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Priming of Attentional Selection in Macaque Visual Cortex: Feature-Based Facilitation and Location-Based Inhibition of Return

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Abstract

Visual search performance varies with stimulus and response history. Priming of pop-out refers to increased accuracy and reduced response time with repeated presentation of particular singleton and distractor features (e.g., a red target among green distractor stimuli), which are abruptly impaired when singleton and distractor features swap (e.g., green target among red distractors). Meanwhile, inhibition of return refers to the slowing of response time when target location repeats. Neurophysiological correlates of both these phenomena have been reported in the frontal eye field (FEF), an area in the frontal lobe contributing to attentional selection and eye movement planning. To understand the mechanistic origin of these adaptive behaviors, we investigated visual cortical area V4, an area providing input to and receiving feedback from FEF, during feature-based priming of pop-out and location-based inhibition of return. Performing a color pop-out task, monkeys exhibited pronounced priming of pop-out and inhibition of return. Neural spiking from V4 revealed earlier target selection associated with priming of pop-out and delayed selection associated with inhibition of return. These results demonstrate substantial involvement of extrastriate visual cortex in behavioral priming and inhibition of return.

Key words: attention; extrastriate cortex; pop-out; V4; vision

Significance Statement

Midlevel attention and visual processing is influenced by recent history of visual stimuli and gaze behavior. Using priming of pop-out visual search, we discovered that neural spiking in extrastriate visual area V4 shows speeded attentional selection when target and distractor features repeat and delayed selection when target location repeats. These neural processes paralleled but did not account for the magnitude of visual search performance changes with stimulus and response history. These new results improve our understanding of how recent experience influences attention and performance.

Introduction

Repetitive performance of cognitive tasks often yields changing behavior. Priming is a salient example of this observation (Tulving and Schacter, 1990). Repeated performance of the same task can lead to behavioral improvements such as in pop-out visual search, where a target that differs in a single visual attribute has to be detected among an array of distractors. The priming of pop-out yields speeded response times and greater accuracy (Maljkovic and Nakayama, 1994). This effect has been replicated in monkeys and extended to show changes in neural processing (Bichot and Schall, 1999, 2002; Purcell et al., 2012; Westerberg et al., 2020). Electrophysiological concomitants of the priming of pop-out have since been shown in humans (Eimer et al., 2010).
The behavioral benefits at the heart of priming of pop-out are based on repeating the target feature (i.e., feature-based facilitation of return). However, not all forms of repetition yield behavioral improvements. Repetition of the target location yields slowed response times while maintaining a similar level of accuracy (i.e., inhibition of return; Klein, 2000; Taylor and Klein, 2000; Bichot and Schall, 2002; Fecteau and Munoz, 2005), albeit not in all instances (Maljkovic and Nakayama, 1996; Fecteau and Munoz, 2003). The neural mechanisms producing these behavioral changes is a matter of debate (Kristjansson and Campana, 2010; Kristjansson and Asgeirsson, 2018). However, investigation into the neural sources of priming in visual search has provided crucial evidence.

The frontal eye field (FEF) and supplementary eye field (SEF) have been targets for neurophysiological investigations of priming in pop-out. While SEF neurons do not show modulation of activity with priming (Purcell et al., 2012; Westerberg et al., 2020), FEF neurons do (Bichot and Schall, 2002). Specifically, target selection (the time point where responses for the target stimulus diverge from that of distractors), measured in the activity of visuomovement neurons, is speeded with the priming of pop-out, while also being slowed following the repetition of target location. This result suggests that the priming of pop-out is based on the priming of attentional selection, which is also supported by studies in humans (Eimer et al., 2010; Brascamp et al., 2011).

FEF is not the only area implicated in priming of pop-out as lesions of V4 attenuate the behavioral improvements caused by priming (Walsh et al., 2000). FEF shares extensive connections with V4 (Schall et al., 1995; Stanton et al., 1995). V4 also shows robust modulation of activity with selective visual attention (Moran and Desimone, 1985; Reynolds and Heeger, 2009), and behavioral measures of selective attention are impaired following lesions of V4 (De Weerdt et al., 1999, 2003). Furthermore, frontal feedback to V4 influences attentional modulation in V4 (Armstrong et al., 2006; Armstrong and Moore, 2007), although it is not entirely necessary (Gregoriou et al., 2014). However, the diversity of information being fed forward into V4 is arguably less than FEF. V4 has been shown previously to lag FEF in selection of the behaviorally relevant stimuli in search (Ogawa and Komatsu, 2006). This suggests that V4 is necessary for the manifestation of priming in FEF by supplying input of relevant information (e.g., features) but does not necessarily represent priming itself. Altogether, there exists anatomic and functional evidence supporting the hypothesis that attentional selection of V4 neurons modulates with the priming of pop-out; however, the alternative, that it does not, is also feasible.

Here we test the hypothesis that priming affects attentional selection in area V4. To do so, we recorded neural activity in V4 of two monkeys performing pop-out visual searches with embedded featural-priming and positional-priming sequences. To evaluate the similarities and differences between FEF and V4 during this task, we performed analyses identical to those of the prior study in FEF (Bichot and Schall, 2002). Specifically, we investigated (1) whether V4 neurons show target enhancement and distractor suppression with feature priming, (2) whether V4 target selection times are speeded with feature priming, and (3) whether V4 shows slowed selection times with repetition of target position.

Materials and Methods

Animal care and surgical procedures

All procedures were performed in accordance with the National Institutes of Health Guidelines and the American Association for Laboratory Animal Care Guide for the Care and Use of Laboratory Animals, and was approved by the Vanderbilt Institutional Animal Care and Use Committee. Two male macaque monkeys (Macaca radiata; monkey C, 7.5 kg; monkey H, 7.3 kg) were implanted with a head post and recording chamber concurrent with a craniotomy positioned over area V4, which was located during a preoperative MR scan (Fig. 1A). Chamber placement was confirmed over V4 by a second, postoperative MR scan. Additionally, in one monkey, electrode penetrations were confirmed by staining two recording sites with diiodine before the animal was sacrificed (Fig. 1B). For chamber placement and craniotomies, monkeys were first tranquilized with ketamine (5–25 mg/kg) for intubation and catheterization before surgery. Surgeries were performed under aseptic conditions with the monkeys under and N2O/O2, isoflurane (1–5%) anesthesia mixture. ECG, temperature, and respiration were monitored continuously throughout the procedure. Expired Pco2 was maintained at ~4%. Postoperative antibiotics and analgesics were administered.

