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Eyespot configuration and predator approach direction affect the antipredator efficacy of eyespots

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Many prey species possess eyespots: paired markings that often consist of two or more concentric circles. Predators are wary of such prey because eyespots are conspicuous and/or mistaken for vertebrate eyes. Here we used naïve domestic chicks as predators of artificial moth-like prey to test the hypothesis that both eyespots configuration and predator approach direction affect the antipredator efficacy of eyespots. We found that when chicks approached prey straight on, eyespots configuration did not influence attack latency. Chicks that approached from either the left or the right, were slower to attack prey in which the central circle of the eyespot was centrally placed or shifted in the direction of the chick's approach, compared to prey in which the central circle had been shifted away from the direction of approach. These findings suggest that eyespots composed of concentric circles may protect prey against predators approaching from a wider range of directions than evespots composed of eccentric circles. They are also consistent with the idea that eyespots are mistaken for eyes, and are perceived to pose a lesser risk when their "gaze" is averted from the approaching predator.

KEYWORDS

antipredator defense, eye gaze, mimicry, predator-prey, protective coloration, wariness

Introduction

Many prey species possess markings that subjectively appear to resemble vertebrate eyes (Janzen et al., 2010). These are known as eyespots and are conspicuous circular or quasi-circular markings (Mukherjee and Kodandaramaiah, 2015). Whilst in some species eyespots appear to play a role in mate choice (Robertson and Monteiro, 2005; Huq et al., 2019), one of their key functions is to protect prey from predators (Blest, 1957; Stevens, 2005; Kjernsmo and Merilaita, 2013; Prudic et al., 2015). They are among the most studied wing color pattern elements in lepidoptera (Stevens, 2005; Kodandaramaiah, 2011), and their antipredator benefits have been predicted or established in various taxa (Stevens, 2005) including Lepidoptera (Lyytinen et al., 2004; Vallin et al., 2005, 2011; Wiklund, 2005; Olofsson et al., 2010, 2012; Merilaita et al., 2011; Ho et al., 2016; Halali et al., 2019; Chan et al., 2021), Orthoptera (Steiner, 1981; Castner, 1995), Coleoptera (Buschman, 1988), and fishes (Altbäcker and Csányi, 1990; Meadows, 1993; Gagliano, 2008; Kjernsmo and Merilaita, 2013; Kjernsmo et al., 2016; Hemingson et al., 2021).

Eyespots reduce the chance of being eaten in a number of ways. Peripherally positioned eyespots, such as those of the squinting bush brown (Bicyclus anynana) and the woodland brown (Lopinga achine) butterflies, can deflect attacks to nonvital body parts (Olofsson et al., 2010; Skelhorn et al., 2014). The eyespots are thought to draw predators' attention, causing them to direct their attacks toward dispensable areas (e.g., wing margins), enhancing the chances that prey will survive the attack (Blest, 1957; Kjernsmo and Merilaita, 2013; Prudic et al., 2015). Larger eyespots such as those of the peacock butterfly Aglais io elicit an aversive response in predators (Vallin et al., 2005, 2010, 2011; Kodandaramaiah et al., 2009). The question of why predators should find eyespots aversive has caused considerable debate (Stevens, 2005; Stevens and Ruxton, 2014; Mukherjee and Kodandaramaiah, 2015). However, there is now evidence for two non-exclusive explanations. Eyespots appear to exploit predators' innate or learned aversions to conspicuously colored prey (Stevens et al., 2008, 2009), and mimic the eyes of the predators' own predators (Blut et al., 2012; De Bona et al., 2015; Kjernsmo and Merilaita, 2017).

