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34

35 **Competing interests**

36 We declare we have no competing interests.

37

38 **Author contribution**

39 AM: Conceptualization, Data curation, Formal analysis, Funding acquisition, Investigation,
40 Methodology, Software, Project administration, Supervision, Visualization, Writing –
41 original draft, Writing – review & editing

42 ML: Data curation, Funding acquisition, Investigation, Methodology, Writing – review &
43 editing

44 PP: Data curation, Investigation, Writing – review & editing

45 YY: Data curation, Investigation, Writing – review & editing

46 MS: Conceptualization, Funding acquisition, Visualization, Writing – review & editing

47 SN: Conceptualization, Funding acquisition, Methodology, Software, Supervision, Project
48 administration, Writing – review & editing

49

50 All gave final approval for publication and agreed to be held accountable for the work
51 performed therein.

52

53 **Data Accessibility**

54 Raw data, analysis script and supplementary materials are available at [https://ayumi-
55 495.github.io/eyespot/](https://ayumi-495.github.io/eyespot/). Once the paper is published, these will all be uploaded to Zenodo.

56 **Abstract**

57 Eyespot patterns have evolved in many prey species. These patterns were traditionally
58 explained by the eye mimicry hypothesis, which proposes that eyespots resembling vertebrate
59 eyes function as predator avoidance. However, it is possible that eyespots are not the mimicry
60 of eyes: according to the conspicuousness hypothesis, eyespots are just one form of vivid
61 aposematic signals where only conspicuousness matters. To test these hypotheses and explore
62 factors influencing predators' responses, we conducted a meta-analysis with 33 empirical
63 papers focusing on bird responses to lepidopterans having conspicuous patterns (eyesspots and
64 non-eyesspots). Supporting the latter hypothesis, the results showed no clear difference in
65 predator avoidance efficacy between eyespots and non-eyesspots. When comparing geometric
66 pattern characteristics, bigger pattern sizes and smaller numbers of patterns were more
67 effective in preventing avian predation. This finding indicates that paired concentric patterns
68 have weaker deterring effects than single ones. Taken together, our study supports the
69 conspicuousness hypothesis more than the eye mimicry hypothesis. Due to the number and
70 species coverage of published studies so far, the generalisability of our conclusion may be
71 limited. The findings highlight that pattern conspicuousness is key to eliciting avian
72 avoidance responses, shedding a different light on this classic example of signal evolution.

73

74 **Keywords**

75 Aves, butterfly, caterpillar, interspecific communication, predator-prey interaction, warning
76 signal

77 **Background**

78 Naturalists have long pondered the evolution and function of the many signals and cues
79 animals use to communicate [1–9]. Visual signals, such as vibrant colours and contrasting
80 patterns, have attracted more interest from researchers than other signals, likely because our
81 species is visually oriented [1, 10, 11]. Eyespot patterns, characterised by concentric rings of
82 different colours with a light outer ring and a dark centre [12], are well-known patterns
83 believed to reduce predation. Although eyespots have been researched for a long time [12–
84 15], researchers continue to debate why eyespots might deter predation.

85 Three hypotheses have been proposed to explain why eyespot patterns can contribute
86 to prey survival (reviewed in [12, 14, 15]; Fig. 1). First, the eye mimicry hypothesis suggests
87 that eyespots play a role in deterring predators from attacking prey and reducing predation
88 risks by mimicking the eyes of vertebrates [16–18]. This hypothesis predicts that if the
89 pattern has specific characteristics (e.g., eye-like shape) and is presented as a pair, predation
90 avoidance will increase, assuming eyespots imitate potential predators. Second, the
91 conspicuousness hypothesis posits that eyespots are simply conspicuous patterns that prevent
92 attacks due to negative predator responses caused by sensory bias, neophobia, or sensory
93 overload [12, 14]. The hypothesis states that the eye-like shape and patterns arranged in pairs
94 have nothing to do with predator deterrence. Eyespots can act as an aposematic signal for
95 potential predators. For example, if the size of the pattern (one of the measures of
96 conspicuousness) increases, the avoidance effect will also increase. Third, the deflection
97 hypothesis suggests that predator attacks should be directed towards eyespots to avoid
98 damage to vital body parts [19–23]. The eye mimicry and conspicuousness hypotheses are
99 usually applied to explain large eyespots, while the deflection hypothesis is used to interpret
100 the function of small ones [12, 14, 15]. Although there seems to be little disagreement in the
101 third hypothesis ([24–26], but see also [27]), why large eyespots can intimidate avian
102 predators has been controversial [12, 14]. This is because while the eye mimicry and
103 conspicuousness hypotheses are not mutually exclusive, the key mechanism that explains
104 why predators react negatively to eyespots is clearly different.

105 Lepidopterans, such as butterflies and moths, have been the leading models for testing
106 the eye mimicry and conspicuousness hypotheses. A typical empirical study has adult
107 individuals, caterpillars, or their models as prey, with birds as predators (reviewed in [12, 14,
108 15]). According to the eye mimicry hypothesis, avian predators perceive the eyespots as the
109 eyes of a potential enemy. For example, great tits (*Parus major*) showed more aversive
110 responses to animated butterflies with a pair of large eyespots than those without, and such
111 eyespots were more effective than modified, less mimetic, but equally contrasting patterns
112 [28]. Although several studies have supported the eye mimicry hypothesis [e.g., 16, 28, 29],
113 many conspicuous patterns other than eyespots, such as dots and stripes, likely deter attacks
114 from predators as well [30–33]. Some field experiments with artificial prey have supported
115 the conspicuousness hypothesis, demonstrating survival rates for both conspicuous (eyespot
116 and non-eyespots) pattern prey stimuli were higher than control prey stimuli [30, 31, 34].
117 Such discrepancies might have arisen from differences in experimental design between
118 studies, such as the size, number, and shape of the presented pattern stimuli or the bird
119 species used as subjects in the experiments [12, 35]. However, there has been no systematic
120 attempt to synthesise and compare earlier studies quantitatively.