Anatomical MRI

Monkeys were anesthetized using the procedure outlined under Animal Care and Surgical Procedures. Anesthetized animals were then placed inside an Intera Achieva 3 T MRI scanner (Koninklijke Philips) at the Vanderbilt University Institute of Imaging Science and remained anesthetized throughout the duration of the scan. Vital signs were monitored continuously. T1-weighted three-dimensional MPRAGE scans were acquired with a 32-channel head coil equipped for SENSE imaging. Images were acquired using a 0.5 mm isotropic voxel resolution with the following parameters: repetition time, 5 s; echo time, 2.5 ms; and flip angle, 7°.

Experimental design and behavior

Monkeys were trained to perform a pop-out visual search task presented on a CRT monitor at 60 Hz, 57 cm
away, where the relevant feature was the color of a stimulus (Fig. 2A). We used the colors red and green for the target and distractor array items. These colors were rendered isoluminant and presented on a uniform gray background. Monkeys began a trial by fixating within 1° of visual angle (dva) around a central fixation cross. The amount of time between fixation acquisition and array onset was randomized and taken from a nonaging foreperiod function (Nickerson and Burnham, 1969; Naatanen, 1970, 1971), with times ranging from 750 to 1250 ms to reduce expectancy. Following the fixation foreperiod, a visual search array, consisting of six items was presented to the monkey. The size of the items in the array scaled with eccentricity at 0.3 dva per 1 dva eccentricity so that they were smaller than the estimated overall receptive field size (Freeman and Simoncelli, 2011). To determine the orientation and eccentricity of the array, each day online multiunit activity was measured during a receptive field mapping task (Cox et al., 2013, 2019; Dougherty et al., 2019; Westerberg et al., 2019), where the monkey fixated while a series of stimuli was presented across the visual field. The size of the items in the array scaled with eccentricity at 0.3 dva per 1 dva eccentricity so that they were smaller than the estimated overall receptive field size (Freeman and Simoncelli, 2011). To determine the orientation and eccentricity of the array, each day online multiunit activity was measured during a receptive field mapping task (Cox et al., 2013, 2019; Dougherty et al., 2019; Westerberg et al., 2019), where the monkey fixated while a series of stimuli was presented across the visual field. The array was then oriented so that its eccentricity coincided with the location of the receptive field (3–10 dva eccentricity), and a single array item was placed at the center of the receptive field. Each trial, one randomized item in the array was of a different color than the others. To acquire juice, the monkey was tasked to saccade to that item within 1000 ms and to maintain fixation within a 2–5 dva space around the target for at least 500 ms. Eye movements were monitored continuously at 1 kHz using an infrared corneal reflection system (SR Research). If the monkey made the saccade to a distractor instead, the monkey did not receive a juice reward on that trial and experienced a short (1–5 s) time-out. Trials were presented in blocks of 5–50 trials with the vast majority of blocks being 5–15 trials in length (median length, 12 trials). The color of the target and distractors were held constant throughout each block (Fig. 2B, top). At the end of a block, the target color and distractor color were swapped. The target item had an equal probability of being located at any one of the six item locations. This led to a portion of sequential trials (~16.67%) having the same target location. These sequences were used for investigation of inhibition of return (Fig. 2B, bottom).

Neurophysiological procedure

Neural spiking data were acquired with 24 kHz resolution (Tucker-Davis Technologies) from V4 of both monkeys (left hemisphere, monkey C; right hemisphere, monkey H) using custom-variant 32-channel linear microelectrode arrays (S-Probe, Plexon) across 38 sessions (n = 31, monkey C; n = 7, monkey H). Spiking activity was derived from the multiunit activity measured at each electrode contact with a significant visual response to the

Figure 2. Behavioral task and trial sequences. A, Order of events in a trial. Monkeys viewed a monitor where a fixation cross appeared at the center. After acquiring fixation and following a variable fixation period, a stimulus array appeared, consisting of a homogenous set of colored distractor disks and a single target of opponent color. Monkeys then were to saccade (denoted with a dashed line) to the target to receive a juice reward. Following reward, there was a short intertrial interval followed by the next trial. B, Example trial sequences used for investigation of priming. Top, Feature priming, sequences of the same target feature (e.g., color, red vs. green) where “1” indicates the first trial where the target took on that feature. Numbers >1 indicate repeated trials of the same target feature. Bottom, Location repetition sequence where 1 indicates that the target appeared in a location different from that of the previous trial. Numbers >1 indicate a repeated target location.
array presentation (for further details, see Data analysis and statistics). This method for deriving population-spiking activity has been described previously (Legatt et al., 1980) and has been demonstrated to be effective across multiple brain areas (Logothetis et al., 2001; Roelfsema et al., 2004; Xing et al., 2009; Self et al., 2013; Shapcott et al., 2016; Tovar et al., 2019). Briefly, the broadband neural signal was low-pass filtered at 3 kHz, high-pass filtered at 300 Hz, full-wave rectified, and, last, low-pass filtered at 150 Hz (Fig. 3). This signal was used for the analysis of spiking activity throughout the study as multiunit activity has been shown to reliably reflect neural population dynamics (Trautmann et al., 2019) and disambiguation of visual and movement cell populations is unnecessary in V4, as was necessary in the previous study in FEF (Bruce and Goldberg, 1985; Bichot and Schall, 2002; Lowe and Schall, 2018). Additionally, multiunit activity in V4 has been shown to reliably reflect attentional modulation (Mehta et al., 2000; Nandy et al., 2017).

**Data analysis and statistics**

Electrode recording sites in V4 were included for subsequent analysis if they had a significant response to the onset of the stimulus array and showed target selectivity. A significant visual response was defined as a change in the mean activity over the time between array onset and saccadic response that exceeded 2 SDs of the baseline activity (defined as the mean activity throughout 300 ms before array onset) for at least 50 ms of the array presentation epoch. Target selectivity was defined as a difference between a response to the target in the receptive field compared with the response to a distractor in the receptive field that exceeded 2 SDs between conditions for at least 25 consecutive milliseconds. A total of 469 visually responsive recording sites were identified, of which 201 (147, monkey C; 54, monkey H) showed target selectivity.