The specific design of eyespots has received much less attention (though see: Stevens et al., 2008; Blut et al., 2012; Hossie et al., 2015; Kjernsmo and Merilaita, 2017 and related papers on responses of birds and reptiles to eye-like stimuli, e.g., Gagliardi et al., 1976; Hennig, 1977). There are lots of ways to be conspicuous and several ways to resemble eyes. However, many (although by no means all) eyespots consist of multiple concentric rings of contrasting colors (Skelhorn et al., 2016). This may well be due to the radial development of pigmented cells from each eyespot's center (Monteiro, 2015). Alternatively, concentric circles with either a bright or dark central region, and the opposite value in its surround may be particularly effective at stimulating vertebrate visual systems and associated brain regions (e.g., the optic tectum; Mey and Thanos, 2000). This common eyespot configuration may be highly conspicuous to the circular receptive fields of vertebrates (as suggested by Stevens, 2005; Stevens et al., 2007, 2008), and conspicuous signals can be especially aversive to birds (Stevens et al., 2008; Halpin et al., 2020). Additionally, concentric circles subjectively resemble forward-facing eyes. Several species of birds and reptiles are sensitive to subtle eye gaze cues, and respond more aversively when gaze is directed toward them compared to when it is averted (Hennig, 1977; Carter et al., 2008; von Bayern and Emery, 2009; Clucas et al., 2013; Garland et al., 2014; Goumas et al., 2019, 2020). Consequently, if eyespots are perceived as predatory eyes, then we might expect forward-facing eyespots to be particularly effective because they appear to gaze at predators approaching from a wide range of different directions. Conversely, the efficacy of eyespots made up of eccentric circles should be heavily influenced by predator approach direction, being most effective when eyespots appear to "gaze" toward the approaching predator and least effective when they "gaze" away from it. Here we test these ideas using naïve domestic chicks *Gallus gallus domesticus* as predators of artificial moth-like prey.

Materials and methods

Subjects and housing

A total of 126 domestic chicks (Gallus gallus domesticus) of the "Hubbard Brown" strain served as predators. Chicks were acquired in two batches (n = 63 per batch) from a commercial hatchery on the day they hatched. They were housed in cages measuring 120 cm \times 50 cm \times 50 cm, and were subject to a 14L: 10D cycle using uncovered fluorescent lights (OSRAM L360Q/965 Biolux). Cage temperatures were maintained at 25-28°C using a room heater. Water was provided ad lib., as were chick starter crumbs except during training and experimenting when food restriction was necessary. When access to food was restricted, chicks had access to water. Chicks received mealworms (Tenebrio molitor) twice a day in their home cages. All subjects were marked with non-toxic Sharpie marker pens which did not have any adverse effects on their behavior. Weights were monitored for welfare purposes throughout the experiment, and all chicks gained weight as the experiment progressed. No specific measures were taken to obscure the eyes of the staff performing routine husbandry. However, during all training and experimental trials, chicks were viewed from above and behind the arena in an attempt to ensure that chicks could not see the experimenter's head.

Preparation of artificial prey

Artificial moths were created by pinning paper triangles (base 44 mm, height 37 mm) over mealworms (*Tenebrio molitor*) to completely obscure them from view. We used Adobe Illustrator 14 (13) to create three types of prey. All three types possessed a pair of eyespots that consisted of a larger white circle containing a smaller gray circle. In left-shifted prey, the smaller circle was shifted to the moth's left; in central prey, the smaller circle was in the center of the larger circle; and in right-shifted prey, the smaller circle was shifted to the moth's right. We also created three training stimuli with the same conspicuousness as

the test stimuli (measured as luminance), but without eyespots (see Figure 1).

All prey consisted of a triangle 100 pt in width and 86.6 pt in height and filled with mid gray (RGB: 164, 164, 164; C: 38 M: 31 Y: 31 K:0). To produce test stimuli that possessed eyespots, we added two white circles 31.4 pt in diameter and two gray circles 16.0 pt in diameter. The centers of the white circles were always positioned 30.5 and 69.5 pt to the right of, and 18.4 pt above, the left apex of the triangle (thus ensuring that each eyespot was equidistant from the triangle's midline). However, the position of the smaller circles differed among the three test stimuli: to produce central prey, we positioned the centers of the gray circles in the same positions as the centers of the white circles; to produce left- and right-shifted eyespots, we shifted the smaller circles 6.6 pt to the prey's left or right, respectively. We then created three training stimuli with similar luminance to the test stimuli, but without eyespots. These had "mottled" patterns, and were created by drawing 10 irregular white shapes of similar area and positioning them so that they were spread over the entire triangle. The combined area of the white shapes matched the area of two white circles minus that of the two gray circles (that is, the white area on a mottled stimulus was the same as that on an experimental stimulus). This was achieved by adjusting the shapes until the luminance of the mottled stimulus matched the luminance of the experimental stimuli. Three training stimuli were produced by rotating and repositioning the shapes (an example of one of the training stimuli and all of the test stimuli can be seen in Figure 1).