121 Here, we conduct a systematic review with meta-analysis to synthesise empirical
122 evidence on the intimidating effects of eyespots and the factors that contribute to predator
123 avoidance responses towards them. To examine the two hypotheses above, we ask three
124 interrelated questions. First, we examine whether conspicuous patterns, namely eyespots and
125 non-eyespot patterns (i.e., conspicuous patterns other than eyespots), influence bird responses
126 or prey survival in a manner that increases the success of predator avoidance. Second, we test
127 whether pattern resemblance to eyes (eye-like shape) is the key to predator avoidance (which
128 differentiates the eye mimicry hypothesis from the conspicuousness hypothesis). Third, we
129 examine what factors promote bird response and increase prey survival, such as pattern size
130 and the number of patterns (i.e., eyespots and non-eyespots; Fig. 1).

131

132 **Materials and Methods**

133 We preregistered our methods and planned analyses before data extraction and analysis in
134 Open Science Framework (<https://osf.io/ymwvb>; [36]). We referenced and followed
135 PRISMA (Preferred Reporting Items for Systematic Reviews and Meta-Analyses; [37]) and
136 PRISMA-EcoEvo (Preferred Reporting Items for Systematic reviews and Meta-analyses in
137 Ecology and Evolutionary biology; [38]) for reporting this study (Table S1).

138

139 **Search protocols**

140 We used the PICO (Population, Intervention, Comparator, Outcome; Table 1) framework
141 [39] to specify the scope of our research questions and to inform our literature searching and
142 screening. We conducted a comprehensive literature search across multiple databases,
143 including Scopus, ISI Web of Science, Google Scholar (for non-English studies), and
144 Bielefeld Academic Search Engine (for unpublished theses; i.e., grey literature). We designed
145 the search strings (see Table S2) to identify studies that used experimental methods to
146 examine the effects of eyespot patterns on birds' predation behaviours. We did not set any
147 temporal restrictions on the database searches. Additionally, we conducted backward and
148 forward reference searches within the Scopus database using four key publications [12–15].
149 The strings were translated for searches in non-English languages, and search results were
150 assessed by reviewers with expertise in the respective languages: AM for Japanese, ML for
151 Polish and Russian, PP for Portuguese and Spanish, and YY for Simplified and Traditional
152 Chinese. We limited Google Scholar searches to the top 100 results in each language, sorted
153 by relevance. In cases of disagreement between the reviewers, discrepancies were discussed
154 and resolved to reach a consensus. The screening process and results are shown in the
155 PRISMA-like flowchart (Fig. 2a).

156

157 **Eligibility criteria**

158 We set specific criteria for including studies in our meta-analysis (according to our pre-
159 registered protocol). Initial screening, including titles, abstracts, and keyword assessment for
160 English-language bibliographic records, was conducted by AM and ML using Rayyan

161 (<https://www.rayyan.ai>; [40]) following predefined inclusion criteria. Subsequently, AM and
162 PP independently screened the full texts of studies that passed the initial screening. To be
163 eligible, a study had to conduct experiments and provide data on bird behavioural responses
164 or prey survival/attacked rates. We excluded studies solely involving non-avian predators,
165 such as fish, insects, mammals, or other species. However, studies that included a mix of
166 species from different taxonomic groups were allowed if the primary focus was on avian
167 predation. In our analysis, we only considered research that presented both conspicuous and
168 control (non-conspicuous) patterns as stimuli. We omitted studies using actual predator or
169 human eyes as stimuli since we focused on understanding how eyespot patterns in butterflies
170 and caterpillars, which are unlikely to resemble specific bird or vertebrate species eyes, affect
171 predation avoidance [41]. We also excluded studies that used bright and contrasting patterns
172 as control stimuli. Furthermore, we focused only on studies that used real or artificial
173 butterflies, moths, caterpillars, or a piece of paper as prey or presented stimuli. We also did
174 not consider research that only investigated avian physiological responses to conspicuous
175 patterns. In addition, we did not include studies that only assessed whether prey with eyespots
176 or conspicuous patterns were less likely to be attacked by birds, based on wing or body
177 damage alone, without including control stimuli.