To compare the responses (e.g., response to target vs distractor, unprimed target to primed target), activity was normalized between the two compared conditions through a z-score conversion, as follows:

$$z_t = \frac{x_t - \text{mean}(x_{\text{baseline}(c1,c2)})}{s(\text{baseline}(c1,c2))}.$$  

All trials were taken for the two conditions (c1 and c2) and transformed at each time point ($t$) into z-score ($z_t$) by subtracting the mean activity during the baseline across all trials encompassed by the two conditions ($x_{\text{baseline}(c1,c2)}$) being compared from the activity at any given time point ($x_t$), then dividing by the SD of the baseline epoch ($s(\text{baseline}(c1,c2)))$. Because this normalization was condition dependent, the magnitudes of responses cannot be directly related across comparisons. For example, when...
comparing target and distractor responses for a given rep-
etition (e.g., 1, 2, 3, . . . n), only trials of that repetition posi-
tion were used in computing the normalized responses.

Target selection time was measured using previously
reported methods (Bichot and Schall, 2002; but see
Bradley et al., 1987; Britten et al., 1992; Westerberg et al.,
2019), rooted in signal detection theory (Green and
Swets, 1966; Macmillan and Creelman, 2009). We per-
formed this analysis at the population level: using 1 ms in-
crements, we compared the activity between the target
and distractor conditions at each time point from 50 ms
before array onset to 250 ms post-array onset by calculat-
ing receiver operating characteristic (ROC) curves. The
area under the curve (AUC) quantifies the degree of sepa-
ration between the conditions at each time point. An AUC
of 0.5 indicates that the conditions are indistinguishable.

As the AUC approaches 0.0 or 1.0, the conditions become
more distinguishable. Therefore, plotting AUC as a func-
tion of time allows for evaluation of the discriminability of
the target and distractor responses throughout the array
presentation epoch. Finally, to derive a point in time
where target selection can be compared across condi-
tions, a cumulative Weibull function was fit to the AUC
data across time, as follows:

\[
P = \gamma - (\gamma - \delta)e^{-\left[\frac{t}{\alpha}\right]^\beta},
\]

where \( P \) is the predicted AUC at time point \( t \), \( \gamma \), and \( \delta \)
are the maximum and minimum asymptotic thresholds,
respectively, \( \alpha \) is the time at which the distribution
reaches 64% the maximum asymptotic threshold, and \( \beta \)
is the slope. All parameters were free in the fits of all con-
ditions. The exact time of target selection was taken as
the point when the cumulative Weibull fit surpassed the
midpoint between the maximum of the minimum asympto-
tic thresholds for the conditions being compared and
the minimum of the maximum asymptotic thresholds. For
example, in the comparison of target selection times for
position in the feature priming sequence we found the
minimum and maximum asymptotic values for all posi-
tions being compared. We then used the minimum of the
maxima and maximum of the minima for those positions
to determine the midpoint for computation of target selec-
tion time.

Results

Behavioral priming in pop-out visual search

Two monkeys were trained to perform a six-item pop-
out visual search task. Sequences of trials in this task
were organized in such a way that the target pop-out fea-
ture remained constant across blocks of trials. These
blocks served to elicit feature-based priming of pop-out,
as reported in other studies (Maljkovic and Nakayama,
1994; Bichot and Schall, 1999, 2002; Purcell et al., 2012;
Westerberg et al., 2020). Briefly, monkeys fixated a cen-
tral fixation cross for a variable amount of time before the
appearance of the search array. The monkey then identi-
fied the color pop-out target (either a red target among
green distractors or vice versa) by making a saccade to
that item as quickly as possible. Successful responses

were rewarded following a 500 ms target fixation period.
Incorrect responses led to a short time-out and no
reward.

Both monkeys showed priming. Response times, meas-
ured as saccade latency, were consistently reduced through
the priming sequences (Fig. 4A), reported here as trial from
feature change (e.g., target changing from red to green or
vice versa). Following a change in the target feature, per-
formance degraded before improving again. Behavioral im-
provement seemed to show an asymptote around five trials
following the feature change. For this reason, trials beyond
this point were combined. The apparent change in response
times was determined to be a statistically significant trend
measured through Page’s L test (Page, 1963) on a session-
by-session basis (L38.5 = 2033, \( p << 0.001 \)). This was also
determined to be statistically significant in each monkey in-
dividually (monkey C: \( L_{31.5} = 1656, p << 0.001 \); monkey H:
\( L_{7.5} = 377, p << 0.001 \)). Accuracy improved through the pri-
m ing sequences as well (Fig. 4B), as determined through
Page’s L test on a session-by-session basis (L38.5 = 2006,
\( p << 0.001 \)). This, too, was determined to be statistically sig-
nificant in each monkey individually (monkey C: \( L_{31.5} = 1639,
 p << 0.001 \); monkey H: \( L_{7.5} = 367, p << 0.001 \)). These results
are consistent with previous reports of feature-based pri-
m ing in humans (Maljkovic and Nakayama, 1994) and mac-
aques (Bichot and Schall, 1999, 2002; Purcell et al., 2012;
Westerberg et al., 2020).
To determine whether V4 shows neural correlates of priming of pop-out, we measured target selection time across our neural population. Previous reports suggest speeded attentional selection may mediate behavioral priming in this task (Bichot and Schall, 2002; Eimer et al., 2010; Kristjánsson and Campana, 2010; Kristjánsson and Ásgeirsson, 2018) and V4 has been shown to be essential for attentional processing (for review, see Reynolds and Heeger, 2009). The neural measure of target selection time was adapted from methods of signal detection theory (Green and Swets, 1966). The details of how target selection times were computed can be found in the Materials and Methods. The threshold for target selection in ROC analysis was 0.5815. Spiking activity averaged across the population of target-selective V4 recording sites was divided up by (1) trial-since-feature-change and (2) whether the target or a distractor was present in the receptive field (Fig. 5A). We then computed the target selection time for the first five trials since feature change (Fig. 5B). We found that feature repetition decreased target selection time. Immediately following the switch of target feature, V4 neurons identified the target at 190 ms. This time decreased to 163 ms in the following trial before reaching the fastest selection time at 126 ms in the fifth trial.