Training

On day one post-hatch, chicks were allowed to acclimate to the laboratory for 1 h, before being randomly assigned to one of three experiments (N = 42 per experiment). Chicks in each of the three experiments were then randomly assigned to one of three experimental groups (N = 14 per experimental group). All birds were trained to attack training stimuli pinned to the back wall of a runway. The protocol was identical in all three experiments, but each experiment used a different runway (Figure 2). All runways were made of reinforced cardboard covered in black paper. The distance between the start point (at which chicks were introduced to the runway) and the center of the rear wall was 50 cm, and the distance between the right and the left wall was 15 cm. However, the angle between the side walls and rear wall differed among experiments. From the inside of the runway, the angle between the left-hand wall (as viewed when facing the rear wall) and the rear wall was 135° in Experiment 1, 90° in Experiment 2, and 45° in Experiment 3. This arrangement meant that chicks were only able to approach prey items pinned to the rear wall from the prey's left, straight on, and from the prey's right, respectively.

On day one post-hatch, chicks were trained to forage alone in the runways in six training trials at regular intervals throughout the day. In all trials, chick crumbs were scattered over the white laminated floor of the experimental runway, and chicks were then placed in the runway for 2 min. In the first two trials, chicks were placed in runways in groups of three; in the following two trials, in groups of two; and in the final two trials chicks were placed in runways individually. Chicks had restricted access to food for 30 min prior to each of the final three trials. By the end of trial six, all chicks were eating chick crumbs in the runways.

On day 2 post-hatch chicks were trained to eat mealworms pinned to the rear wall of the experimental runways. Each chick received four trials at regular intervals throughout the day. In each trial, a single mealworm was pinned to the center of the rear wall 10 cm from the runway floor. Chicks had restricted access to food for 30 min and were then placed in the runway and left there until they attacked the mealworm. This process was repeated on day 3 post-hatch, but in these trials a training stimulus (mottled paper triangle, Figure 1) was pinned over the mealworm partly obscuring it from view. Finally, on day 4 post-hatch, each chick received a further four trials in which the training stimulus completely obscured the mealworm from view: the base of the stimulus was always positioned in the center of the back wall, 10 cm from the runway floor. We used three training stimuli (as described above) during trials on days 3 and 4 to ensure that our findings were not restricted to a specific training regimen. Each regimen was received by two birds from each of the three experimental groups in each of the three experiments. Birds received either stimulus one, stimulus two, stimulus three, stimuli one and two, stimuli one and three, stimuli two and three, or all three stimuli. All chicks attacked the mealworm within 10 min in every training trial, and by the end of the final trial, all chicks attacked the artificial prey item within 2 s of entering the arena.

Test trial

On day 5 post-hatch, chicks were randomly assigned to one of three experimental groups (n = 14 per group). All chicks received a single test trial. Each chick had restricted access to food for 30 min before being placed into the same runway used in training. A single prey item was pinned to the center of the rear wall 10 cm above the floor. The position of the central circles of the eyespot (analogous to the iris of a real eye) differed among experimental groups. In each experiment, one group of chicks encountered prey with circles shifted to the prey's left, one encountered prey with centered circles, and one encountered prey with circles shifted to the prey's right (see **Figure 1**). To human observers positioned directly in front of the prey, these eyespots appeared to "gaze" to the



FIGURE 1

Images of the artificial moth stimuli. From left to right: training prey, left-shifted prey (the central circle is shifted to the moth's left), central-circle prey, and right-shifted prey (the central circle is shifted to the moth's right).



prey's left, straight on, and right, respectively. Chicks remained in the runway until they attacked the prey or until 10 min had elapsed (whichever came first). We recorded the latency to attack the prey, and the presence/absence of approachretreat behavior: repeatedly approaching and retreating from the prey (Skelhorn et al., 2014, 2015). Only five chicks failed to attack the prey in 10 min (two in experiment 1, two in experiment 2, and one in experiment 3), and these were included in the analyses by awarding them an attack latency of 601 s. There were no qualitative differences between these analyses and analyses in which these chicks were excluded (see **Supplementary material**). Chicks were trained and tested in the same random order throughout the experiments.

All procedures were in accordance with UK Home Office regulations and the Association for the Study of Animal Behavior's Guidelines for the Treatment of Animals in Research and Teaching. The University Psychology Ethics Committee approved the experiment (proposal 2013/319). Chicks were donated to a free-range smallholding at the end of the study.