178

179 **Data collection**

180 We extracted four types of information from each study. First, we collected citation
181 information, such as title, author name, and publication year. Second, we gathered the details
182 of the presented stimuli used in each experiment within studies: type of control pattern (plain
183 neutral-coloured or camouflaged), type of treatment pattern (eyespot or non-eyespot
184 patterns), pattern area (mm^2 : area per shape comprising the pattern), total pattern area (mm^2 :
185 when multiple patterns exist on the presented stimulus, it denotes the total area of all patterns;
186 for stimuli with single eyespot or distinct pattern, the value equals the pattern area), linear
187 size of the pattern (mm: e.g., maximum diameter or length of pattern), number of shapes in
188 pattern, total area of prey surface (mm^2 : e.g., butterfly wings and caterpillar bodies), prey
189 material type (i.e., whether a real butterfly or a complete imitation of a particular butterfly

190 was used as prey), and prey shape type (a further subdivision of the former). For non-eyespot
191 patterns, we also noted pattern shapes (e.g., circles, stripes, and triangles). In each study, bird
192 responses to control and treatment pattern stimuli and prey survival/attacked rates when these
193 patterns were present were reported. Bird responses contained a variety of measures,
194 including the number of attacks and escape behaviours, latency to attack, latency to approach,
195 and the proportion of birds attacking the presented stimuli. Henceforth, we refer to these
196 measures and responses as 'predator avoidance'. Third, we obtained data for calculating effect
197 sizes (e.g., mean, standard deviation or standard error, and sample size of control and
198 treatment group) from plots using WebPlotDigitizer 4.6.0
199 (<https://automeris.io/WebPlotDigitizer>), detailed tables, texts, or raw data. In survival
200 analysis plots, we extracted data at the point in time when the difference between the
201 'survival' or 'attacked' rates of the intervention and comparison groups was greatest as
202 outcomes. Study design (i.e., whether experiments were done independently or dependently
203 between the control and treatment group) was also recorded. Fourth, we gathered predator
204 and prey information, specifically, the study species (common English name and scientific
205 name) and predator diet type. In some cases, studies did not use a specific bird species as a
206 predator or a specific lepidopteran species as prey. We contacted authors when such
207 information was ambiguous or missing. When the paper did not report the pattern area and
208 diameter of the treatment stimulus or the presented stimulus surface area, AM calculated or
209 measured them from available images using ImageJ v.1.53i [42].

210 The dataset was originally divided into two parts. The first part involved the data from
211 presenting eyespot patterns to avian predators and directly observing their responses
212 (predator dataset). The sample size or unit of analysis in this part was based on the number of
213 individual avian predators. The second part involved the data from using real or artificial
214 abstract butterflies, moths, or caterpillars with eyespots or non-eyespot patterns as stimuli or
215 prey, and observing their survival/attacked probabilities in the field (prey dataset). The
216 sample size or unit of analysis in this part was based on the number of real or artificial
217 abstract prey. However, we also used the combined dataset that included both predator and

218 prey datasets, as detailed in the “**Meta-analysis and meta-regressions**” and “**Publication**
219 **bias**” sections.

220

221 **Effect size calculation**

222 To obtain the effect size point estimates and sampling variances, we used the natural
223 logarithm of the response ratio ($\ln RR$) between the means of the treatment and the treatment
224 control stimulus groups [43–45]. Positive $\ln RR$ values indicate heightened aversion in birds
225 and enhanced prey survival, while negative $\ln RR$ values signify diminished bird aversion and
226 increased prey mortality. The point estimate and sampling variance (var) of $\ln RR$ can be then
227 calculated in:

$$\ln RR = \ln\left(\frac{M_T}{M_C}\right) \quad (1)$$

$$\text{var}(\ln RR) = \frac{SD^2_T}{N_T M^2_T} + \frac{SD^2_C}{N_C M^2_C} - 2r \sqrt{\frac{SD^2_T}{N_T M^2_T}} \sqrt{\frac{SD^2_C}{N_C M^2_C}} \quad (2)$$

228 where M_T and M_C are mean responses of treatment and control groups (e.g., total frequency
229 of attacking prey, latency of approach, or prey survivability), respectively. SD and N are
230 (sample) standard deviations and sample size, respectively. The term, r is the correlation
231 coefficient between responses of the two groups. Some of our eligible studies used the paired
232 (dependent) study design where treatment and control samples originated from the same
233 individuals, and sample sizes between the two groups were the same. None of these studies
234 provided an estimate of r . Thus, when calculating our effect sizes, we assumed that this
235 correlation was 0.5, which is conservative [46]. For the other studies that used independent
236 study design, we set $r = 0$.

237 We note that our dataset included proportion (percentage) data (e.g., predator attack
238 rate or prey survival probability), which are bounded at 0 (0%) and 1 (100%). Therefore, we
239 transformed group means (M) and group standard deviations (SD) for proportion data using

240 Equations (3) and (4) before applying (1) and (2) to calculate lnRR and the sampling
241 variance:

$$f(M) = \arcsine(\sqrt{M}) \quad (3)$$

$$SD(f(M)) = \sqrt{\frac{SD^2}{4M(1-M)}} \quad (4)$$

242 where f indicates a function, in our case, the arcsine transformation. The standard deviation
243 (SD) related to this transformation was derived using the delta method before calculating
244 lnRR and the sampling variance [47]. We have also assumed that the standard deviation was
245 $SD(f(M)) = 1/\sqrt{8}$ if SD was not available.

246

247 **Meta-analysis and meta-regressions**

248 We used the *rma.mv* function from the package metafor v.4.4.0 [48] in R v.4.3.1 [49] for our
249 analyses. We started by fitting multilevel, mixed-effect meta-analytic models to the predator
250 and prey datasets. These meta-analytic models explicitly incorporated random factors, Study
251 ID, Cohort ID (groups of the same subjects), and Shared control ID (indicating effect sizes
252 sharing control groups) [50] along with Observation ID, fitted by the above function [48].

