

Electrophysiological Indices of Distractor Processing in Visual Search Are Shaped by Target Expectations

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Abstract

■ Although in many cases salient stimuli capture attention involuntarily, it has been proposed recently that under certain conditions, the bottom-up signal generated by such stimuli can be proactively suppressed. In support of this signal suppression hypothesis, ERP studies have demonstrated that salient stimuli that do not capture attention elicit a distractor positivity (P_D), a putative neural index of suppression. At the same time, it is becoming increasingly clear that regularities across preceding search episodes have a large influence on attentional selection. Yet to date, studies in support of the signal suppression hypothesis have largely ignored the role of selection history on the processing of distractors. The current study addressed this issue by examining how electrophysiological markers of attentional selection (N2pc) and suppression (P_D) elicited by targets and distractors, respectively, were modulated when the search target randomly varied instead of being fixed across trials. Results showed that although target selection was unaffected by this manipulation, both in terms of manual response times, as well as in terms of the N2pc component, the P_D component was reliably attenuated when the target features varied randomly across trials. This result demonstrates that the distractor P_D , which is typically considered the marker of selective distractor processing, cannot unequivocally be attributed to suppression only, as it also, at least in part, reflects the upweighting of target features.

INTRODUCTION

The understanding that attentional selection is not only determined by the interaction between top-down and bottom-up processes, but is also strongly influenced by previous selection episodes has revived the attentional capture debate (Luck, Gaspelin, Folk, Remington, & Theeuwes, 2021; van Moorselaar & Slagter, 2020). This debate is centered around the perplexing puzzle that on the one hand salient, yet irrelevant, stimuli often appear to involuntarily capture attention, while at the same time such involuntary capture would make simple everyday tasks nearly impossible. Recent research demonstrating that learning about regularities in the environment does not only prioritize target properties, such as the location and features of the target (Chun & Jiang, 1998), but can also result in the suppression of task irrelevant information (van Moorselaar & Theeuwes, 2022; Sauter, Liesefeld, & Müller, 2019; Ferrante et al., 2018; Wang & Theeuwes, 2018; Leber, Gwinn, Hong, & O'Toole, 2016), may provide a missing piece to this puzzle.

As an initial resolution, the signal suppression hypothesis (Gaspelin & Luck, 2018b; Sawaki & Luck, 2010) was put forward, which elegantly incorporates the seemingly conflicting sides of the attentional capture debate. Specifically, the signal suppression hypothesis posits that while salient

stimuli all generate a bottom-up "attend-to-me" signal and hence compete for attention, this signal can be proactively suppressed if the attentional system is appropriately configured (Luck et al., 2021). This idea is supported by empirical findings that salient stimuli automatically capture attention when the target is defined as a unique element in the display and therefore there is no clear search goal (e.g., the target is defined as the unique shape in the search display; singleton detection mode). By contrast, when the target is a specific shape embedded in a display of heterogeneous shapes and observers hence engage in what is called feature search mode (Bacon & Egeth, 1994), allowing them to impose top-down control, which not only eliminates attentional capture (Folk, Remington, & Johnston, 1992), but under certain conditions can even result in below baseline suppression (Gaspelin & Luck, 2019). Although support for this hybrid model of attentional capture has accumulated through converging evidence of psychophysics (e.g., Adam, Patel, Rangan, & Serences, 2021; Gaspelin, Leonard, & Luck, 2015) and eye movement studies (e.g., Gaspelin, Gaspar, & Luck, 2019; Gaspelin, Leonard, & Luck, 2017), some influential studies relied on electrophysiological indices related to attentional selection (the N2pc) and suppression (the P_D; Eimer, 1996; Luck & Hillyard, 1994a, 1994b) and suppression (Hickey, Di Lollo, & McDonald, 2009).

Although the N2pc, which is a negative-going deflection occurring around 200–300 msec after stimulus presentation that is larger over the hemisphere contralateral to the attended location, is unequivocally considered as an

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index of covert attention (Luck, 2012; Woodman & Luck, 2003), the P_D is less well characterized. Nevertheless, the P_D , which is much like the inverse of the N2pc, as it presents itself as a larger contralateral positive deflection to a to-be-ignored rather than a task-relevant stimulus, is commonly presented as a putative neural index of suppression (Gaspelin & Luck, 2018b; Hickey et al., 2009) and has become a tool to study whether salient distractors can be suppressed proactively (Stilwell, Egeth, & Gaspelin, 2022; Drisdelle & Eimer, 2021; van Moorselaar, Daneshtalab, & Slagter, 2021; van Moorselaar, Lampers, Cordesius, & Slagter, 2020; van Moorselaar & Slagter, 2019; Wang, van Driel, Ort, & Theeuwes, 2019). Indeed, various strands of evidence link the P_D to a suppressive mechanism; the P_D often appears exclusively in response to distractors in the absence of a N2pc (e.g., Gaspar & McDonald, 2014; Jannati, Gaspar, & McDonald, 2013; Sawaki & Luck, 2010), it has a larger amplitude on a subset of trials with the fastest responses (McDonald, Green, Jannati, & Di Lollo, 2013), and it is no longer found when observers fail to avoid an eye movement toward the distractor (Weaver, van Zoest, & Hickey, 2017). However, arguably the most convincing evidence thus far linking the P_D to suppression comes from a study by Gaspelin and Luck (2018a) that observed a correlation between the magnitude of below baseline behavioral suppression and the magnitude of the P_D component (see also Feldmann-Wüstefeld, Brandhofer, & Schubö, 2016; Gaspar & McDonald, 2014).