**Figure 5.** V4 target selection during priming of pop-out. 
**A.** Spiking activity aligned on array presentation, averaged across all recording sites (n = 201) when the target was in the receptive field (black) versus when a distractor was in the receptive field (gray). Clouds represent 95% confidence intervals (CIs). Data are broken out by trial since feature change (right) from top to bottom. **B.** AUC as a function of time from array onset where gray dots are the population average experimental data for each time point, and black curves are the cumulative Weibull fits. Computed target selection times (for details, see Materials and Methods) are indicated with an arrow pointing at the abscissa for each plot.

**Speeded attentional selection in V4 with feature-based priming**

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trial following the feature change. To confirm that the change in target selection time was statistically significant, we performed a Page’s L test. To do so we subsampled 80% of the recorded sites and computed the target selection time, via the same ROC methods, 25 times. This method allows for the evaluation of the variability in target selection time across the population by repeatedly subsampling the data (Sato and Schall, 2003). We found a significant decrease in the target selection time of the whole population through these methods ($L_{25.5} = 1363, p < 0.001$) as well as for the individual monkeys (monkey C: $L_{25.5} = 1360, p < 0.001$; monkey H: $L_{25.5} = 1172, p < 0.05$). These results suggest that attentional selection of the target stimulus in V4 hastens with feature-based priming.

**Relationship between feature-based priming of behavior and selection in V4**

We next sought to determine whether there was any correlational relationship between the changes observed in the behavior of the monkeys and the changes in target selection in V4, as previous work suggests that causal inactivation of V4 through lesions degrades the potency of priming effects (Walsh et al., 2000). We first plotted the relationship between changes in response time (averaged across sessions) against the population selection times (Fig. 6A). We performed a least-squares linear regression to obtain a fit and slope. We found a significant relationship, determined through a comparison against a constant model, between the behavioral and neural measures ($F_{1,4} = 84.9, p = 0.0029$). The slope of the fit, 0.59, was significantly different from 0 ($t_3 = 9.22, p = 0.0027$). The slope measures the relationship between neural selection time and response time. If they changed perfectly proportionally (e.g., a 10 ms decrease in neural response time corresponded with a 10 ms decrease in response time), the slope would be 1. Slopes >1, as plotted, reveal a larger change in response time relative to the change in neural selection time (e.g., a 10 ms decrease in neural selection time corresponding to a 15 ms decrease in response time would yield a slope of 1.5). Slopes <1, but >0 reveal a smaller change in response time relative to the change in neural selection time (e.g., 10 ms decrease in neural selection corresponding to a 5 ms decrease in response time would yield a slope of 0.5). The slope measured here indicates that the variation in target selection time in V4 exceeds the concomitant change in response time.

While there appears to be a systematic relationship between the neural target selection time in V4 and behavioral response times, we did not find a relationship between the probability of a correct response and the magnitude of discriminability between target and distractor in V4 (Fig. 6B). The probability of correct responses was taken as the mean across sessions, and the magnitude of distinguishability was derived from the maximum AUC across time using the Weibull fit of the experimental data. This value was adjusted for the number of items in the array as ROC analysis is usually performed for two-alternative forced choice tasks while our arrays featured six possible choices.

The conversion was done through a conversion table provided by Hacker and Ratcliff (1979; see also Smith, 1982; Macmillan and Creelman, 2009). The least-squares fit for the behavioral and neural measures of accuracy was not significant ($F_{1,4} = 0.0097, p = 0.875$), nor was the slope ($t_3 = -0.17, p = 0.874$). In contrast to the change in performance as a function of position in priming sequence (Fig. 4), the discrimination of target and distractors in V4 was invariant across position in priming sequence. Hence, as observed in Figure 6B, the unprimed trial associated with the worst performance did not have the lowest adjusted area under the ROC.

We then sought to determine whether these relationships between neural and behavioral measures could be found at the individual recording site level. To do so, we performed the Weibull fitting procedure (for details, see Materials and Methods) for each recording site as a function of trial since target position change. We then performed two linear regressions. One regression was performed between the measured target selection time of the individual recording site and the behavioral response time for the session (Fig. 7A,B). Another regression was performed on the measured adjusted maximum area under the ROC for the individual recording site and the behavioral accuracy for the session (Fig. 7C,D). Both regressions were performed as a function of the number of trials since the target feature change. Only units showing a significant difference in target selection time in the initial Weibull fit as a function of feature priming condition are shown in Figure 7. By bootstrapping the Weibull fits 25 times and performing an ANOVA ($p < 0.05$) on the results for the target selection times as a function of trial since target feature change, of 201 units, 103 satisfied this criterion.

**Figure 6.** Effect of feature priming on the relationship between behavior and neural processing in V4. **A**, Response time as a function of target selection time (see Materials and Methods). Dashed vertical and horizontal lines around each data point represent the 95% confidence intervals. The five points plot data from each of the five trials since feature change. Goodness-of-fit through adjusted $R^2$ for the least-squares regression denoted in the bottom right. **B**, Probability of a correct response as a function of adjusted maximum area under the ROC. Dashed vertical and horizontal lines around each data point represent the 95% confidence interval for probability of correct response and adjusted maximum area under the ROC, respectively. Goodness-of-fit through adjusted $R^2$ for the least-squares regression is denoted in the bottom right.
To quantify the relationship between the neural and behavioral measures, we measured the slopes for each of the regressions (Fig. 7). The median slope of the relationship between neural target selection time and response time was 0.47, which was significantly different from 0 (Wilcoxon signed-rank test: mean = 0.47; \( Z = 8.81 \); \( p = 1.8 \times 10^{-18} \)). This number is but not, like the slope measured at the population level, indicating more variation over feature priming in target selection time than in response time. The median slope of the relationship between adjusted maximum area under the ROC and the behavioral accuracy was 0.29, which was not significantly different from 0.

Figure 7. Relationship between neural and behavioral measures for individual units. A, Linear regressions of neural target selection time and behavioral response time for each unit with a measurable change in target selection as a function of feature priming (\( n = 103 \)). B, Distribution of the slopes plotted in A. The median slope of 0.47, which was significantly different from 0 (black dashed line, ***), is indicated by the red dashed line. C, Linear regressions of adjusted maximum area under the ROC and behavioral accuracy for each of the units (\( n = 103 \)). D, Distribution of the slopes in C. The median slope was 0.29, which was not significantly different from 0.

