BRIEF REPORT

Spatial context learning in Pigeons (*Columba Livia*)

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In a seminal paper in the cognitive sciences, Chun & Jiang (1998) described the Contextual Cuing (CC) paradigm in which they used artificial stimuli and showed that people became faster to locate a target when the background predicted the location of a target compared to when it did not. Here examined contextual cuing in pigeons for the first time using artificial stimuli and procedures similar to those of Chun & Jiang. In the first test we had pigeons search for a target among a display of 7 distractors; during one condition the position of the distractors predicted the location of the target and in the second condition there was no relationship between the two. In a second test we presented the pigeons with the predictive displays from Test 1 and a second set of displays that also predicted the location of a target to see if learning about one set of predictive backgrounds disrupted learning about a second set. The pigeons were quick to acquire context based knowledge and retain that information when faced with additional contexts. The results suggest that contextual cuing can occur for a variety of stimuli in non-human animals and that it may be a common mechanism for processing visual information across a wide variety of species.

Keywords: contextual cuing, visual search task, pigeons, predictive scene learning

When an animal moves through an environment it may perceive a tremendous variety of colors, shapes, textures, and line orientations that comprise the elements of a visual scene. Yet, despite the complexity of an observed scene animals are able to identify objects that are important for their survival (e.g., prey, mate) from the background features of the scene and respond appropriately. Animals use cognitive mechanisms such as perception, attention, and memory, to help them efficiently identify these important stimuli (see Shettleworth, 2010 for a review).

The investigation of the cognitive mechanisms that animals use to visually search for targets like cryptic prey has long been the basis for interdisciplinary work by both psychologists and evolutionary biologists. For instance, Luuk Tinbergen (1960) proposed that a foraging bird that repeatedly encountered the same prey type (more likely for populations with high densities) developed a search image – a remembered representation of the features of the prey - that allowed it to be more efficient during search. That is, the birds were more sensitive (primed) to the features of a prey after recently seeing similar conspecifics. Subsequent laboratory based work using multiple avian species and methods confirmed the search image hypothesis (Blough, 1989, 1991, 1992; Bond, 1983; Langley, Riley, Bond & Goel, 1995; Pietrewicz and Kamil 1979, 1981; Reid & Shettleworth, 1992).

The emphasis of researchers studying the search image and related phenomena has largely been on the features of the prey/target and how they are encoded into memory. However, several sources of information
may be available to a predator in an environment to indicate the presence and perhaps the location of prey. One additional source of information that might be used is the nature of the habitat comprising the background of the visual scene. The information in a background is not typically random and tends to show regularities (Gibson, 1963). Researchers have historically been less interested in the background of the search scene and have treated it more as ‘noise’ that can be used to increase or decrease the crypticity of prey during the search process (though see Kono, Reid & Kamil, 1998 for example). Our knowledge about how the background or contextual information is acquired and how it can direct search in non-human animals, independent of the search image, is surprising limited.

In a seminal paper in the cognitive sciences, Chun & Jiang (1998) described the Contextual Cuing (CC) paradigm in which they studied how visual background/context information is acquired and how that learning is expressed. Chun and Jiang employed a visual search task in which human participants searched for a rotated letter T, the bottom of which pointed to the left or right (i.e., ← or →). The target was surrounded by an array of distractors (letter Ls), which were each randomly rotated. Participants were asked to find the target and discriminate, via manual response, whether the T was rotated to the left or right. Chun and Jiang created conditions in which participants would learn that repeated spatial configurations of distractors were paired with target locations. For each participant, the investigators devised a set of 12 invariant or “old” context configurations in which the spatial locations of target and distractor locations were fixed. These 12 configurations were each inserted, in random order, into every block of 24 trials. For the remaining 12 trials in each block, the spatial locations of targets and distractors were generated randomly (“new” context). Chun and Jiang hypothesized that, after repeated exposure to the “old” configurations, participants would learn that each of these configurations reliably predicted the target location – and that the participants would use these associations to facilitate search, compared to the “new” condition. Indeed, the results showed that participants became faster to locate the targets in the old context condition compared to the new context condition after a few exposures (though later tests indicated that participants had no explicit knowledge of the old context displays). Thus, contextual information provided by the spatial layout of a set of distractors can cue attention to a target and direct a behavioral response (see also Chun, 2000; Chun & Jiang, 1999, 2003; Chun & Phelps, 1999). The contextual cuing effect has also been extended to research with more naturalistic stimuli. For example, Brockmole and Henderson (2006) had participants search for a target in images of real-world scenes. The participants showed a steady decline in the amount of time it took to find the target during repeated images with the target, but not for novel target-image configurations. Thus, regularities in background features of natural scenes also can produce contextual cuing.

