we observed in the firing rates of single neurons (Figure 7). LFP polarization on false alarm trials was statistically indistinguishable from that on RF_correct trials but was significantly
more negative than that on $\text{RF}_{\text{correct}}^D$ trials ($t(12) = 2.6, p < 0.05$). The polarization on correct catch trials and late catch trial responses was not different. A different pattern was evident in the m-N2pc. The polarization in $\text{RF}_{\text{correct}}^T$ correct catch and late catch trials were statistically indistinguishable (Figure 7). Meanwhile, significantly greater positivity was measured on false alarm trials relative to the voltage on $\text{RF}_{\text{correct}}^T$ trials ($t(7) = -4.07, p < 0.01$), and the polarization on $\text{RF}_{\text{correct}}^D$ trials was significantly different from that in all other conditions ($F(1,7) = 44.62, p < 0.001$).

**Discussion**

*N2pc and FEF signals during visual search errors*

The N2pc is generated by areas in parietal and occipito-temporal cortex (Hopf et al. 2004; Hopf et al. 2000) that are reciprocally connected with FEF (Pouget et al. 2009; Schall et al. 1995b). Thus, the N2pc could be considered a signature of the signals received by FEF. However, the projections of FEF to extrastriate visual cortex (Pouget et al. 2009) provide for the possibility that FEF influences visual processing (Moore and Armstrong 2003). In a recent study, we demonstrated that visually responsive neurons in macaque frontal eye field (FEF) could be a source of the feedback to that generates the m-N2pc measured over posterior visual areas (Cohen et al. 2009a). Specifically, we found that the spiking activity and LFP in FEF selected visual search targets significantly earlier than did the m-N2pc. Thus, the timing and form of FEF activity when visual search is correctly performed are consistent with the hypothesis that signals from
FEF drive the extrastriate attentional selection mechanism manifest as the m-N2pc. This hypothesis is bolstered by the recent observation that the enhancement of neural activity due to attention occurs earlier in FEF neurons than in V4 neurons (Gregoriou et al. 2009; Ogawa and Komatsu 2006) as well as the demonstration that subthreshold microstimulation of FEF produces an enhancement of V4 neuron activity (Armstrong et al. 2006; Moore and Armstrong 2003).

Although the direction of influence cannot be determined by the present data, the results do strengthen the link between target selection in FEF and the covert orienting of attention. Previous studies reveal that single-unit responses in FEF and superior colliculus incorrectly select distractor stimuli prior to an errant response (Shen and Paré 2007; Thompson et al. 2005). That is, single units fire at a higher rate when distractor stimuli fall in the receptive field (RF) prior to an incorrect saccade to that distractor compared to when a correct response is made to a target outside the RF. We found that the macaque homolog of the human N2pc component mirrors activity in FEF even before errors. To our knowledge, such a pattern has never been reported in the human or nonhuman primate literature.

Alternatively, it was entirely possible that the m-N2pc would be absent during errant responses. Because the m-N2pc is a lateralized component in humans and nonhuman primates, it can only emerge under specific circumstances, and previous research has documented a number of task conditions that fail to elicit it (Luck and Hillyard 1994). If errors during visual
search entailed a nonsystematic orienting of attention to items across the visual field, the m-N2pc would not develop on error trials though present before correct responses. Similarly, if FEF activity did not reflect the orienting of attention, or if the error occurred during response mapping, one might expect to see accurate orienting of attention reflected in the m-N2pc despite the patterns exhibited by FEF. Indeed, when response mapping errors are more probable, FEF neurons correctly identify target stimuli both when report is correct and in error (Trageser et al. 2008). The present data make clear that errors during this visual search task stemmed from a systematic misorienting of attention and further support the identification of target selection in FEF with attention allocation.

Our analysis of the FEF LFPs provides further support for the relationship between the m-N2pc and FEF activity. First, like single unit responses, the FEF LFP incorrectly selected distractor stimuli prior to an error. This is perhaps not surprising as LFPs are thought to be a result of afferent post-synaptic potentials (Katzner et al. 2009; Mitzdorf 1985) and are related to local spiking (Fox and O'brien 1965). Second, the amplitude of FEF activity was correlated with m-N2pc amplitude on a trial-by-trial basis. While this can neither establish the direction of influence nor discount common input as a potential mediator, it is clear that the cortical areas responsible for the m-N2pc are related to activity local to FEF. Cortical inactivation studies will be useful in exploring these possibilities. For instance, the viewpoint that FEF drives the attention-related effects observed in extrastriate cortex predicts the absence of the m-N2pc following FEF inactivation.
Contingency of FEF signals