Target enhancement and distractor suppression during feature-based priming

Changes in attentional selection times could come about in a variety of different ways, such as enhancement of the target stimulus, suppression of distractors, or a combination of both. Previous work has shown that in FEF, both target enhancement and distractor suppression contribute to the effect (Bichot and Schall, 2002). We determined whether V4 shows target enhancement and distractor suppression with priming by comparing the mean spiking activity in the 50 ms preceding the average response time for early trials in the priming sequence to those later in the sequence (i.e., trials 1 and 2 since feature change and trial 3 or more, respectively). Results on the population level are shown in Figure 8A.

We found a significant difference between the mean spiking activity in the unprimed target (mean = 5.91; SD = 5.58) and primed target (mean = 6.45; SD = 6.14) conditions (\( t_{200} = -9.68, p = 2e^{-10} \)), which is in line with target enhancement during feature priming. Additionally, there was a significant difference between the mean activity in the unprimed distractor (mean = 5.09; SD = 4.70) and primed distractor (mean = 4.73; SD = 4.73) conditions (\( t_{200} = 7.15, p = 2e^{-11} \)), indicative of distractor suppression during feature priming. Both results were robust in each monkey (monkey C: target enhancement, \( t_{146} = -10.85, p = 2e^{-20} \); distractor suppression, \( t_{146} = 7.23, p = 3e^{-11} \); monkey H:...
target enhancement, $t_{53} = -2.77, p = 0.0077$; distractor suppression, $t_{53} = 2.26, p = 0.028$). Furthermore, we found no significant difference between the magnitudes of target enhancement (mean = 0.70; SD = 7.96) and distractor suppression (mean = 0.57; SD = 7.06) across the population ($t_{200} = 1.41, p = 0.16$).

We also performed a linear regression on the difference between target and distractor responses during the 50 ms preceding the average response time between the feature-primed and feature-unprimed trials (Fig. 8B). The regression significantly exceeds the unity line. This suggests a greater target–distractor response difference in the feature-primed trials. This is confirmed by the significant shift from zero of the intercept of the regression (intercept $= 0.20, p = 0.033$) and the slope being $> 1$ (slope $= 1.18, p = 2e^{-91}$). These results parallel the findings in FEF and suggest that the changes in target selection times are mediated by a combination of changes in V4 processing of targets and distractors, rather than just one or the other.

Baseline activity remains stable with feature-based priming

Spontaneous firing rates in area V4 have been shown previously to modulate with allocation of attention when locations are predictable (Luck et al., 1997). This sort of attentional modulation could account for the hastened target selection time observed in V4 with feature priming. By enhancing the spontaneous firing rate in V4 following a change in target feature, it is feasible that the selectivity for the feature is enhanced. While the monkeys are not cued to the location of the target stimulus before array onset in this task used in this study, we investigated whether baseline activity varied systematically following a change in target feature in the feature-based priming sequences. To do so, we measured the average neural activity in the 300 ms before presentation of the search array. Values were compared across the initial five trials in the priming sequence (Fig. 9). A one-way ANOVA revealed no significant difference ($F_{(4,1000)} = 1.45, p = 0.21$). Thus, the spontaneous firing rate in V4 during the baseline period does not change as a function of feature priming.

Inhibition of return with target position repetition

The second form of repetition-induced behavioral modification that we were interested in is the changes associated with repetition of the target position in the search array. Previous work has shown two different behavioral outcomes under these circumstances. One is priming, where response times hasten when the target appears in the same location (Maljkovic and Nakayama, 1996). The other finding is inhibition of return, where repetition of target location leads to slowed behavior (Bichot and Schall, 2002; for review, see Klein, 2000). To determine the behavioral consequence of target position repetition in our task, we identified sequences where the target position changed and compared these with instances where target position was constant across subsequent trials. In our paradigm, the target had equal probability of appearing at any one of the six item locations that lead to repetition.
approximately one-sixth of the time, with sequences of three repetitions approximately one-thirty-sixth of the time.

Figure 10 summarizes the behavioral results of target position repetition. Qualitatively, it appears that response times increase with position repetition while accuracy remains largely unchanged. These results would replicate previous findings in monkeys performing this task (Bichot and Schall, 2002). Quantitative analysis tells a different story. Response time does significantly increase, as indicated by a Pages’ L test ($L_{38.3} = 506, p < 0.001$), and this finding is robust for both monkeys individually (monkey C, $L_{31.3} = 413, p < 0.001$; monkey H, $L_{7.3} = 93, p = 0.01$). However, while the increase in accuracy is small (0.43% from 1 to 2 and 1.53% from 2 to 3), it was a significant trend across both monkeys ($L_{38.3} = 484, p = 0.001$). However, when breaking out the data by monkey, it was only monkey C that showed this effect ($L_{31.3} = 399, p < 0.001$), while monkey H did not ($L_{7.3} = 85, p > 0.05$).

**Slowed attentional selection in V4 with target position repetition**

We next sought to determine whether neural correlates of inhibition of return were present in V4 as FEF has previously been shown to exhibit neural correlates of this behavioral phenomenon (Bichot and Schall, 2002). Evaluation of the neural correlates of inhibition of return in V4 was performed in a manner similar to that for feature-based priming. First, we plotted the spiking activity for each of the conditions comparing when the target was in the receptive field to when a distractor was present instead (Fig. 11A). We then evaluated target selection time as a function of trials since target position change using the same ROC methods. The threshold for target selection in the ROC analysis was 0.5703 for the analysis of target position repetition. We compared the trials where the previous trial had a different target location than the current trial (trial 1) to trials where the target had been in the same location for one to two previous trials (trials 2–3; Fig. 11B). We found that target selection times slowed by an average of 11 ms following target position repetition. This difference was found to be significant through a paired t test on the bootstrapped data where we subsampled 80% of the recording sites 25 times and computed the statistic ($t_{200} = -11.72, p = 2e^{-11}$). This finding was confirmed at the individual monkey level (monkey C, $t_{146} = -16.83, p = 8e^{-10}$; monkey H, $t_{53} = -6.29, p = 2e^{-5}$).