Limited recent work on how context can direct search has been conducted with non-human primates and pigeons. Goujon & Fagot (2013) have explored contextual cuing in Baboons using search task procedures comparable to Chun & Jiang. Baboons, like the human participants reported by Chun and Jiang, were faster to respond to the target (T) when the background was predictive compared to when it was not predictive. Some researchers also have recently examined if non-human animals also show contextual cuing for more naturalistic stimuli. Wasserman, Teng, & Castro (2013) had pigeons find a target that was presented on a solid color background or a colored image of a real-world photograph. The target could be located in one of the four corners of the display on the background. On 80% of the trials the particular background (either color or real-world image) was paired with an invariant location; on the other 20% of the trials the background-target location relationship was determined randomly. The birds were faster to respond to the target when it was predicted by the background (context) than when the target location was determined randomly, demonstrating a predictive effect of context. Additionally, the real-world background exerted higher rates of contextual cuing compared to the solid color background.

In the current study we used procedures comparable to those of Chun and Jiang (1999) in two separate tests to examine if pigeons show contextual cuing. We used Chun and Jian’s procedures so that we could isolate the spatial configuration information potentially driving contextual cuing, rather than other factors like conspicuousness or scene categorization that might occur when using natural scenes (see Chun & Jian, 1998).
The distinction is similar to work in animal navigation. Some researchers examine navigation in the natural environment, but it can be difficult to isolate which cues control behavior. In contrast, other researchers study spatial cognition using laboratory-based tests such as the Morris water arena where spatial cues presented to the animal can be better controlled (see Shettleworth, 2010, Chapter 8). The current study is important for several reasons. As mentioned above, work on contextual cuing in non-human animal is limited. Contextual cuing is an important mechanism for people to effectively search visual scenes (Chun, 2000). Pigeons have an acute visual system that follows a different lineage than that of mammals, and they utilize it to search for food in a rich visual environment. While mammals and their avian counterparts have different visual systems, both groups may demonstrate contextual cuing given common search demands in their respective environments. Researchers have successfully used pigeons in a variety of visual search tasks and past work has shown that people and non-human animals like pigeons may process visual information in qualitatively the same way. The same may be true for contextual cuing. The current work, represents an important step in address these questions.

**Method**

**Animals.** We used seven individually housed adult white Carnoux pigeons. We restricted the weight of each pigeon during the course of the study to 85% of its ad lib value. During daily experimental sessions the birds obtained mixed grain as reinforcement. The experimenter weighted the birds following the conclusion of an experimental session and provided a supplement of mixed grain if necessary to keep a bird at its target weight. The birds had free access to water and grit.

**Apparatus.** We trained and tested the pigeons in an operant chamber that had a 17” high-resolution (1024x768) monitor inserted through the front wall. An infrared touchframe (Elotouch, Menlo Park, CA) surrounding the monitor recorded the Cartesian location of the bird’s responses to stimuli presented on the monitor. We used a feeder in the back wall of the chamber to deliver momentary (~2s) access to mixed grain following selected responses. We paired a tone with the delivery of food to indicate its availability and to serve as a secondary reinforcer.

**Discrimination Training.** We presented a black letter T (40 x 40 pixels) on a light grey background on the monitor. The target was rotated on center so that the tail (i.e., bottom part) of the T was either to the left or the right (see Figure 1) during a trial. The direction of the tail of the target was block randomized. Following a peck to this stimulus, left and right rectangular choice keys appeared to the left and right side of the target, respectively. For half of the birds a peck to the left key was considered a correct choice when the tail of the T was to the right, whereas a response to the right key was considered incorrect. When the tail of the T was to the left the opposite relationship held true, that is a response to the right key was correct and a response to the left incorrect. For the other half of the birds the relationship between the direction of the tail of the target and the side of the key that was correct was reversed. Reinforcement was delivered following a correct response (always the tone, but food with 50% probability determined randomly). An incorrect response resulted in the screen going black for 5 seconds, followed by the initiation of a correction trial, which was a repetition of the same stimulus display used on the previous incorrect trial. The pigeon encountered correction trials until it made the correct response. Each daily session consisted maximally of 20 blocks of 16 trials for a total of 320 trials. Training continued until the birds made 80% correct responses to both the left and right tailed T during a single session. The birds were also presented with the target among a seven Os on the display that served as distractors. The procedures for the intermediate phase were identical to those described above and were designed to prepare the birds for encountering more challenging distractors during testing. Because the target was quite different than the distractors the birds only required one or two sessions of this intermediate training before they reached criterion and were move onto testing.