253 The model for the predator dataset included Species ID and a correlation matrix related to
254 phylogenetic relatedness for the species as random factors [51]. This is because we had data
255 on the bird species used in the experiment in the predator dataset, and we needed to control
256 for phylogenetic relationships between birds. We also quantified the total I^2 (a measure of
257 heterogeneity not attributed to sampling error [52]) and how much each random factor was
258 explained (partial I^2), calculated by the *i2_ml* function from the package orchaRd v.2.0.0
259 [53]. After running both meta-analytical models, we found that phylogeny and Species ID did
260 not need to be controlled for in the predator dataset, as their partial I^2 were zero ($I^2 = 0.00\%$).

261 That is, these factors explained little heterogeneity between effect sizes.

262 Therefore, we merged predator and prey datasets (i.e., full dataset) without
263 considering phylogenetic information and used them for the following models. We had, as
264 random effects, Study ID, Cohort ID, Shared control ID, and Observation ID for our meta-
265 analytic model using the full dataset. The Cohort ID and Shared control ID were removed
266 from our subsequent meta-regressions because they both explained little heterogeneity (both
267 partial $I^2 < 0.001\%$). This intercept-only (meta-analytic) model tested the conspicuous
268 patterns (eyesspots and non-eyesspots) that affected predator avoidance (i.e., our first question).

269 Next, we tested whether eyespots and non-eyespot patterns differ in the magnitude
270 and direction of the effect of elicited bird predator avoidance and what factors contribute to
271 the deterring effects of conspicuous patterns. We performed uni-moderator meta-regression
272 models with each of eight moderators: treatment stimulus pattern types (eyesspots vs. non-
273 eyesspots), pattern area, the number of pattern shapes, prey material type, maximum pattern
274 diameter/length, total pattern area, total area of prey surface, and prey shape type. We also
275 ran a multi-moderator meta-regression model, including the first four of the eight variables
276 mentioned in the uni-moderators, due to moderator correlations. We used log-transformed
277 data for pattern area, total pattern area, total area of prey surface, and pattern maximum
278 diameter/length in our analysis to normalise these moderators. We created all result plots in
279 the *orchard_plot* and *bubble_plot* functions from the package *orchaRd* [53].

280

281 **Publication bias**

282 We used three approaches to assess the presence of publication bias in our study. First, we
283 visually assessed the funnel plot asymmetry by examining the residuals from a meta-analytic
284 model, which included all the random factors utilised in our study. These residuals were
285 plotted against the precision of the effect sizes. Secondly, we performed an alternative
286 method to Egger's regression. This method used the inverse of the effective sample size as a
287 moderator within a multilevel meta-analytic model [54]. Third, we examined the possibility
288 of time-lag bias by including publication year as a moderator in our multilevel meta-analytic
289 model. Uni-moderator models were run for each inverse of the effective sample size and

290 publication year, and a multi-moderator model was carried out with the full model including
291 both inverse of the effective sample size and publication year as moderators.

292

293 **Additions and deviations**

294 We made two changes to the pre-registration: the addition of four new moderators and the
295 removal of two moderators. The new moderators were pattern area, total pattern area, total
296 area of prey surface, and prey shape types, although similar moderators were in the pre-
297 registration such as the number of eyespots (patterns) and diameter of an eyespot (a pattern).
298 These *post-hoc* decisions were taken to refine our initial moderators. We subsequently used
299 them in our meta-regression analyses. We originally intended to include the broad outcome
300 categories of predator avoidance measure as a moderator in the models, but the diversity of
301 reported results made categorisation impossible. Therefore, we did not include it as a
302 moderator. We also collected information on bird diet but decided not to include it. This
303 decision was because six of the seven bird species in our study were omnivores, resulting in a
304 lack of variability needed to detect diet effects in our data (for more details, please see
305 **Results**).

306

307 **Results**

308 **Screening outcomes and dataset characteristics**

309 We obtained 270 effect sizes from 33 studies for our analysis. The screening process and
310 reasons for exclusion at the full-text screening stage are summarised in the PRISMA-like
311 flowchart (Fig. 2a), with additional details available in Table S3, which comprises a list of
312 included/excluded studies. Of the dataset, 68.9% of effect sizes came from eyespot
313 presentation experiments (Fig. 2b). The remaining 31.1% of effect sizes came from non-
314 eyespot pattern presentation experiments (Fig. 2b). The latter category encompassed various
315 shapes, including circles (71.4%), rectangles (16.7%), diamonds (6.0%), complex patterns
316 (combinations of circles and diamonds; 4.8%), and stripes (1.1%); 93.7% of the control
317 stimuli used in these experiments involved the removal of the pattern used in the treatment
318 stimuli; the remaining stimuli were camouflage patterns (6.3%). Prey shape type used for

319 stimulus presentation varied from real or imitation of a particular butterfly (24.4%) to simply
320 a piece of paper (21.5%) (Fig. 2b). The number of pattern shapes varied between studies from
321 one to 11, but in most experiments, they were two (i.e., a pair of shapes; Fig. 2c).