**Relationship between location-based inhibition of return and selection in V4**

As we found a correlation between feature-based priming enhancements of behavior and target selection times in V4, we wanted to investigate whether a similar phenomenon could be observed with target position repetition. To do so, we plotted the mean response time as a function of target selection time in V4 (Fig. 12). As we had only two data points, we did not perform a least-squares fit. However, we nonetheless estimated the slope of the relationship by measuring the change in response times between the two points. We measured the slope of the line to determine the relationship between the change in selection time and response time. Again, a slope of 1 (a one-to-one relationship) would suggest that response time and selection time change at the same rate. Where the feature-based priming had a slope <1 between response time and target selection time, the inhibition of return condition did in fact have a slope near to 1. This
Target suppression with target position repetition

A slowing of attentional selection time can happen in several ways, such as suppression of the target, enhancement of distractors, or a combination of both. Hence, we determined whether V4 shows target suppression and distractor enhancement concurrently with inhibition of return. This was accomplished by comparing the mean spiking activity in the 50 ms preceding the average response time for trials in which the target position did repeat and when it did not repeat (i.e., trial 1 since feature position change and trial 2 or more, respectively). Results on the population level are shown in Figure 13A. Mean spiking activity was significantly different between the target position change (mean = 9.08; SD = 6.90) and the target position repeat (mean = 8.72; SD = 6.74) conditions for the target response ($t_{200} = 4.60, p = 7e^{-5}$), consistent with target suppression contributing to inhibition of return. Mean spiking activity was not significantly different between the mean activity in the target position change (mean = 5.30; SD = 3.81) and target position repeat (mean = 5.41; SD = 3.76) conditions for the distractor response ($t_{200} = -1.17, p = 0.24$), indicating no contribution of distractor enhancement to inhibition of return. The target suppression result was robust in each monkey (monkey C, $t_{146} = 5.04, p = 1e^{-6}$; monkey H, $t_{53} = 2.15, p = 0.033$). Neither monkey showed significant distractor enhancement (monkey C, $t_{146} = -0.63, p = 0.53$; monkey H, $t_{53} = -1.40, p = 0.16$).

We also performed a linear regression on the difference between target and distractor responses during the 50 ms preceding the average response time between the target position change and target position repeat trials (Fig. 13B). Here we find that the regression line lies below the unity line. This suggests a greater target–distractor response difference in the target position change trials. This is confirmed by the significant shift from zero of the intercept of the regression ($y = -0.31, p = 0.037$) and the slope measured to be below 1 ($slope = 0.96, p = 6e^{-90}$). These results suggest that the change in response time is mediated by a suppression of targets in the receptive field alone, rather than a combination of changes in target and distractor responses like that in the feature priming case.

Baseline activity remains stable with target position repetition

While we did not observe a change in the spontaneous neural activity with feature-based priming, we sought to determine whether there was a change associated with target position repetition. To do so, we compared the baseline activity during the 300 ms leading up to array
onset (while the monkey was fixating) of trials where the prior target position was outside the receptive field to trials where the target position repeated in the receptive field. Through a paired $t$ test, we found no difference in the average activity between the target position change (mean = 0.0021; SD = 0.54) and target position repeat (mean = $-0.058$; SD = 0.62) conditions ($t_{200} = 1.29, p = 0.20$). This suggests that like feature-based facilitation, inhibition of return does not change the spontaneous firing rate in V4.

**Discussion**

Behavioral priming can be elicited through feature-based facilitation of return in visual search under pop-out conditions. Previous work suggests that attentional priming mediates these changes (for review, see Kristjánsson and Campana, 2010; Kristjánsson and Ásgeirsson, 2018) as well as that frontal area FEF demonstrates neural correlates of these behavioral changes (Bichot and Schall, 2002). Noninvasive electrophysiological measures demonstrate altered activity in an area over posterior regions of the brain (Eimer et al., 2010), and a lesion study implicates visual cortex in priming (Walsh et al., 2000). To investigate whether visual cortex exhibits neural correlates of the behavioral changes observed in priming, we recorded population-spiking responses in area V4 of two macaque monkeys. We primarily sought to determine whether attentional selection in area V4 modulates with priming in feature-based priming of pop-out. We also wanted to determine whether neural correlates of location-based inhibition of return could be observed in V4 neurons. We found that attentional selection, measured as target selection time, modulated with both feature-based priming and location-based inhibition of return. These results suggest an explicit role for visual cortex in priming in pop-out and inhibition of return.

**Evidence for priming of pop-out as attentional priming**

The mechanism underlying the behavioral improvements associated with priming of pop-out in visual search has been a matter of some debate in psychology (Kristjánsson and Campana, 2010; Kristjánsson and Ásgeirsson, 2018). While there is psychophysical evidence suggesting that priming of pop-out is mediated by memory mechanisms (Hillstrom, 2000; Huang et al., 2004; Huang and Pashler, 2005), electrophysiological evidence largely supports the idea that it is mediated through attentional priming (Bichot and Schall, 2002; Eimer et al., 2010). More recent psychophysical evidence also supports the attentional priming hypothesis (Brascamp et al., 2011). The findings of this study further support the idea that priming of pop-out is mediated through attentional priming mechanisms, as we found that the speed of attentional selection was speeded in V4 in this task. While it is possible that other cognitive phenomena, such as performance monitoring (Westerberg et al., 2020), may modulate as a function of priming in this task, they are likely secondary to the changes in attentional selection (Kristjánsson and Ásgeirsson, 2018).