**Test 1:** Each bird was presented with the target (tail to the left or right) that must be found in a group of 7 L-shaped distractors (see Figure 1). During each block of 16 trials we presented the birds with two different types of trials. During the eight Novel condition trials we randomly positioned the seven distractors and the target in eight cells of a virtual 64 cell square grid overlaying the monitor. The position of the distractors and the target
within a cell was adjusted slightly (~ 10 pixels) to the north, south, east or west (randomly determined). The 64-cell grid was divided into four quadrants (up, down, left and right); the position of the target was randomly assigned to a cell in one of the four quadrants. The position of the target was block randomized in each of the four quadrants across the trials of a daily session. The location of the seven distractors was randomly determined from the remaining 63 locations without restriction for each trial. To make the search task more challenging we randomly rotated the L-shaped distractors by 0°, 90°, 180°, or 270°. For the eight Repeated condition trials we created eight configurations of the target and the seven L distractors. The target location within each quadrant was randomly determined for each display, as well as the orientation of the target (i.e., \( \square \) or \( \mathbf{1} \)). The target position was balanced across the four quadrants of the screen. The location of the seven distractors (also rotated on their center as described for the new context condition) was randomly determined from the remaining 63 locations without restriction for these eight displays. These eight repeated configurations then were randomly mixed with the eight new random condition trials to determine the presentation order of each block of the session. A peck of the target resulted in the appearance of the choice keys. A correct response to the target initiated the reward sequence, and errors prompted a correction trial as described above. Each daily session consisted of between 12-20 such blocks. We recorded the latency (ms) of the first peck to the target as a dependent measure. The median RT was calculated for each bird and then averaged across birds. These mean RT measures were log transformed prior to performing statistical analysis. We conducted a repeated measures Analysis of Variance (ANOVA) using Condition (Novel, Repeated) and Block as within-subject variables and RT (ms) as a dependent measure. Confidence intervals are reported after the partial eta squared value \( (\eta^2_p) \) for each effect where appropriate.

Five birds encountered 800 trials during Test 1, one bird 544 trials, and one bird 256 trials. The performance of these later two birds was comparable to the other five birds. These two birds were shifted into Test 2 more rapidly as they served as initial pilots through the Test 1 procedure.

Test 2: One goal of conducting Test 2 was to examine if whether learning about a context - if it occurred in Test 1- is continuous or if it stops once regularities about the background have been acquired. Five of the birds continued into Test 2, as some of the birds in the first study were used in a neural manipulation as part of

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Figure 1. An example of the target, distractors, and choice buttons used during testing.
another study and did not continue forward. Note, that our choice of which birds continued in the following tests was not based on their performance in earlier tests, but determined by randomization. As in Test 1 the birds encountered the same eight Repeated trials as in Test 1 and eight Novel trials during each block. We also presented the pigeons with a second set of eight repeated configurations (Repeated 2) of the seven distractors relative to the target (as described above) that they had not encountered previously. Thus, during each block of testing the pigeons encountered eight Repeated trials (the same eight configurations from Test 1), eight Novel trials, and eight Repeated 2 configurations for a total of 24 trials in each block. If the birds continued to learn about background regularities with new configurations, then the pigeons might be expected to show a RT advantage to the second population of repeated configurations. Likewise, since the pigeons may have already learned about contextual regularities in Test 1, learning about new contexts might be expected to be faster than in Test 1.

Each daily session consisted of approximately 10 blocks for a total of 240 trials. Four of the birds completed 432 trials, whereas one bird completed 96 trials (the bird that completed 256 trials in Test 1). We conducted a within subjects ANOVA using Block and Condition (Novel, Repeated, Repeated 2) as variables, and RT (ms) as a dependent measure to examine for differences between the conditions.