There is thus good reason to believe that in many situations, the P_D can be linked to a suppressive mechanism (van Moorselaar & Slagter, 2020). At the same time, it should be noted that studies examining below baseline suppression through feature search by means of the P_D thus far have largely ignored the potential modulation by regularities across search displays (Theeuwes, Bogaerts, & van Moorselaar, 2022; van Moorselaar & Slagter, 2020). In this respect, it is noteworthy that in the work by Gaspelin and colleagues (Stilwell et al., 2022; Gaspelin & Luck, 2018a) examining proactive distractor suppression via the P_D not only the distractor feature (i.e., its color) but also the target features (i.e., both its color and shape) were fixed across trials. In other words, the experimental designs used by Gaspelin and colleagues strongly induced, be it via implicit learning (Theeuwes et al., 2022) or an explicit top-down process (Wolfe, 1994), predictions regarding both targets and distractors. It is therefore unclear to what extent the observed electrophysiological markers of distractor processing reflect pure distractorfeature suppression, as typically assumed, or alternatively, at least partly, also reflect target-feature upweighting (Chang & Egeth, 2019).

Although the P_D is exclusively elicited by to-be-ignored items, which seems to favor an interpretation in terms of suppression, there is reason to believe that it may in part also reflect upweighting of target features. As also acknowledged by updated versions of the signal

suppression hypothesis (Luck et al., 2021), distractor inhibition often results not only from proactive suppression of static and hence predictable distractor features (Vatterott & Vecera, 2012), but in part also reflects upweighting of predictable target features. For example, by modifying the capture-probe technique, a technique used to read out attentional processing across search displays, such that it could independently dissociate between distractor suppression and target feature upweighting, Chang and Egeth (2019, 2021) demonstrated that both distractor suppression and target feature upweighting contributed to distractor inhibition. Moreover, on the basis of a more systematic manipulation of color similarity between the target and neutral filler items in the display, Oxner et al. (2023) even concluded that apparent proactive distractor suppression could be entirely explained by global target feature enhancement. Although future work is necessary to further understand the interplay between suppressive and upweighting mechanisms in the context of distractor suppression, to the very least, these findings illustrate the importance of disambiguating target and distractor effects in P_D research. Given that in recent years the P_D and its modulation has been widely used by both sides of the attentional capture debate, it is critical to establish to what extent it represents pure distractor processing independent from learned expectations regarding task-relevant features, which to the best of our knowledge has not yet been done experimentally.

To examine the extent to which, if at all, the P_D component also reflects upweighting of target features, we adopted the paradigm used by Gaspelin and colleagues that reliably produces below baseline suppression and a robust distractor P_D (Stilwell et al., 2022; Gaspelin & Luck, 2018a), but varied whether target features were static across trials. Specifically, whereas in half of the experiment both the target and the distractor feature were fixed across trials (as in previous studies), in the other half of the experiment, target features (i.e., shape and color) varied randomly across trials (see Figure 1A). This design allowed us to establish whether the P_D as observed in previous studies reflects pure distractor suppression, or alternatively, at least partly, is driven by upweighting of static target features. Critically, if the P_D purely reflects distractor suppression, under the current conditions, the P_D component should not be modulated by cross-trial variation of the target features.

Open Practice Statement

Experiment and analyses were based on the OSF Preregistration template at Open Science Framework (https://osf .io/9827w/). Analyses that diverge from the preregistration are described as exploratory. De-identified data for all experiments along with the data-analysis scripts (custom Python 3 scripts) will be posted alongside the preregistration upon publication. All code for running the experiment will also be made available here.



Figure 1. Schematic of the experimental procedure and behavioral results. (A) On each that, a heterogeneous set of shapes was presented in a circular configuration around fixation. The target shape (i.e., diamond or circle) was cued by the shape of the fixation marker (see top left of the figure). In singleton present displays, one of the shapes had a unique color (red or green; counterbalanced across participants), which was fixed throughout the entire experiment. Participants (n = 24) were instructed to report the orientation of the line inside the shape cued by the fixation marker (i.e., diamond or circle). In the fixed-features condition (half of the experiment), this target shape, as well as the target color was static across trials. By contrast, in the mixed-features condition, both the target shape (diamond or circle) as well as the target color (red/green or gray) varied randomly across trials. Search stimuli were taken from the Stilwell et al. (2022) study on which the current study was based. (B) The singleton present (dashed bar) versus singleton absent (solid bar) displays ($m_{absent} = 792.0$; $m_{present} = 775.2$; $\Delta m = 16.8$; n = 24; two-tailed p < .001; d = 1.2; 95% CI [15.3, 31.1]) resulting in a unreliable Block Type × Singleton Presence interaction, F(1, 23) = 1.3, p = .27, $\eta_p^2 = .052$; BF_{excl} = 3.1. The height of each bar reflects the population average, and error bars represent 95% within-subject confidence intervals (Morey, 2008). Data from each participant are represented as gray dots, connected by solid lines (i.e., singleton presence benefit) or dashed lines (i.e., attentional capture).