322 Additionally, we found that the size of these patterns, both area and maximum
323 diameter/length, exhibited considerable variation across studies (Fig. 2c). The total area of
324 the patterns and stimulus also varied widely (Fig. 2c). The studies reported responses to
325 conspicuous pattern stimuli by seven bird species (Fig. 2d). Chickens (*Gallus gallus*) and
326 common starlings (*Sturnus vulgaris*) were the most studied birds in our dataset. Apart from
327 chickens (eight studies) and Eurasian blue tits (*Cyanistes caeruleus*; five studies), effect sizes
328 were available from just one or two studies per species. Six of the seven species were
329 omnivores, and one (yellow bunting; *Emberiza sulphurata*) was a granivore [55].

330

331 **Does the presence of conspicuous patterns affect predator avoidance?**

332 The overall mean effect size was statistically significant, showing a 21.86% (this percentage
333 value is the back-transformed values of lnRR) increase in the probability of predator
334 avoidance, such as higher prey survival rates or eliciting fewer attacks from birds (estimate =
335 0.20, 95% CI = [0.08, 0.31], $t_{[df = 268]} = 3.40$, $p = 0.0008$), in prey with conspicuous patterns
336 than in prey without such patterns (Fig. 3a). Total heterogeneity across effect sizes was high
337 ($I^2 = 96.50\%$); more specifically, observation ID (representing the within-study effect)
338 accounted for the most heterogeneity, 79.88%, with study ID (representing between-study
339 effect) accounting for the remaining 16.61%.

340

341 **Is there a difference in predator avoidance between eyespots and conspicuous patterns?**

342 There was no statistically significant difference between the effects of eyespots and non-
343 eyespot patterns ($F_{[df1 = 1, df2 = 268]} = 0.33$, $p = 0.57$, $R^2 = 0.27\%$; Fig. 3b). On average, eyespot
344 patterns resulted in 24.37% (estimate = 0.22, 95% CI = [0.08, 0.35], $t_{[df = 268]} = 3.17$, $p =$
345 0.002) and non-eyespot patterns in 17.11% (estimate = 0.16, 95% CI = [-0.02, 0.34], $t_{[df = 268]}$
346 = 1.71, $p = 0.09$) increases in predator avoidance compared with control stimuli, although this
347 trend was not statistically significant for non-eyespots (Fig. 3b).

348

349 **What factors promote predator avoidance?**

350 Our uni-moderator meta-regression model with pattern area (individual shape area) showed
351 that larger patterns were associated with an increase in predator avoidance (estimate = 0.11,
352 95% CI = [0.03, 0.19], $t_{[df=268]} = 2.71$, $p = 0.007$, $R^2 = 8.56\%$; Fig. 4a). The total pattern area
353 also promoted predator avoidance (estimate = 0.09, 95% CI = [0.004, 0.17], $t_{[df=268]} = 2.07$, p
354 = 0.04, $R^2 = 5.18\%$; Fig. S1a). Similarly, the maximum diameter/length of the pattern
355 positively influenced predator avoidance (estimate = 0.19, 95% CI = [0.04, 0.35], $t_{[df=268]} =$
356 2.46 , $p = 0.01$, $R^2 = 6.62\%$; Fig. S1b). In contrast, an increased number of pattern shapes
357 significantly reduced the effect of predator avoidance (estimate = -0.06, 95% CI = [-0.11, -
358 0.008], $t_{[df=268]} = -2.29$, $p = 0.02$, $R^2 = 2.46\%$; Fig. 4b). We found no significant effects of
359 total prey surface area on predator avoidance (estimate = -0.03, 95% CI = [-0.15, 0.09], $t_{[df=}$
360 $268]} = -0.48$, $p = 0.63$, $R^2 = 0.42\%$; Fig. S1c). Predator avoidance was not statistically
361 significantly affected by differences in whether the presented prey looked like a real
362 lepidopteran species ($F_{[df1=1, df2=268]} = 0.12$, $p = 0.72$, $R^2 = 0.13\%$). Both types of prey
363 material (real/imitation and abstract butterfly) had similar positive trends (Fig. 3c), with the
364 former increasing predator avoidance by 25.55% (estimate = 0.23, 95% CI = [0.03, 0.43], $t_{[df=}$
365 $268]} = 2.24$, $p = 0.03$) and the latter by 20.07% (estimate = 0.18, 95% CI = [0.04, 0.33], $t_{[df=}$
366 $268]} = 2.44$, $p = 0.02$). Further, when also considering prey type (Fig. S2), abstract and real
367 butterflies significantly exhibited increased predator avoidance by 37.98% (estimate = 0.32,
368 95% CI = [0.11, 0.53], $t_{[df=268]} = 3.04$, $p = 0.003$) and by 25.40% (estimate = 0.23, 95% CI =
369 [0.03, 0.42], $t_{[df=268]} = 2.25$, $p = 0.03$), respectively, but artificial abstract caterpillars
370 (estimate = 0.07, 95% CI = [-0.18, 0.31], $t_{[df=266]} = 0.53$, $p = 0.60$) and artificial abstract prey
371 (estimate = 0.01, 95% CI = [-0.35, 0.37], $t_{[df=266]} = 0.06$, $p = 0.95$) did not, respectively.
372 When comparing each prey type (e.g., abstract butterfly vs. real butterfly), none of the
373 differences was statistically significant (Fig. S2).

374 The multi-moderator (full) regression model showed that only pattern area positively
375 affected predator avoidance (estimate = 0.10, 95% CI = [0.009, 0.18], $t_{[df=266]} = 2.16$, $p =$
376 0.03; Table S4). Contrary to the uni-moderator regression model, the number of patterns

377 showed no significant effects on predator avoidance, although the consistent trend remained
378 (estimate = -0.05, 95% CI = [-0.11, 0.004], $t_{[df=266]} = -1.84$, $p = 0.07$; Table S4). The full
379 model accounted for 8.33% of the variation in the dataset. The complete output of the multi-
380 moderator model is displayed in Table S4.