Results and Discussion
The birds became faster in finding the target during the course of Test 1, resulting in a significant effect of block, $F(23, 92) = 9.324, p < 0.0001, n_p^2 = 0.700, 117.038, 283.582$ (Figure 2, top). The ANOVA also revealed a significant interaction between condition and block, $F(23, 92) = 1.827, p = 0.023, n_p^2 = 0.314, 0.251, 49.819$. The effect of the interaction appears to play out in two ways. First, response times for the Random and Repeated conditions appeared comparable during the first two blocks but diverged after Block 3 until Block 16. We conducted simple planned comparisons examining response times for the first three blocks. As anticipated, there was no reliable difference in response times between the two groups during Block 1, $t(6) = 1.170, p = 0.286, -0.059, 0.168$, and Block 2, $t(6) = 0.413, p = 0.694, -0.303, 0.426$, but the responses times for the two groups started to diverge during Block 3, $t(6) = 0.413, p = 0.694, -0.303, 0.426$. The second phase of the interaction was that by Block 16 the response times for the two groups converged and appeared similar to each other, as at the start of testing. Indeed, comparisons indicated that there was no difference between the groups starting at Block 16, $F(1,4) = 0.435, p = 0.682, n_p^2 = 0.643, 0.000, 6.851$. The ANOVA we conducted also revealed a marginally significant main effect of condition, $F(1,4) = 7.189, p = 0.055, n_p^2 = 0.643, 0.000, 27.626$, with the Repeated group showing faster responses times compared to the Random condition when compared across all blocks. Notably, the five birds that were designated to continue onto Test 2 (see below) showed a similar pattern of performance to the entire set of birds used in Test 1 (Figure 2, middle).

When the birds were first introduced to the search task they learned to search the display and locate the target and their responses times across both conditions improved, as expected. The birds were also faster in finding the target during the Repeated compared to the Novel condition for the large balance of Test 1. Thus, the pigeons demonstrated the contextual cuing effect and were able to use the repeated arrangement of distractors in the displays (Repeated condition) to more effectively locate a target than when such contextual information was not present (Novel condition). The pigeons acquired knowledge about the target-background relationships rapidly, as the difference in RTs between the two conditions was apparent during the first block of trials. The faster RTs for the Repeated condition were observed both during the learning phase and continued when performance had stabilized. The fact that the pigeons had some intermediate training prior to Test 1 may have failed a familiarity of the procedures and a rapid acquisition of the cuing effect. Interestingly, the RT advantage for the Repeated compared to Novel condition disappeared during the final blocks of Test 1. We are uncertain why this occurred, but discuss a few possibilities in more detail in the discussion below.

During Test 2 we presented pigeons with the same repeated displays as in Test 1 (Repeated-1), a new set of repeated displays (Repeated-2) and novel stimulus arrangements (Novel). As can be seen in Figure 2 (bottom), the RTs for the Novel displays were consistently longer across all blocks compared to either the Repeated-1 or
Figure 2. The mean Log Response Time (RT) in milliseconds to find the target across blocks of trials during the two tests. During Test 1 (top) the pigeons were presented with Novel and Repeated displays. The middle panel shows the results from the subset of the five birds that continued onto Test 2. Note that the performance of this subset of birds is comparable to the pattern for all seven birds shown in the top panel. During Test 2 (bottom) we presented the pigeons with Novel, Repeated, and Repeated-2 displays. Note, the HE column on the far left indicates the scores for the first half Epoch of Epoch 1. Every other Epoch contains a full 48 trials per Epoch. Please see the text for a description of the stimulus conditions for each test. Error bars show SEMs.
Repeated-2 display conditions. Correspondingly, ANOVA revealed a significant main effect of condition, $F(2,6) = 15.452, p = 0.004, \eta^2_p = 0.837, 2.654, 82.395$. Least significant difference (LSD) comparisons indicated that RTs for the Novel condition were significantly longer compared to the Repeated-1 condition ($p = 0.008$) and marginally significant when compared to the Repeated-2 condition ($p = 0.072$). We subsequently performed comparisons of response times for the Novel and the Repeated-2 conditions during the first four blocks. The response times for the Novel and Repeated-2 conditions were comparable during Block1, $t(4)=0.593, p = 0.585, -410.629, 633.657, \text{Block2, } t(3)=0.794, p = 0.485, -349.605, 582.089, \text{Block3, } t(3)=0.786, p = 0.489, -360.354, 596.843, \text{but diverged at Block4, } t(3)=-8.557, p = 0.003, -169.115, -77.421$. Neither the effect of Block, $F(8,24) = 1.051, p = 0.428, \eta^2_p = 0.259, 0.000, 16.051$ nor the interaction between Condition and Block, $F(16,48) = 0.359, p = 0.986, \eta^2_p = 0.107, 0.000, 0.000$ reached significance.

The pigeons were faster to respond to the repeated-1 displays compared to the novel displays indicating that they retained their learning about the relationship between the distractors and the goal during Test 2. The pigeons also learned the new Repeated-2 displays, show divergence with the novel condition starting around block 4, again demonstrating an effect of contextual cuing. Finally, the RTs for the Novel condition were comparable to those observed at the end of Test 1, but the RTs for Repeated conditions were faster.