Data analysis

We square root transformed the latency data for positive skew in the raw data, and analyzed the effect of eyespot design on the latency to approach with a general linear model with a Gaussian distribution and identity link function. We calculated standardized effect sizes (Cohen's d) using Package 'rstatix' (Kassambara, 2021). We analyzed the approach retreat behavior with Fisher's exact tests. We calculated standardized effect sizes (Cohen's *d*) using the package rstatix. The GLM and associated *post-hoc* tests were conducted using R version 4.1.2. The Fisher's tests were conducted using GraphPad¹.

Results

By the end of training, all chicks readily attacked the training prey within 2 s. In the test trial, however, all chicks took longer than 2 s to attack the test stimulus, indicating that the eyespots had a deterrent effect. In Experiment 1, when chicks approached from the prey's left, there was a significant difference in the average latency to approach the prey (main effect: $F_{2, 39} = 14.06$, p < 0.0001; see Figure 3). The average attack latency did not differ between the leftshifted group and the central group (estimate = -0.09 ± 1.74 , df = 39, t = -0.049, P = 0.9987). However, chicks in both these groups took longer to attack prey than chicks in the right-shifted group (left: estimate = 7.97 \pm 1.74, df = 39, t = -4.567, P = 0.0001, Cohen's d = 1.16; central: estimate = 8.06 ± 1.74 , df = 39, t = -4.617, P = 0.0001, Cohen's d = 1.52). In Experiment 2, when chicks approached prey straight on, there was no significant difference in the average attack latency among our experimental groups (main effect: F_{2} , $_{39} = 0.009$, P = 0.992; see Figure 3). Finally, in Experiment 3, when chicks approached from the prey's right, there was a significant difference in the average latency to approach the three prey types (main effect: F_{2} , $_{39} = 15.91$, p < 0.0001; see Figure 3). The average attack latency did not differ between right-shifted group and the central group (estimate = -1.24 ± 1.79 , t = -0.696, p = 0.7669). However, chicks in both these groups took longer to attack prey than chicks in the left-shifted group (right: estimate = -9.28 ± 1.79 , df = 39, t = -5.196, P < 0.0001, Cohen's d = -1.88; central: estimate = -8.03 ± 1.79 , df = 39, t = 4.50, P = 0.0002, Cohen's d = -1.44).

Similar trends were seen in approach retreat behavior: an established measure of wariness, where chicks repeatedly



Time taken to attack the test stimuli for birds in each of the three experiments. light gray bars represent groups given left-shifted prey; mid-gray bars groups given prey with central circles; and dark gray bars groups given right-shifted prey (N = 14 in each group). Boxplots show the median, interquartile range, and the whiskers represent the largest and smallest value within 1.5 times the 25th and 75th percentile. Outliers are represented by small black dots.

approach and retreat from prey before making contact with it (Skelhorn et al., 2014, 2015). In Experiment 1, when chicks approached prey from the prey's left, there was no significant difference between the left-shifted group and the central group in the number of chicks that demonstrated this behavior (P = 0.4815; see Figure 4). However, in both these groups more chicks demonstrated this behavior than in the right-shifted group (right vs. left: $P = \langle 0.0001;$ right vs. straight: P = 0.0004). In Experiment 2, when chicks approached prey straight on, there was no significant difference in the number of chicks demonstrating this behavior among our experimental groups (P > 0.1 in all cases; see Figure 4). Finally, in Experiment 3, when chicks approached from the prey's right, the rightshifted group and the central group did not differ in the number of chicks that demonstrated this behavior (Fisher's exact test, P = 0.4815; see Figure 4). However, more chicks in the right-shifted group demonstrated this behavior than in the left-shifted group (left vs. right: P = 0.0006), and there was a similar trend for the central group that was not significant after correcting for multiple comparison (left vs. central: P = 0.0183).

Discussion

Our study shows that the effectiveness of artificial eyespots is dependent on their configuration and the approach direction of the predator. When chicks approached prey straight on, eyespot configuration had no significant effect on our measures of chick behavior. However, when chicks approached prey from either the left or right, chicks were slower to attack prey with eyespots shifted in the direction of their approach

¹ https://www.graphpad.com/quickcalcs/contingency1/



compared to when they encountered prey with the central circle of the eyespot shifted away from the direction of their approach. Chicks were also more likely to demonstrate aversive responses (approach-retreat behavior), when they encountered prey with the central circle of the eyespot centrally placed or shifted in the direction of the chick's approach. This suggests that eyespots consisting of concentric circles could offer greater protection than eyespots with left- or right-shifted central circles in situations where individual predators approach from a range of different directions, as is likely the case for moths and butterflies (Stevens, 2005), but not for caterpillars (Hossie et al., 2015).