METHODS

Participants

A planned number of 24 participants (mean age = 21 years, range = 19-26; 21 women), participated in the experiment, in exchange for course credit or monetary compensation (10 \in per hour). Sample size was based on our previous work (van Moorselaar et al., 2021) and previous studies on which the current study was based (Stilwell et al., 2022; Gaspelin & Luck, 2018a). Eight participants were replaced because they failed to maintain fixation during the window of interest in a large subset of trials (n = 6), because automatic artifact rejection resulted in removal of too many trials (n = 1), or because their accuracy deviated more than 2.5 SDs from the group mean (n = 1). All participants gave their informed consent before the start of the study, which was approved by the ethical review committee of the Faculty of Behavioural and Movement Sciences of the Vrije Universiteit Amsterdam.

Apparatus, Material, and Procedure

The experiment, which took place in a dimly lit room on a 23.8-in. ASUS ROG STRIX XG248 LED monitor with a 240-Hz refresh rate, was created using OpenSesame (Mathôt, Schreij, & Theeuwes, 2012) utilizing PsychoPy functionality (Peirce, 2009). Participants were positioned 60 cm away from the screen using a desk-mounted chinrest. The eyes were tracked on- and offline using an Eyelink 1000 (SR Research) eye tracker tracking the left eve with a 1000-Hz sampling frequency (one participant had a sampling frequency of 2000 Hz), and participants heard a beep each time fixation was broken by more than 2° of visual angle. At the start of the experiment, the eves were calibrated via a five-dot calibration procedure until spatial error for each dot position was smaller than 1° of visual angle. Drift correction was applied every 80 trials (i.e., at the start and halfway a block); when deemed necessary, the calibration procedure was repeated. EEG data were recorded at a sampling rate of 512 Hz with default settings using a 64-electrode cap with electrodes placed according to the 10-10 system (Biosemi ActiveTwo system; biosemi.com). VEOG/HEOG were recorded via external electrodes placed ~2 cm above and below the eve, and ~ 1 cm lateral to the external canthi, respectively.

The paradigm was modeled after Stilwell and colleagues (2022). Each trial started with a randomly jittered black display (100–400 msec) followed by a randomly jittered fixation display (750–1000 msec). This display contained either a circular or a diamond shape with an embedded cross hair, a combination that has been shown to

improve stable fixation (Thaler, Schütz, Goodale, & Gegenfurtner, 2013). Critically, the shape of the fixation marker signaled the target shape in the subsequent search display. Each search display contained eight shapes in a circular configuration (radius 3°) around the fixation marker, each with a black line tilted left or right (14° around the vertical plane) in their center. Individual shapes were selected from a stimulus pool of triangles (radius 0.7°), hexagons (1.2° by 1.2°), ovals (1.5° by (0.9°) , crosses $(1.2^{\circ} \text{ by } 1.2^{\circ})$, diamonds $(1.3^{\circ} \text{ by } 1.3^{\circ})$, and circles (radius 0.6°). Selection was such that each display contained the target shape (i.e., diamond or circle) and seven shapes randomly selected from the remaining shapes in the stimulus pool with replacement, with the restriction that each individual nontarget shape appeared 2 times at the most to ensure high display heterogeneity.

Individual shapes were either red (red, green, blue [RGB]: 253, 34, 34), green (RGB: 90, 174, 20), or gray (RGB: 146, 147, 153). In the fixed-feature condition, both the shape (circle or diamond; counterbalanced across participants) and the color of the target (red or green; counterbalanced across participants) were held constant. By contrast, in the mixed-feature condition, both the target shape and target color varied randomly across trials (counterbalanced across trials in a block) such that in half of the displays, the target color was gray, whereas in the other half, the target color matched the target color in the fixed-feature condition. Whereas in 25% of trials all stimuli in the search display had the same color (i.e., singleton absent displays), in the remaining trials (i.e., singleton present displays), one of the nontarget shapes was rendered in a unique color (red or green; counterbalanced across participants) that was held constant throughout the experiment. These singleton absent and present displays were not randomly intermixed, but instead singleton absent displays were grouped together at the start or at the end of an experimental block (alternating between blocks) such that any singleton presence benefit could not be attributed to a surprise induced by infrequent distractor absent displays. Across all display configurations, targets and singleton distractors appeared with equal probability selectively at positions along the vertical and horizontal axis.

At the beginning of each session, it was made explicit that the distractor singleton was irrelevant to the task at hand and should thus be ignored. Participants were instructed to keep their eyes at fixation and covertly search for the shape that matched the fixation shape on the current trial and indicate the orientation of the line segment within this target shape via button press (i.e., "Z" or "/" button) as quickly as possible while keeping the number of errors to a minimum. In case of an incorrect response or missing response, a 200-Hz tone lasting 300 msec was played, which was accompanied by the text "too slow!" in case participants did not respond within 2000 msec.

The experiment consisted of 12 experimental blocks of 160 trials (six consecutive blocks for each condition; order

counterbalanced across participants) preceded by sequences of 32 training trials in the mixed-feature condition, which repeated until mean accuracy was above 70%. At the start of each new block, participants were informed about the dynamics of the upcoming block (mixed-feature or fixed-feature condition) and whether distractor present/absent displays were grouped at the start or at the end of the upcoming block. Halfway between each block, there was a 15-sec mandatory break to rest the eyes. After each block, participants received feedback on their performance (i.e., mean RT and accuracy).