381

382 **Publication bias**

383 The funnel plot showed no visual sign of funnel asymmetry (Fig. 5a). The meta-regression
384 analysis, which included the square root of the inverse of the effective sample size, further
385 supported this observation by showing that the effective sample size did not significantly
386 predict the effect size values (estimate = -0.09, 95% CI = [-0.83, 0.65], $t_{[df=266]} = -0.24$, $p =$
387 0.81; Fig. 5b). There was no detectable trend suggesting that more recent publications
388 consistently showed lower or higher effect size values, which would have indicated the
389 presence of time-lag publication bias (estimate = -0.0008; 95% CI = [-0.01, 0.01], $t_{[df=266]} =$
390 -0.12, $p = 0.90$; Fig. 5c). We obtained the same trends from multi-moderator meta-
391 regressions (Fig. S3).

392

393 **Discussion**

394 Eyespots and non-eyespot patterns did not differ significantly in the magnitude of deterring
395 effects (Fig. 3b). Avian predators showed similar avoidance responses to the conspicuous
396 patterns compared to control ones (Fig. 3a). Specifically, larger pattern sizes played a crucial
397 role in eliciting negative responses from birds (Fig. 4a). Further, negative responses from
398 birds showed the tendency to decline with increasing pattern number: single patterns were
399 likely more intimidating than a group of patterns (Fig. 4b). Taken together, our results
400 support the conspicuousness hypothesis rather than the eye mimicry hypothesis.

401

402 **Eye mimicry or conspicuous hypothesis?**

403 Overall, our meta-analysis showed that conspicuous patterns could increase predator
404 avoidance by over 20%. Specifically, our results indicate that conspicuousness per se can be
405 advantageous in avoiding bird predation (Fig. 3ab, Fig. 4). The evidence favouring the

406 conspicuousness hypothesis comes mainly from a series of field experiments by Steven and
407 his colleagues [30, 31, 34]. They showed that both eyespots and non-eyesspots improved the
408 prey survival similarly compared to non-conspicuous patterns [30, 31, 34]. In addition, their
409 research showed prey with more conspicuous patterns (i.e., large-size patterns) tended to
410 survive more than others [30, 31, 34], and eye resemblance (e.g., number or pattern shapes)
411 did not significantly affect the prey's survival [30, 31, 34]. Given that these pattern stimuli
412 used in the experiments are rarely or never found in natural environments [34], the most
413 parsimonious explanation for these results is neophobia or dietary conservatism in birds [56–
414 58]. Both phenomena appear to diminish with habituation and/or learning. A few studies
415 investigated such factors for intimidating effects, and they showed that repeated encounters
416 made birds more habituated to eyespot patterns [16, 59, 60]. We need more systematic tests
417 of bird habituation to aposematic-coloured patterns to better understand the evolution and
418 function of such patterns in Lepidoptera.

419 While our meta-analytic results favour the conspicuousness hypothesis, several
420 empirical studies support the eye mimicry hypothesis. For example, De Bona et al. [28] found
421 that a pair of eyespots of *Caligo martia* was as effective as true owl eyes and more efficient
422 in eliciting predator avoidance responses than less mimetic but equally contrasting circles.
423 Blut and Luau [61] created artificial eye-spotted prey with different similarities to the
424 vertebrate eyes and checked their survival rates in a field experiment. They revealed that the
425 prey with the most mimetic pattern had the highest survival rate [61]. Although studies on
426 Lepidoptera larvae are relatively limited, caterpillar eyespots are considered part of snake
427 mimicry [14]. Some research examined the benefit of eyespots by presenting artificial
428 caterpillars (marked with eyespots and control) made from dyed pastry to wild birds and
429 showed that eyespots improved survival [60, 62, 63]. Despite these convincing pieces of
430 empirical evidence, our meta-analytic results showed that eye resemblance did not improve
431 predator avoidance. If the eye mimic hypothesis was true, we would have seen a clear
432 difference between studies investigating eyespots and non-eyesspots.

433 However, we observed little heterogeneity among studies, despite finding high
434 heterogeneity within individual studies. This finding implies that if each study followed

435 similar experimental procedures within studies, our main result on predator avoidance would
436 be more generalisable. The high within-study heterogeneity can be caused by varying
437 stimulus characteristics contributing to the effect size variations, even in the same studies.
438 Bird phylogenetic relatedness explained little heterogeneity in our predator dataset, but this
439 may have occurred because a limited number of subject bird species (i.e., chickens, common
440 starlings, Eurasian blue tits) dominated our dataset (Fig. 2d). While we cannot exclude the
441 possibility of species differences in birds' responses to the conspicuous patterns, our analysis
442 indicated that bird species identity did not explain the observed variation in predator
443 avoidance.