**Putative circuitry for attentional priming**

Few publications describe neural mechanisms of priming in pop-out visual search. FEF, a frontal area with known functional relevance to attention, eye movements, and vision shows speeded target selection during visual search (Bichot and Schall, 2002). SEF shows no changes during the period when attentional selection would occur (Purcell et al., 2012) and is therefore unlikely to contribute. Lesions of V4 and TEO suggest that those areas are necessary for priming effects (Walsh et al., 2000), and our results suggest that the speed of attentional selection is increased in V4 under these priming conditions. Together, these results suggest two plausible hypotheses regarding how attentional priming comes about in this task: (1) attentional selection is accomplished in visual cortex in pop-out visual search and repetition of target feature primes that selection, which is then inherited by FEF leading to shorter response times; and (2) attentional selection of the behaviorally relevant stimulus occurs in FEF, and the target selection signal is then fed back to visual cortex, leading to the observations in this study. To distinguish between these hypotheses, one could determine whether priming of pop-out and inhibition of return are evident in the feedforward input layers (granular layer 4) of V4 or only in the extragranular layers (layers 2/3 or 5/6). The vast majority of feedback to V4 from FEF occurs outside of the granular layers. Therefore, if the priming effects are first observed in the granular layers, it would suggest that these effects are generated in visual cortex. If the signals are exclusively observed in the extragranular layers, the history effects would either be from feedback to V4 or through processes occurring locally within V4. More definitively, one must record simultaneously from FEF and V4 and perform trial-by-trial analyses of target selection to determine the origins of the priming effect.

However, one hypothesis seems more likely given the quantitative results of these studies. In the study of FEF, there was no relationship between the target selection time in that area and the response time across priming conditions. This indicates that as selection time decreased in FEF, the behavioral response time decreased equally as much. In V4, we observe a less than one-to-one relationship where target selection time decreased more strongly than response time. This relationship perhaps suggests that FEF receives crucial visual input from V4, but in resolving the behaviorally relevant stimulus, it does not perfectly integrate that information. This would mean that although V4 can more rapidly identify the target during priming, FEF acts as a bottleneck, in some manner. This combination leads to speeded responses, but not as speeded as would be predicted by the V4 activity. This hypothesis suggests that FEF target selection is necessary for generating the response but is not the sole origin of the behavioral effect.

The reason FEF does not speed target selection as rapidly as V4 may be because FEF receives input from many other visual cortical areas, which could add noise to the process (Schall et al., 1995; Stanton et al., 1995). The V4 lesion study discussed above perhaps provides some evidence for this as well (Walsh et al., 2000). Lesioning either
V4 or TEO (both providing input to FEF) reduces priming while Lesioning either V4 or TEO (both providing input to FEF) reduces priming. Additionally, FEF shows a relationship between neural measures of discriminability and response accuracy, while V4 does not. This may suggest that V4 feeds FEF relatively invariant selection signals, whereas behavior more closely follows FEF neural measures of discriminability, which may be noisy, as previously suggested. It is important to remember that this notion is speculative, but it would suggest that attentional priming originates in sensory cortex before being inherited by the frontal area that then plans and initiates the response.

Further evidence for this hypothesis is available in comparing the measured target selection times in FEF previously and V4 now. Unfortunately, the FEF data are no longer readable, so this comparison is limited to published values. On the first trial in feature priming sequence, target selection time was earlier in V4 (190 ms; Fig. 5b) than in FEF (217 ms; Bichot and Schall, 2002). In subsequent trials, target selection times in V4 (trials 2–5: 163, 146, 147, and 126 ms; Fig. 4b) were somewhat later than those in FEF (trials 2–5: 131, 125, 117, and 120 ms; Bichot and Schall, 2002). These results seem reasonable in the context of the visual input filtering hypothesis for FEF target identification. We note that directly comparing the results of FEF to those of V4 may be unfair given that the data were obtained from different monkeys, which contributes unknown, subtle differences in behavior. Also, the summary times from Bichot and Schall (2002) were taken from a single representative unit rather than from the population. To draw more confident conclusions regarding how attentional priming comes about across cortical areas, simultaneous recordings or inactivation studies must occur.

The rationale behind this hypothesis lends itself to an interpretation of the inhibition of return result as well. While we found a less than one-to-one relationship with feature-based priming, we did find a one-to-one relationship between behavioral and neural processing for location-based inhibition of return. Whereas the priming of attentional selection may originate in visual cortex, this observation would suggest that V4 may inherit inhibition of return. Because of the task design (e.g., no predefined blocks of target position repetition, unlike the feature priming blocks), we are unable to determine whether this is a purely linear relationship as it may change further into the target position repetition sequence. Again, it is important to remember that these are hypotheses generated by the results of this study that must be tested more directly before a complete model of priming of attentional selection can be drawn.

Comparative relevance

The finding of this study that priming of attentional selection is observed in V4, taken in consideration with the same observation in FEF (Bichot and Schall, 2002), may provide some insight into the nature of attentional selection signals observed through noninvasive measures in humans. More specifically, these findings may provide insight into the N2pc event-related potential (ERP) component, a measure of covert spatial attentional selection (Luck and Hillyard, 1994; Eimer, 1996; Woodman and Luck, 1999). The magnitude of the N2pc has been shown to be largest over occipital and parietal cortices (Luck and Hillyard, 1994), with evidence from a magnetoencephalography study showing potential sources of the N2pc in parietal and occipital cortex in humans (Hopf et al., 2000).

Monkeys performing the same task show the same attentional selection signals in the ERP (Woodman et al., 2007) as well as similar, but earlier, concurrently recorded attentional selection signals in the spiking activity in FEF (Cohen et al., 2009; Heitz et al., 2010; Purcell et al., 2013). Together with the fact that FEF is extensively connected to V4 (Schall et al., 1995; Stanton et al., 1995), it is conceivable that FEF sends attentional selection signals to V4 where the N2pc is then generated. The priming results seem to corroborate this hypothesis. The N2pc in humans is speeded with feature-based priming in pop-out visual search (Eimer et al., 2010), as is attentional selection in FEF (Bichot and Schall, 2002). With the finding that FEF and V4 show speeded attentional selection, it is conceivable that V4 might be the generator of the N2pc that also shows speeded selection. Further work is needed to determine whether the biophysical properties of V4 could support the generation of the N2pc, however these functional correlates support the hypothesis. Further work is also necessary to determine whether FEF is necessary for the N2pc and whether it is necessary for speeding through priming or whether V4 alone is sufficient for this to occur.

References

Armstrong KM, Moore T (2007) Rapid enhancement of visual cortical response discriminability by microstimulation of the frontal eye field. Proc Natl Acad Sci U S A 104:9499–9504.

Armstrong KM, Fitzgerald JK, Moore T (2006) Changes in visual receptive fields with stimulation of frontal cortex. Neuron 50:791–798.