Behavioral Analysis

All data were preprocessed in a Python environment (Python Software Foundation, https://www.python.org/). Analyses were limited to RT data of correct trials only. RTs were filtered in a two-step trimming procedure: Trials with RTs shorter than 200 msec were excluded, after which data were trimmed based on a cutoff value of 2.5 *SD*s from the mean per participant. Remaining RTs were analyzed with repeated-measures ANOVAs with within-subject factors Block Type (fixed-features, mixed-features) and Singleton Presence (present, absent), followed by planned comparisons with paired *t* tests using JASP software (JASP Team, 2023). In case of insignificant interactions, we also report BF_{excl}, which reflects the comparison between the interaction and equivalent models stripped of the effect.

EEG Preprocessing

EEG data, which was rereferenced offline to the average of the left and right earlobe, was first high-pass filtered using a zero-phase "firwin" filter at .1 Hz to remove slow drifts. Continuous EEG was subsequently epoched from -700 to 1100 msec relative to search display onset (to avoid filter artifacts during automatic artifact rejection; see below). Before trial rejection, malfunctioning electrodes as identified during recording (M = 0.6, range = 0-2) were temporarily removed. As a first artifact removal step, independent component analysis as implemented in MNE (method = "picard") was performed on 1-Hz filtered epochs to remove eye-blink components selectively from the 0.1-Hz filtered data. Next, noise-contaminated epochs within a -200 to 600 msec were identified using an adapted version of an automatic trial-rejection procedure. To specifically capture muscle activity, the EEG signal was filtered using a 110- to 140-Hz band-pass filter and subsequently transformed into z scores. A subject-specific z-score threshold was then set based on within-subject variance of z scores (de Vries, van Driel, & Olivers, 2017). Moreover, to reduce the number of false alarms, rather than immediate removal of epochs exceeding the z-score threshold, per marked epoch, the five electrodes that contributed most to accumulated z score within the time period containing the marked artifact were identified.

Then in an iterative procedure, the worst five electrodes per marked epoch were interpolated using spherical splines (Perrin, Pernier, Bertrand, & Echallier, 1989) one by one, checking after each interpolation whether that epoch still exceeded the determined *z*-score threshold. Epochs were selectively dropped when, after this iterative interpolation procedure, the *z*-score threshold was still exceeded. Finally, malfunctioning electrodes were interpolated using spherical splines (Perrin et al., 1989).

ERP Analyses

ERP analyses were limited to trials without identified eve movements. For this purpose, trials with a fixation deviation $> 1^{\circ}$ of visual angle correction in a segment of continuous data of at least 40 msec in the time window -200 to 400 msec after drift correction using prestimulus data (van Moorselaar & Slagter, 2019) were excluded. We focused the analysis of distractor- and target-elicited ERP waveforms on electrodes PO7/8, which were chosen a priori based on previous studies examining the P_D (Stilwell et al., 2022; Gaspelin & Luck, 2018a). Epochs were baseline corrected using a -200- to 0-msec prestimulus baseline period. To enable isolation of lateralized distractor- and target-specific components, the analyses focused on trials in which the stimulus of interest (distractor or target) was presented to the left or right of fixation, whereas the other stimulus was presented on the vertical meridian or absent. Waveforms evoked by the various search displays were collapsed across left and right visual hemifield and left and right electrodes to produce separate waveforms for contralateral and ipsilateral scalp regions. Lateralized difference waveforms were then computed by subtracting the ipsilateral waveform from the corresponding contralateral waveform. Time windows of interest were centered around the positive peak $(\pm 55 \text{ msec})$ of the grand mean waveform (i.e., averaged across conditions) in distractor-tuned analyses, and around the negative peak $(\pm 37.5 \text{ msec})$ of the grand mean waveform in target-tuned analyses.

RESULTS

Search Times

Exclusion of incorrect responses (7.5%) and data trimming (2.9%) resulted in an overall loss of 10.5% of behavioral data. Figure 1A depicts the mean RTs for singleton present and absent displays in the two search conditions. Efficiency of target selection appeared to not differ as a function of whether target features were static or varied randomly across trials, F(1, 23) = 2.57, p = .12, $\eta_p^2 = .10$; BF_{excl} = 0.24. Critically, there was a clear benefit when the search array contained a singleton distractor (main effect Singleton Presence: F(1, 23) = 26.84, p < .001, $\eta_p^2 = .54$), independent of whether the target features (i.e., shape and color) were fixed or varied randomly

across trials (interaction Singleton Presence and Block Type: F(1, 23) = 1.26, p = .27, $\eta_p^2 = .052$; BF_{excl} = 3.1). Replicating previous studies using heterogeneous search displays (Stilwell et al., 2022; Chang & Egeth, 2019; Gaspelin & Luck, 2018a), in the fixed-feature condition, there was a reliable ~17-msec singleton-presence benefit, t(23) = 3.01, p = .006, d = 0.61. This singletonpresence benefit was not reduced, but, if anything, larger (~23 msec) and more reliable, t(23) = 6.05, p < .001, d = 1.24, when target features varied randomly across trials and, hence, the trial structure provided less opportunity to upweight relevant features for the upcoming search. Error rates did not reliably differ between singleton present and singleton absent trials, neither in fixedfeatures nor in mixed-feature conditions (interaction Search Condition and Block Type: F(1, 23) = 0.038, p =.85, $\eta_p^2 = .002$; BF_{excl} = 3.56).