444 We also note that conspicuous patterns can also be important for conspecific
445 communication in butterflies [12, 64]. Eyespots on *Bicyclus anynana* are known to function
446 as sexual signals. For example, males choose females depending on eyespot size and
447 reflectance [65]. Regarding the non-eyespot patterns, males of *Heliconius cydno* and *H.*
448 *pachinus* can recognise conspecific females by the bright colour of wing patches [66, 67].
449 Conspicuous patterns can also act as social signals in other taxa (e.g., birds: [68]), but this
450 function remains unclear in butterflies. Therefore, the diversity of patterns on wings could be
451 shaped by intra-specific and inter-specific communication. We should simultaneously
452 consider the influence of anti-predator and sexual/social signalling functions on the evolution
453 of butterfly conspicuous patterns [cf., 65, 69, 70].

454

455 **What factors explain the observed heterogeneity?**

456 The indicators of pattern size, including each pattern area (Fig. 4a), total pattern area (Fig.
457 S1a), and maximum diameter/length (Fig. S1b), were the most important moderators of effect
458 sizes, overall indicating that large patterns could promote predator avoidance. Notably, these
459 size metrics were correlated, so they are not independent of each other. Several studies
460 suggested that the pattern size difference is related to the difference in prey survival [21, 26,
461 30]. For example, eyespots larger than 6.0 mm may have a strong deterrent effect with
462 increasing size [26], but such patterns may increase the visibility of lepidopterans, and their
463 presence may increase predation rates as well [71]. Indeed, small conspicuous patterns tend

464 to attract predators' attention, as explained by the deflection hypothesis [12, 72]. The effect
465 may contribute to the observed negative overall effect sizes (Fig. 3, Fig. 4). Considering
466 studies on *B. anynana* with eyespots with a deflecting effect (maximum diameter is about 5.0
467 mm; Table S5), a size of at least 6.0 mm is required to avoid predator approach. However, it
468 is uncertain whether the effect would linearly increase with size or whether an optimal size
469 exists. Although eyespot sizes on actual Lepidoptera may be restricted by their body or wing
470 size ([e.g., 73], but see also [21]), it would be interesting to find a maximum threshold for
471 patterns that promote predator avoidance responses in birds.

472 Among other moderators tested (prey material type, total pattern area, and prey shape
473 type), the only moderator that seemed to explain heterogeneity was the number of patterns
474 (Fig. 4b; yet it is likely inconclusive; see Table S4). Previous studies predominantly
475 employed a single pattern or a pair of patterns, leading to limited variations. Nonetheless, our
476 findings indicate that a single eyespot is equally or more effective than a pair of eyespots.
477 Consequently, the resemblance to a pair of eyes, a crucial aspect of the eye mimicry
478 hypothesis, may not be essential for effective predator avoidance. To disentangle the two
479 hypotheses, we recommend conducting the following experiments with two key features [30,
480 35, 74]: a set of stimuli that (1) have the same size (area or diameter/maximum length of each
481 pattern or total pattern area) but with different numbers of patterns ranging from a few
482 usually found in Lepidoptera to numerous patterns unlike those seen in them, and (2) are
483 presented with the same number of patterns and the same size but different pattern shapes.
484 Results from these experiments could deepen our current knowledge, allowing us to inch
485 toward a more definitive answer.

486

487 **Knowledge gaps and future opportunities**

488 Along with other conspicuous patterns, eyespots are believed to deter bird predation,
489 and our meta-analysis supports this function. However, five major gaps remain in the current
490 literature and our knowledge. First, birds and humans likely perceive eye-like shapes
491 differently based on the interspecific diversity of bird vision [75]. Thus, it could be premature
492 to conclude that eyespots on Lepidoptera resemble vertebrate eyes universally. For example,

493 most bird species can detect ultraviolet light, which is invisible to humans, and the ultraviolet
494 reflection of the butterflies' eyespots may contribute to predator avoidance [e.g., 20, 22].

495 Second, some lepidopterans present conspicuous patterns to potential predators in
496 combination with other elements, such as sounds and movements [13, 16, 17, 76, 77],
497 presumably to emphasise the conspicuousness of the patterns. Most of the current literature
498 does not take these effects into account in experiments, although some studies argue in favour
499 or against their importance [e.g., 16, 17]. We should also consider how factors other than
500 those constituting the pattern (e.g., colour, number, and size) are involved in the predator
501 avoidance function of eyespots. The location of the butterfly's eyespot patterns varies from
502 species to species as well; eyespots exist on the wings' ventral, dorsal, or both sides. Not only
503 the dorsal eyespot patterns, which were used in most studies, but also the ventral eyespot
504 patterns should be explored. In addition, we need to avoid presenting patterns unnaturally
505 when using real butterflies in experiments. For example, many owl butterflies (family Caligo)
506 have a pair of eyespot patterns on the ventral side. Their eyespots are usually visible to birds
507 when the wings are closed and would not present side by side as in the eyes of the owl's
508 frontal face.

509 Third, recent studies have shown that birds are sensitive to the gaze of other
510 individuals and may respond more aversively when their gazes are directed at them [e.g., 78-
511 80]. Skelhorn and Rowland [81] showed that the anti-predation effect may be further
512 enhanced if the inner circle of the eyespot is in a more gazing-like position for subject birds.
513 However, further research is needed to investigate the importance of the position of the inner
514 circle.

515 Fourth, as mentioned above, studies focusing on caterpillar eyespots are much more
516 scarce compared to butterflies; Hossie and Sherratt [82] have shown similarities between
517 caterpillars and snakes, but the response of birds to actual caterpillars has not been
518 experimentally tested. Conversely, in butterflies, similarities between the eyespot patterns on
519 wings and the eyes of birds of prey have not been investigated.