This work demonstrates a new dissociation in neural activity related to the type of error produced. In FEF (single units and LFPs), catch trial false alarm saccades to a distractor in the receptive field were associated with activity closely resembling that recorded on trials when the target appeared in a neurons’ receptive field ($RF^T_{\text{correct}}$). This is intuitive, as both trial types involve saccades into the same RF. However, when monkeys, after receiving reward for a successful catch trial, made an accurate but late self-initiated, visually-guided saccade to a distractor located in the RF, neural activity was significantly reduced. Unlike subcortical structures such as central thalamus (Schlag-Rey and Schlag 1984), cortical areas associated with eye movements (e.g., FEF, supplementary eye fields) exhibit little modulation during spontaneous saccades in the dark (Bruce and Goldberg 1985; Schall 1991). This is the first demonstration of a lack of FEF neural activity during seemingly purposive saccades. Surprisingly, visuomovement neuron activity aligned to saccade onset also demonstrated this pattern. In other words, neurons that are normally very active before a task-related saccade were silent when a saccade of the same direction was made after receiving reinforcement. It is possible that FEF may have been preparing a guess, and one signature of this may be the slight elevation of single unit activity prior to late catch responses (Figure 5B). Many classic models of decision formation assume that guesses are either preselected or evolve during a trial (Meyer et al. 1988; Ollman and Billington 1972), to be
enacted according to some internal deadline corresponding to the length of time
allowed for stimulus processing. Such strategies become useful when trials are
time-limited, as in the present task. Probabilistically, monkeys stand to receive
reinforcement at a higher rate by making a random guess than by withholding a
saccade altogether. Because a guess by definition would not depend on the
behavioral significance of any item, and because FEF neurons are sensitive to
context, it stands to reason that activity will be attenuated. Alternatively, this may
be a situation which dissociates attention and reward – constructs that are often
necessarily confounded (Maunsell 2004), since the attenuation in neural activity
was observed after reward delivery. However, as we cannot be sure of the
nature of attentional orienting preceding such responses, any further conclusions
would be speculative. In either case, it is interesting that FEF unit activity is
nearly absent during the same eye movements that otherwise produce strong
responses. These findings add to the evidence that the visual activity in FEF is
not directly related to saccade preparation or planning (Hanes et al. 1998; Juan

The current work supports the role of FEF in the production of covert
attention and further establishes a link between FEF and the cortical drivers of
the m-N2pc measure of selective attention employed in both monkeys and
humans. Further, these data make clear that errors during this visual search task
were systematic mis-deployments of attention rather than a breakdown in
attentional mechanisms per se. That the m-N2pc arises at all before behavioral
errors reflects a consistency of perceptual processing and attentional selection, even when that processing is flawed.
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Figure Legends

Figure 1. A: Saccade trajectories during visual search. Monkeys made saccades to a target $T$ shape in 4-element array. For display, target and distractor items are colored red. The RF for all signals in this session was directly to the left of the fixation point, indicated by the dashed semicircle, not to scale. Below each plot are the x-dimension eye traces recorded from the eye coil signal, converted to degrees visual angle. Gray traces are errors 180° from the target, black traces are correct trials to the target. B: Mean RT and error rate during the visual search task. Vertical bars represent $\pm$ 1 s.e.m.

Figure 2. Representative single-session averages for M-N2pc (top, electrode OR), single neuron from FEF (middle), and LFP from FEF (bottom).

Figure 3. Grand average m-N2pc (top), spike-density functions (middle), and LFP (bottom) for monkey S (left) and monkey Q (right). Traces are aligned to stimulus onset. Shaded region indicates the interval submitted to statistical test. Horizontal bars indicate times in which neural activation for $RF^T$ was significantly different from that for $RF^D$.

Figure 4. Magnitude and direction of selectivity on correct (abscissa) and error (ordinate) trials across the population. Each point represents the average difference between $RF^T$ and $RF^D$ activity over the period 150-250 ms post stimulus onset for a given signal and session. Points falling in the upper-right or
lower-left quadrants do not reverse polarity on correct and error trials. Those falling in the lower-right and upper left quadrants do reverse polarity (see text).

Figure 5. A: Saccade trajectories and x-dimension eye traces (converted to degrees visual angle) for one representative session. Left, saccade trajectories and landing points during false alarms (red) and late catch responses (blue) are identical. Right, eye traces during correct catch trials, where no eye movement was detected. Small, non-zero voltages are slow drifting eye-movements typically observed during the inter-trial interval. B: Grand average spike-density function on target-present and catch trials aligned to array onset. Shaded area indicates region submitted to statistical test. C: Grand average spike-density function on target-present and catch trials aligned to saccade initiation.

Figure 6. A: Memory-guided response profile for a typical visuomovement FEF neuron aligned to stimulus onset (left) and saccade onset (right). The neuron fired at elevated rates during the 1000 ms memory interval before increasing again prior to movement. B: Saccade trajectories for RF\textsuperscript{correct} trials (black), false alarms (red), and late catch responses (blue) into the RF of neuron in A. C: Response-aligned spike density functions for the same single neuron.

Figure 7. Grand average m-N2pc (top) and LFP (bottom) on correct target-present and catch trials. Shaded area indicates region submitted to statistical test.
References


Figure 1
Figure 2
Figure 3

Monkey S

Neuron (spikes/s)

Monk Q

RF

T

correct

RF

error

RF

D

correct

LFP (μV)

RF

D

error

Time from array (ms)

Figure 3
Figure 4
Figure 5
Figure 6

A. Memory-Guided Activity

B. Saccades into RF

C. Correct vs. False Alarm vs. Late Response during Memory-Guided Activity
Figure 7