To further explore the effect of randomly switching target features, in the mixed-features condition, we separated trials in which none of the target features, the target color, the target shape, or both target features repeated from one trial to the next. This analysis yielded no evidence that the singleton presence benefit was modulated by intertrial target feature priming (interaction Singleton Presence and Target Feature Repetition: F(3, 69) = 0.95, $p = .42, \eta_p^2 = .040; BF_{excl} = 5.17).$ Altogether, the behavioral results indicate that selection of the target, as well as ignoring the static singleton distractor color, remained equally effective when target features varied randomly across trials compared with a condition with static target features. Given the putative roles of the N2pc, reflecting covert attentional selection (Luck, 2012; Woodman & Luck, 2003), and the P_D , reflecting suppression (Gaspelin & Luck, 2018a, 2018b), we should thus also not expect differences in the N2pc elicited by targets and, of especially interest here, the P_D elicited by distractors between mixed- and fixed-feature conditions.

Electrophysiological Results

We first examined the waveforms elicited by lateralized targets to characterize attentional selection across conditions. Previous studies examining target selection in heterogeneous search displays identified no reliable differences between difference waveforms elicited by lateral targets either with or without singleton distractors (Stilwell et al., 2022; Gaspelin & Luck, 2018a), as was also the case here. Therefore, in target-tuned analysis (see Figure 2), we collapsed across singleton absent trials and trials with a distractor on the vertical midline (individual waveforms for distractor absent and distractor present displays are shown in Appendix Figures A1 and A2). As expected, lateralized targets elicited a more negativegoing deflection in the contralateral compared with the ipsilateral waveform beginning at approximately 200 msec after search display onset. In line with behavior, this N2pc





attenuated, but nevertheless reliable, when target features varied randomly relative to a condition with static target features ($m_{\text{fixed}} = 0.57 \,\mu\text{V}$; $m_{\text{mixed}} = 0.25 \,\mu\text{V}$; $\Delta m = 0.32$; n = 24; two-tailed p = .031; d = 0.47; 95% CI [0.04, 0.89]).

component appeared with approximately the same amplitude and time course when the target features were fixed across trials (i.e., fixed-features condition), and when they varied randomly across trials (i.e., mixed-features condition).

The N2pc components were measured as the mean amplitudes from 261 to 336 msec post stimulus and subsequently analyzed using a repeated-measures ANOVA with within-subject factors Block Type (mixed-features, fixed-features) and Hemifield (contralateral to target, ipsilateral to target). This analysis confirmed that although the N2pc was reliable across conditions (main effect Hemifield: $F(1, 23) = 28.58, p < .001, \eta_p^2 = .55$), there was no evidence that it differed between block types (interaction Block Type and Hemifield: $F(1, 23) = 0.019, p = .89, \eta_p^2 < .001$; BF_{excl} = 3.91). Indeed, planned pairwise comparisons comparing contralateral versus ipsilateral waveforms yielded reliable differences both in the fixed-features, t(23) = 4.27, p < .001, d = 0.87, and the mixed-features, t(23) = 4.97, p < .001, d = 1.01, conditions.

Although the target-tuned analysis, mimicking the observed behavior, thus did not identify marked differences between conditions, somewhat surprisingly waveforms elicited by distractors appeared to be modulated by target feature regularities (see Figure 2). Like previous studies (Stilwell et al., 2022; Drisdelle & Eimer, 2021; Gaspelin & Luck, 2018a; Sawaki & Luck, 2010), the positive difference elicited by lateral distractors, corresponding to the P_D , had an earlier onset than the N2pc elicited by targets, seemingly consistent with the idea that the P_D signals proactive suppression (Gaspar & McDonald, 2014; Jannati et al., 2013; Sawaki & Luck, 2010). Yet, at odds with this idea, the P_D appeared to be attenuated when target features were no longer fixed across trials, suggesting that processing of singleton distractor as signaled by the P_D , at least to some extent, also reflects target feature upweighting rather than a pure suppressive process.

The P_D components were measured as the mean amplitudes from 174 to 284 msec post stimulus onset and subsequently analyzed in the same way as the N2pc components. Replicating previous studies, there was a reliable lateralized positivity elicited by singleton distractors (main effect hemifield: F(1, 23) = 11.60, p = .002, $\eta_p^2 =$.34. Critically, although planned pairwise comparisons showed that this P_D component was evident in both the fixed- (0.5 µV; t(23) = 3.38, p = .003, d = 0.69) and the mixed-features condition $(0.2 \ \mu\text{V}; t(23) = 2.49, p = .022, d = 0.50)$, a Block Type × Hemifield interaction confirmed that the P_D was reliably attenuated when target features were not static, but instead varied randomly across trials, $F(1, 23) = 5.29, p = .031, \eta_p^2 = .19)$.¹ These findings demonstrate that when observers could not predict the target color in advance and were not consistently searching for the same shape, the P_D elicited by singleton distractors was reduced, but nevertheless reliable.

Exploratory Analyses

Visual inspection of Figure 2 suggests that the observed modulation of the P_D was especially apparent in the early time window of the P_D. Given that distractor positivity often contain an early and a late component (van Moorselaar et al., 2021; Feldmann-Wüstefeld & Vogel, 2019; Weaver et al., 2017), which have been speculatively linked to different cognitive processes, we explored whether temporal dynamics of the PD differed between conditions. For this purpose, we analyzed the area under the curve in the contralateral versus ipsilateral difference waveform with a repeated-measures ANOVA with withinsubject factors Time Window (early: 100-200 msec, late: 200-35 = 00 msec) and Block Type (fixed-features, mixed-features). Here, we focused on area under the curve rather than mean amplitude as this method is less sensitive to potential latency differences between individuals (note that mean amplitude yielded the same pattern of results). This analysis yielded no reliable interaction, $F(1, 23) = 0.002, p = .97, \eta_p^2 < .001; BF_{excl} = 3.70, sug$ gesting that the observed modulation was uniform across time. This was also confirmed by an exploratory jackknife procedure (Miller, Patterson, & Ulrich, 1998) that did not identify a difference in onset latency between fixed- and mixed-feature conditions (threshold = 50% of maximum

amplitude; onset_{fixed} = 140 msec, onset_{mixed} = 130 msec; t(23) = 0.08, p = .94).