Bichot NP, Schall JD (1999) Saccade target selection in macaque during feature and conjunction visual search. Vis Neurosci 16:81–89.

Bichot NP, Schall JD (2002) Priming in macaque frontal cortex during popout visual search: feature-based facilitation and location-based inhibition of return. J Neurosci 22:4675–4685.

Bradley A, Skottun BC, Ohzawa I, Sclar G, Freeman RD (1987) Visual orientation and spatial frequency discrimination: a comparison of single neurons and behavior. J Neurophysiol 57:755–772.

Brascamp JW, Blake R, Kristjansson Á (2011) Deciding where to attend: priming of pop-out drives target selection. J Exp Psychol Hum Percept Perform 37:1700–1707.

Britten KH, Shadlen MN, Newsome WT, Movshon JA (1992) The analysis of visual motion: a comparison of neuronal and psycho-physical performance. J Neurosci 12:4745–4765.

Bruce CJ, Goldberg ME (1988) Primate frontal eye fields. I. Single neurons discharging before saccades. J Neurophysiol 53:603–635.

Cohen JY, Heitz RP, Schall JD, Woodman GF (2009) On the origin of event-related potentials indexing covert attentional selection during visual search. J Neurophysiol 102:2375–2386.

Cox MA, Schmid MC, Peters AJ, Saunders RC, Leopold DA, Maier A (2013) Receptive filed focus of visual area V4 neurons determines
responses to illusory surfaces. Proc Natl Acad Sci U S A 110:17095–17100.

Cox MA, Dougherty K, Westerberg JA, Schall MS, Maier A (2019) Temporal dynamics of binocular integration in primary visual cortex. J Vis 19:1–21.

De Weerd P, Peralta IM, Desimone R, Ungerleider L (1999) Loss of attentional stimulus selection after extrastriate cortical lesions in macaques. Nat Neurosci 2:753–758.

De Weerd P, Desimone R, Ungerleider L (2003) Generalized deficits in visual selective attention after V4 and TEO lesions in macaques. Eur J Neurosci 18:1671–1691.

Dougherty K, Cox MA, Westerberg JA, Maier A (2019) Binocular modulation of monocular V1 neurons. Curr Biol 29:381–391.

Eimer M (1996) The N2pc component as an indicator of attentional selectivity. Electroencephalogr Clin Neurophysiol 99:225–234.

Eimer M, Kiss M, Cheung T (2010) Priming of pop-out modulates attentional target selection in visual search: behavioural and electrophysiological evidence. Vision Res 50:1353–1361.

Fecteu JH, Munoz DP (2003) Exploring the consequences of the previous trial. Nat Rev Neurosci 4:435–439.

Fecteu JH, Munoz DP (2005) Correlates of capture of attention and inhibition of return across stages of visual processing. J Cogn Neurosci 17:1714–1727.

Freeman J, Simoncelli EP (2011) Metamers of the ventral stream. Nat Neurosci 14:1195–1201.

Green DM, Swets JA (1966) Signal detection theory and psychophysics. New York: Wiley.

Gregoriou GG, Rossi AF, Ungerleider LG, Desimone R (2014) Lesions of prefrontal cortex reduce attentional modulation of neuronal responses and synchrony in V4. Nat Neurosci 17:1003–1011.

Hacker MJ, Ratcliff R (1979) A revised table d’ for M-alternative forced choice. Percept Psychophys 26:168–170.

Heitz RP, Cohen JY, Woodman GF, Schall JD (2010) Neural correlates of correct and errant attentional selection revealed through N2pc and frontal eye field activity. J Neurophysiol 104:2433–2441.

Hillstrom AP (2000) Repetition effects in visual search. Percept Psychophys 62:800–817.

Hopf J-M, Luck SJ, Girelli M, Hagner T, Mangun GR, Scheich H, Heinze H-J (2000) Neural sources of focused attention in visual search. Cereb Cortex 10:1233–1241.

Huang L, Pashler H (2005) Expectation and repetition effects in search for featural singletons in very brief displays. Percept Psychophys 67:150–157.

Huang L, Holcombe AO, Pashler H (2004) Repetition priming in visual search: episodic retrieval, not feature priming. Mem Cogn 32:12–20.

Klein RM (2000) Inhibition of return. Trends Cogn Sci 4:138–147.

Kristjánsson Á, Campana G (2010) Where perception meets memory: a review of repetition priming in visual search tasks. Atten Percept Psychophys 72:5–18.

Kristjánsson Á, Åsgeirsson ÁG (2018) Attentional priming: recent insights and current controversies. Curr Opin Psychol 29:71–75.

Legatt AD, Arezzo J, Vaughan HG Jr (1980) Averaged multiple unit activity as an estimate of phasic changes in local neuronal activity: effects of volume-conducted potentials. J Neurosci Methods 2:203–217.

Logothetis NK, Pauls J, Augath M, Trinath T, Oeltermann A (2001) Neurophysiological investigation of the basis of the fMRI signal. Nature 412:150–157.

Low KA, Schall JD (2018) Functional categories of visuomotor neurons in macaque frontal eye field. eNeuro 5:ENEURO.0131-18.2018.

Luck SJ, Hilliard SA (1994) Electrophysiological correlates of feature analysis during visual search. Psychophysiology 31:291–308.

Luck SJ, Chelazzi L, Hilliard SA, Desimone R (1997) Neural mechanisms of spatial selective attention in areas V1, V2, and V4 of macaque visual cortex. J Neurophysiol 77:24–42.

Macmillan NA, Creelman CD (2009) Detection theory: a user’s guide. New York: Psychology Press.
for adaptation in response to visual repetition. J Neurophysiol 121:1938–1952.

Westerberg JA, Maier A, Woodman GF, Schall JD (2020) Performance monitoring during visual priming. J Cogn Neurosci 32:515–526.

Woodman GF, Luck SJ (1999) Electrophysiological measurement of rapid shifts of attention during visual search. Nature 400:867–869.

Woodman GF, Kang MS, Rossi AF, Schall JD (2007) Nonhuman primate event-related potentials indexing covert shifts of attention. Proc Natl Acad Sci U S A 104:15111–15116.

Xing D, Yeh CI, Shapley RM (2009) Spatial spread of the local field potential and its laminar variation in visual cortex. J Neurosci 29:11540–11549.