In the current study we used procedures comparable to those of Chun and Jiang (1999) in two separate tests to examine if pigeons show contextual cuing. We used Chun and Jian’s procedures so that we could isolate the spatial configuration information potentially driving contextual cuing, rather than other factors like conspicuousness or scene categorization that might occur when using natural scenes. The results from our tests suggest that pigeons demonstrated contextual cuing and that the effect was due to the features of the distractors rather than other display features. Like people, pigeons were faster to respond to displays in which the position of the distractors predicted the target location, compared to displays in which there was no reliable spatial relationship between the two sets of items (e.g., Chun & Jiang, 1998). Pigeons, like people, also showed the contextual cuing effect early during testing (Chun, 2000; Chun & Jiang, 1998). Similar results have also been reported by Brooks, Rasmussen, Hollingworth, & Wasserman (2008) with pigeons and by Brooks, Dai, & Sheinberg (2011) with monkeys using procedures similar to that of Chun and Jiang. Knowledge about the configurations was retained over time across the tests by the pigeons in the current study and the addition of new configurations did not appear to impair performance based on earlier contextual learning (Test 2), effects which have also been observed for people.

The current findings are also consistent with other work documenting contextual cuing like effects in non-human animals. As mentioned previously, Goujon & Fagot (2013) have explored contextual cuing in baboons using search task procedures comparable to Chun & Jiang (1999). Baboons, like pigeons in the current study and people in past work, were faster to respond to the target (T) when the background was predictive compared to when it was not predictive.

In addition the current results support other work on contextual cue with pigeons that have used simple images or scenes. Wasserman, Teng, & Castro (2013) have explored contextual cuing like effects using somewhat different procedures. These researchers had pigeons find a target (simple image) that was presented on a background that had a unique solid color or real-world photograph. The birds were faster to respond to the target when it was predicted by the background (context) than when the target location was determined randomly. Additionally, Wasserman, Teng & Brooks (2014) using similar procedures recently found that contextual cuing could occur with up to an 8 $s$ delay between the appearance of the scene and the target onset. These authors also found that more global features of the predictive scenes tended to control search rather than more local features of the context.

As mentioned previously, the contextual cuing effect disappeared at the end of Test 1 before reappearing again for the exact same displays during Test 2. This lapse in spatial cuing is somewhat surprising given that humans retain information about spatial context in memory for at least one week (e.g., Chun & Jiang, 2003). The lapse in cuing is unlikely to be due to a floor effect, since when cuing reappears in Experiment 2 response
times are faster than those observed when contextual cuing was present during Test 1. Also, it is unlikely that the birds failed to retain information about spatial context in memory since the effect returns quite prominently at the beginning of Test 2. Why then did the pigeons fail to demonstrate contextual cuing at the end of Test 1?

There are several possible explanations. One possibility is that while the birds retained knowledge about the spatial context of the repeated displays, they had encountered a change in motivation toward the end of Experiment 1 that prevented them from expressing such knowledge. Another related possibility is that the pigeons had shifted their attention primarily to the Novel displays after having well learned the Repeated displays by the end of Test 1. Extended repeated testing with a small number of displays may have exacerbated the shift in attention. Ultimately, we have strong evidence that the spatial contexts were robustly learned and retained across sessions, but we remain puzzled about why the expression of this learning temporarily subsided at the end of Test 1 and hope to address this in future research.

One important facet of contextual cuing in humans is that the process is implicit. Participants do not report knowledge of the spatially correlated displays even though they encounter them repeatedly in a session. Awareness in non-human animals is controversial and difficult to assess (see Shettelworth, 2010 for a review). Thus, while the performance of people and non-human animals appears quite comparable based on their performance, it is unclear whether non-human animals have knowledge of the repeated displays or whether the effects are operating in the same way.

Past work has revealed that participants show a decrement in learning about background relationships when natural scenes serving as the background are inverted compared to when the images are upright (Brockmole & Henderson, 2006). This suggest that semantic encoding of the background information may be important for people, particularly for natural scenes. Future work also might compare whether pigeons and other non-human animals might also show a similar effect, or whether performance on a comparable contextual cuing tasks that differs in only the nature of the background information (e.g., scenes vs. artificial stimuli) show different rates of learning.

Finally, the fact that both non-human primates and now pigeons have been shown to demonstrate the contextual cuing during a complex visual search task speaks to the ubiquitous nature of the effect. Using background information increases efficiency during the search processes that many animals engage in for survival. Given that pigeons have very different visual systems than primates, different animals are likely using different neural architectures to solve the same search process. Future work exploring these possibilities would be of considerable interest.

References


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