The results thus far indicate that varying the target features (i.e., color and shape) randomly across trials uniformly attenuates the positivity elicited by lateral distractors in the typical P_D window. This raises the question to what extent the P_D in fixed-feature conditions as typically observed reflects sensitivity to target regularities across longer time scales above and beyond intertrial priming effects. Although our experiment was not designed to specifically target this question, in an exploratory analysis, we aimed to address this by examining the effects of feature repetition in the mixed-feature condition. Although behaviorally there was little to no difference between different forms of intertrial feature priming on the singleton presence benefit, in terms of the P_D, the effect of intertrial color repetitions appeared most pronounced. For this purpose, we focused on the comparison of trials with and without repetition of the target color in the mixedfeatures condition. As visualized in Figure 3, the observed pattern of results was reminiscent of the main pattern of results comparing fixed- and mixed-feature conditions (see Figure 2), with no modulation of the N2pc, but an apparent attenuation of the P_D when the target color did not repeat from one trial to the next. Indeed, whereas the N2pc elicited by lateral distractors was highly reliable irrespective of intertrial color priming (all ts > 4.33, all ps <.001, all ds > 0.88), the distractor P_D reached significance on trials where target colors repeated (0.34 μ V; t(23) =2.51, p = .019, d = 0.51), but not on trials where the target color switched (0.16 μ V; t(23) = 0.98, p = .34, d = 0.20). Note however that this pattern of results should be interpreted cautiously as the Prime (color repeat, color switch) × Hemifield interaction did not reach significance, $F(1, 23) = 0.60, p = .45, \eta_p^2 = .025; BF_{excl} = 1.95, and$ future research is thus necessary to establish whether

Figure 3. Target color repetitions appear to modulate the distractor $P_{\mathrm{D}}\boldsymbol{.}$ Waveforms depict difference between contra- and ipsilateral waveforms at PO7/8 for targettuned (solid lines) and distractor-tuned analyses (dashed lines), separately for color repeat (black) and color switch (gray) displays within the mixed-feature condition. Shaded areas reflect the time windows of interest for the N2pc (blue) and the P_D (red) analyses. The N2pc elicited by lateral targets did not differ between color repeat and color



switch display ($m_{\text{repeat}} = -1.0 \,\mu\text{V}$; $m_{\text{switch}} = -1.0 \,\mu\text{V}$; $\Delta m = 0.0$; n = 24; two-tailed p = .95; d = 0.014; 95% CI [-0.41, 0.39]). By contrast, the P_D elicited by distractors appeared to be reliably attenuated when the target color switched, although this was not supported by a reliable difference between mean amplitudes in the P_D window ($m_{\text{repeat}} = 0.34 \,\mu\text{V}$; $m_{\text{switch}} = 0.16 \,\mu\text{V}$; $\Delta m = 0.18$; n = 24; two-tailed p = .45; d = 0.16; 95% CI [-0.25, 0.56]). The waveforms in this plot were smoothed by a third-order polynomial (window length = 51) to improve the visibility of the effects but were analyzed using unsmoothed waveforms.

target color repetitions are by itself sufficient to modulate the P_D in response to a static color singleton.

DISCUSSION

The aim of the present study was to better characterize the P_D , which has become one of the major tools to study suppression of salient distractors. Specifically, we aimed to examine the extent to which the P_D is driven by taskrelevant regularities in designs typically used to study proactive distractor suppression (Stilwell et al., 2022; Chang & Egeth, 2019; Gaspelin & Luck, 2018a). To this end, we modified a heterogenous shape version of the additional singleton paradigm (Theeuwes, 1991, 1992), which has been shown to reliably produce both behavioral and electrophysiological signatures of distractor suppression (Stilwell et al., 2022; Gaspelin & Luck, 2018a), such that target features were static in one half of the experiment (i.e., fixed-features condition), whereas they varied randomly in the other half of the experiment (i.e., mixedfeatures condition). This design made it possible to examine how processing of a fixed distractor color in a display of heterogenous shapes is shaped by expectations regarding the upcoming target features. We found that target selection was equally effective in both conditions, both in terms of manual response times, as well as in terms of the N2pc components. In contrast, although both conditions resulted in a singleton-presence benefits, the P_D elicited by lateralized distractors was reliably attenuated when the target features varied randomly across trials. This result demonstrates that the $P_{\rm D}$ elicited by distractors cannot unequivocally be attributed to a suppressive process, as it can also, at least when the experimental design allows, reflect upweighting of target features.