Although we did not directly test whether chicks mistook our artificial eyespots for eyes, our findings are consistent with this idea. Chicks may have found prey with the central circles shifted away from the direction of approach less aversive than other prey because they were perceived to have eyes that gazed away from, rather than toward, the approaching chick. This would reduce the risk of the chick being detected by the predator, competitor, or prey that the eyes were perceived to belong to. This explanation is in line with previous work demonstrating that birds respond more aversively to eyes that gaze toward rather than away from them (Carter et al., 2008; von Bayern and Emery, 2009; Clucas et al., 2013; Garland et al., 2014; Goumas et al., 2019, 2020). We didn't, however, find a difference in protection between central eyespots and those shifted in the predator approach direction. This could be because eyespots with concentric circles appear to gaze directly at observers approaching from the side. In humans this perceptual phenomenon is known as the "Mona Lisa" effect (Bruce and Young, 1998), because the forward-facing eyes of that portrait appear to follow viewers around the room. Alternatively, the central circle of our eyespots may have been shifted too far toward the direction of approach, causing the prey to "look" slightly past the birds rather than directly at them.

It is more difficult to explain our findings in terms of prey conspicuousness as our prey were designed to be equally conspicuous by all commonly used measures. The number of gray and white pixels, the size and shape of boundaries between light and dark areas, and minimum, maximum and mean luminance were consistent across prey types. It is possible that the central gray circle appeared to merge with the outer gray area of the prey when it was shifted away from the direction of approach, thus reducing perceived internal contrast. However, human observers perceived a distinct inner circle when viewing the prey from the start of the experimental runway. In addition, our findings provide no support for the idea that concentric circles are particularly aversive because they are more effective at stimulating vertebrate visual systems (Stevens et al., 2007, 2008). Predators approaching prey straight on were equally reluctant to attack prey with concentric and eccentric circles.

It could be argued that we should have included control groups that were given prey without eyespots. We decided against this because our aim was to establish how the relative aversiveness of the stimuli changed with viewing angle (i.e., across experiments), not how aversive eyespots were compared to a specific control stimulus. We also chose to use artificial stimuli in a laboratory environment as this allowed us to manipulate prey appearance and predator approach direction in a controlled manner. The use of paper prey with achromatic eyespots has been fundamental in establishing the antipredator benefits of eyespots (Blest, 1957; Scaife, 1976; Coss, 1979), yet such prey may not capture the nuances of real prey (De Bona et al., 2015). Whilst future work comparing how predators' responses to artificial and live prey would be beneficial, we predict that predators are likely to have a broad view of what is considered a conspicuous/eye-like stimulus because the cost of failing to respond to such stimuli is potentially very high. It could also be argued that the question we address is irrelevant because eyespots design is not the result of predator-driven selection, but rather developmental constraints (Monteiro, 2015). This, however, misses two key points. Conspicuous eyespots are only likely to evolve if they are beneficial to prey, and we cannot fully understand the benefit of eyespots without establishing how, and under what circumstances, they work. Moreover, not all eyespots consist of concentric circles, meaning that in some species developmental constraints are either absent or have been overcome.

Irrespective of these limitations, our results clearly demonstrate that eyespot design and predator approach direction can influence the antipredator benefits of eyespots. Consequently, the microhabitat in which prey position themselves and the feeding ecology of predators could affect eyespot efficacy (and perhaps evolution), and these factors should be carefully considered when designing future experiments.

Data availability statement

The original contributions presented in this study are included in the article/Supplementary material, further inquiries can be directed to the corresponding author.

Ethics statement

This animal study was reviewed and approved by the University of Exeter Psychology Ethics Committee (proposal 2013/319).

Author contributions

HR: conceptualization, methodology, formal analysis, data curation, writing—original draft, and visualization. JS: conceptualization, methodology, investigation, resources, data curation, writing—original draft, and funding acquisition. Both authors contributed to the article and approved the submitted version.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/ fevo.2022.951967/full#supplementary-material

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