The present findings have important implications for the attentional capture debate, and specifically for studies that examine proactive distractor suppression via the P_{D} component. As noted in the introduction, below baseline suppression, a defining marker of proactive suppression, to date has been selective to experimental designs using heterogeneous shapes in which not only the distractor color but also the defining target features (i.e., shape and color) are fixed across trials. Although it is notoriously difficult to empirically distinguish between feature-specific upweighting and suppression (Gaspelin & Luck, 2018b), the consensus, as also acknowledged by recent formulations of the signal suppression hypothesis (Luck et al., 2021), is that under such conditions, both upweighting and suppression concurrently guide attention (Hamblin-Frohman, Chang, Egeth, & Becker, 2022; Chang & Egeth, 2019, 2021; Vatterott & Vecera, 2012). Here, for the first time, we demonstrate an electrophysiological correlate of this interplay between suppression an upweighting, by showing that the P_D is attenuated when target features are no longer static but instead vary randomly across trials. This attenuation of the P_D under conditions that discourage target feature upweighting is consistent with a hybrid

model of distractor processing, in which distractor inhibition is driven by two independent mechanisms. At the same time, it should be noted that it has even been argued that observed distractor inhibition can be exclusively explained by target feature upweighting (Oxner et al., 2023; Saenz, Buracas, & Boynton, 2002). In the current experimental design, when target features varied, they only did so between two possible options, leaving open the possibility that the remaining part of the P_D was still driven by target feature upweighting, but simply less pronounced upweighting given that in mixed-feature conditions, it now had to be distributed across color space.

Although our results are consistent with the idea that the P_D elicited by distractor in heterogeneous search displays with static target features is sensitive to task-relevant regularities, the exact underlying mechanism remains elusive. As outlined above, one possibility is that the P_D reflects both enhancement and suppression, such that the amplitude increases when the task also allows for secondary inhibition resulting from an upweighting of predictive target features (van Moorselaar & Slagter, 2020; Noonan, Crittenden, Jensen, & Stokes, 2018). Indeed, there is evidence that target representations can be strategically shifted off-veridical to optimally distinguish targets from distractors (Geng & Witkowski, 2019). Alternatively, it could be argued that the P_D purely reflects suppression, if one assumes that task-relevant regularities influence how the distractor is encoded in relation to the other items in display (Becker, 2010). Whereas in the mixed-feature condition observers cannot rely on a relational distractor code and hence must rely on an absolute code, in the fixed-feature condition, relational coding is possible to potentially strengthen distractor suppression. We believe this less likely, however, given the behavioral evidence in support of target enhancement driving distractor inhibition in the current paradigm (Oxner et al., 2023; Chang & Egeth, 2019, 2021). Irrespective of the underlying mechanism, however, the current results highlight the importance of taking target regularities into consideration when examining distractor suppression via the $P_{\rm D}$.

The observation that the electrophysiological response elicited by salient distractors is modulated by target predictability is also consistent with previous ERP studies relying on homogeneous instead of heterogeneous search displays. When the target is defined as the unique shape in the display (e.g., a diamond among circles or vice versa), there is some disagreement on the presence of a distractor P_D when the target shape varies unpredictable across trials, with some studies actually reporting an N2pc (Wang et al., 2019; Burra & Kerzel, 2013; Hickey, McDonald, & Theeuwes, 2006) suggesting attentional capture rather than distractor suppression (but see van Moorselaar et al., 2021; McDonald et al., 2013). By contrast, when the unique target shape is fixed, favoring feature search mode (Bacon & Egeth, 1994), not only the target N2pc increases, but distractors also reliably elicit a P_D (van Moorselaar et al., 2021; Burra & Kerzel, 2013). It should be noted, however, that although a P_D signals suppression of distractors (Forschack, Gundlach, Hillyard, & Müller, 2022), in these displays, distractors typically continue to interfere with attentional selection. To resolve this apparent discrepancy, one could argue that suppression was in place, but insufficiently so to counteract bottom-up attentional capture. Alternatively, the P_D may not necessarily signal suppression of a salient stimulus, but instead processing of a salient, yet irrelevant feature that does not require a further read out. Although highly speculative, such a framework would predict that any salient distractor would generate a P_D, as long as the distractor does not resemble the target. In these circumstances, the initial capture by the salient distractor does not require the formation of an object representation of the distractor as it can be immediately discarded as "not being the target." Indeed, there is evidence that attention can be disengaged very rapidly if the distractor does not resemble the target (Born, Kerzel, & Theeuwes, 2011; Mulckhuyse, Van der Stigchel, & Theeuwes, 2009). In this framework, the P_D can still be envisioned as the mirror image of the N2pc, but not because they reflect covert shifts of attention (Eimer, 1996) or suppression (Gaspelin & Luck, 2018b), but instead indexing object individuation (Foster, Bsales, & Awh, 2020; Mazza & Caramazza, 2015) on the one hand in case of an N2pc and ignoring of specific features on the other hand in case of the P_D .

An important caveat in the interpretation of our results is that targets and neutral nontargets always shared the same color and, hence, their effects cannot be disambiguated. Therefore, rather than attributing the observed P_D attenuation to an upweighting of static target features, an alternative explanation is to assume that the neural adaptation to repeated colors over time results in a reduced the inter-item competition between nontarget items (Adam & Serences, 2021; Solomon & Kohn, 2014). According to such a framework, however, the singleton should become increasingly less salient over time in the fixed-feature condition, which appears at odds with the observation that behaviorally the effect did not differ between conditions, and the fact that PD decreased rather than increased in the mixed-feature condition where neural adaptation across trials should be less pronounced. Nevertheless, future work examining to what extent the PD is sensitive to task-relevant regularities should also take nontarget regularities into account and also consider that the early positivity that we chose to label P_D in the current study overlaps with the Ppc component, which is proposed to be sensitive to feature discontinuities within search displays (Gokce, Geyer, Finke, Müller, & Töllner, 2014; Fortier-Gauthier, Moffat, Dell'Acqua, McDonald, & Jolicœur, 2012).

Our behavioral results indicate a finding what has been considered to be "a striking reversal of the capture effect" (Chang & Egeth, 2019) as observers were significantly faster on distractor present trials than on absent trials. This reversal of the capture effect has now been reported in several recent studies (Ma & Abrams, 2023; Lien, Ruthruff, & Hauck, 2022; Stilwell & Gaspelin, 2021; Gaspelin et al., 2015). Although some may consider this reversal as surprising, this finding is consistent with our claim that during feature search in which the target is typically nonsalient, the attentional window is adjusted to keep the discriminability of the target to an acceptable signal-tonoise-ratio (Liesefeld & Müller, 2021; Theeuwes, 2004, 2010). Because of the small attentional window, search proceeds serially and during serial search, there is no attentional capture by the salient distractor singleton. Because they are in serial search, participants can immediately discard the distractor that gives them one less item to inspect in distractor present than in absent trials (see Theeuwes, 2010, 2022 for a discussion). Consistent with this notion is our speculation that in these circumstances, in which search is serial, the distractor will generate a P_D because it can be immediately discarded as irrelevant and "not being the target."

In summary, the current study clearly demonstrates that in displays that encourage feature search mode, the P_D cannot unequivocally be attributed to suppression of distractor features, as at least in part, it is also sensitive to target regularities. This finding has large implications for future studies that use the P_D to examine whether specific distractor features can be proactively suppressed.

APPENDIX A



Figure A1. Target elicited waveforms in distractor absent and distractor present displays in the fixed-features condition. (A) Electrophysiological results from search trials with lateral targets without a distractor (left) and with lateral targets accompanied by a distractor on the vertical midline (right). Ipsilateral (dashed lines) and contralateral (solid lines) waveforms reflect activity at electrode sides PO7/8. (B) Difference waveforms between contra- and ipsilateral waveforms for target tuned waveforms on distractor absent (solid lines) and distractor present displays (dashed lines). The shaded area reflects the time windows of interest for the N2pc analyses). Despite the apparent numerical difference, the N2pc elicited by lateral targets did not differ between distractor absent and distractor present displays ($m_{absent} = -0.54 \ \mu$ V; $m_{present} = -1.0 \ \mu$ V; $\Delta m = -0.5$; n = 24; two-tailed p = .12; d = 0.33; 95% CI = -1.14-0.14).



Figure A2. Target elicited waveforms in distractor absent and distractor present displays in the mixed-features condition. (A) Electrophysiological results from search trials with lateral targets without a distractor (left) and with lateral targets accompanied by a distractor on the vertical midline (right). Ipsilateral (dashed lines) and contralateral (solid lines) waveforms reflect activity at electrode sides PO7/8. (B) Difference waveforms between contra- and ipsilateral waveforms for target tuned waveforms on distractor absent (solid lines) and distractor present displays (dashed lines). The shaded area reflects the time windows of interest for the N2pc analyses). The N2pc elicited by lateral targets did not differ between distractor absent and distractor present displays ($m_{absent} = -0.90 \ \mu$ Y; $m_{present} = -0.77 \ \mu$ Y; $\Delta m = -0.13$; n = 24; two-tailed p = .52; d = 0.14; 95% CI = -0.54-0.28).

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Data Availability Statement

Experiment and analyses were based on the OSF Preregistration template at Open Science Framework (https://osf .io/9827w/). Analyses that diverge from the preregistration are described as exploratory. Deidentified data for all experiments along with the data-analysis scripts (custom Python 3 scripts) have been posted alongside the preregistration upon publication. All code for running the experiment has also been made available here.

Author Contributions

Dirk van Moorselaar: Conceptualization; Data curation; Formal analysis; Investigation; Methodology; Resources; Software; Visualization; Writing—Original draft. Changrun Huang: Data curation; Investigation; Methodology; Project administration; Writing—Review & editing. Jan Theeuwes: Conceptualization; Funding acquisition; Methodology; Supervision; Writing—Review & editing.

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Diversity in Citation Practices

Retrospective analysis of the citations in every article published in this journal from 2010 to 2021 reveals a persistent pattern of gender imbalance: Although the proportions of authorship teams (categorized by estimated gender identification of first author/last author) publishing in the *Journal of Cognitive Neuroscience* (*JoCN*) during this period were M(an)/M = .407, W(oman)/M = .32, M/W = .115, and W/W = .159, the comparable proportions for the articles that these authorship teams cited were M/M = .549, W/M = .257, M/W = .109, and W/W = .085 (Postle and Fulvio, *JoCN*, 34:1, pp. 1–3). Consequently, *JoCN* encourages all authors to consider gender balance explicitly when selecting which articles to cite and gives them the opportunity to report their article's gender citation balance.

Note

1. The same pattern of results was obtained when time windows of interest were centered around condition-specific positive peaks in distractor-tuned waveforms (177–287 msec and 171–282 msec for fixed- and mixed-features conditions, respectively), or alternatively when rather than a data-driven approach, time-windows matched the P_D window (i.e., 115–225 msec) reported in Stilwell and colleagues (2022).

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