520 Finally, birds are generally considered as potential predators of butterflies and
521 caterpillars. Although other taxa species, such as invertebrates [83–85], lizards [27, 86, 87],

522 and rats [88–91], are also known to prey on lepidopterans, there are much fewer studies using
523 non-avian species as predators. Therefore, we should expand the range of taxa used for
524 experiments to get a better and more generalisable understanding of the eyespots' function
525 and evolution in butterflies and caterpillars.

526 Knowing the effects of conspicuous patterns may contribute to creating a world where
527 birds and humans can live more harmoniously. Both eyespots and conspicuous patterns have
528 already been used to control birds, particularly in agriculture, although their effectiveness has
529 been questioned [e.g., 92, 93]. Such uncertainty may reflect our limited understanding of why
530 birds avoid eyespots and conspicuous patterns. Nevertheless, visual stimuli are less likely to
531 harm birds or affect the natural environment than others (e.g., nest/egg destructions or toxic
532 chemicals; reviewed in [94]). Therefore, when proven effective, they could be used for better
533 pest control, population management and conservation [95].

534

535 **Conclusion**

536 We have shed light on a traditional but controversial research topic that has fascinated
537 behavioural ecologists for decades. Our findings provide a better understanding of the
538 evolution of signal designs, but also show that more work is needed to understand the
539 function of the eyespot patterns in Lepidoptera, such as whether eyespot patterns evolved due
540 to mimicry or conspicuousness.

541

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544

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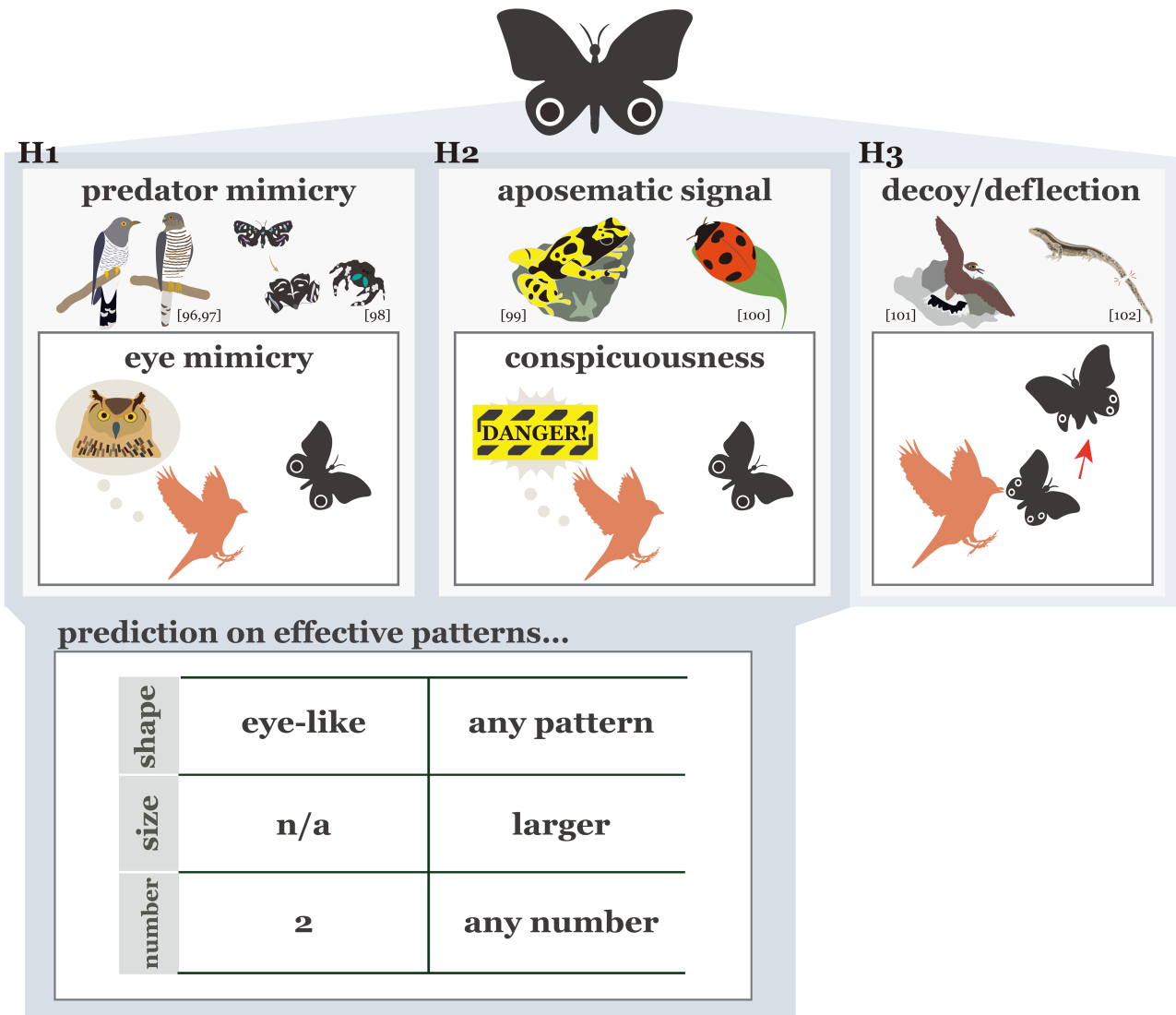
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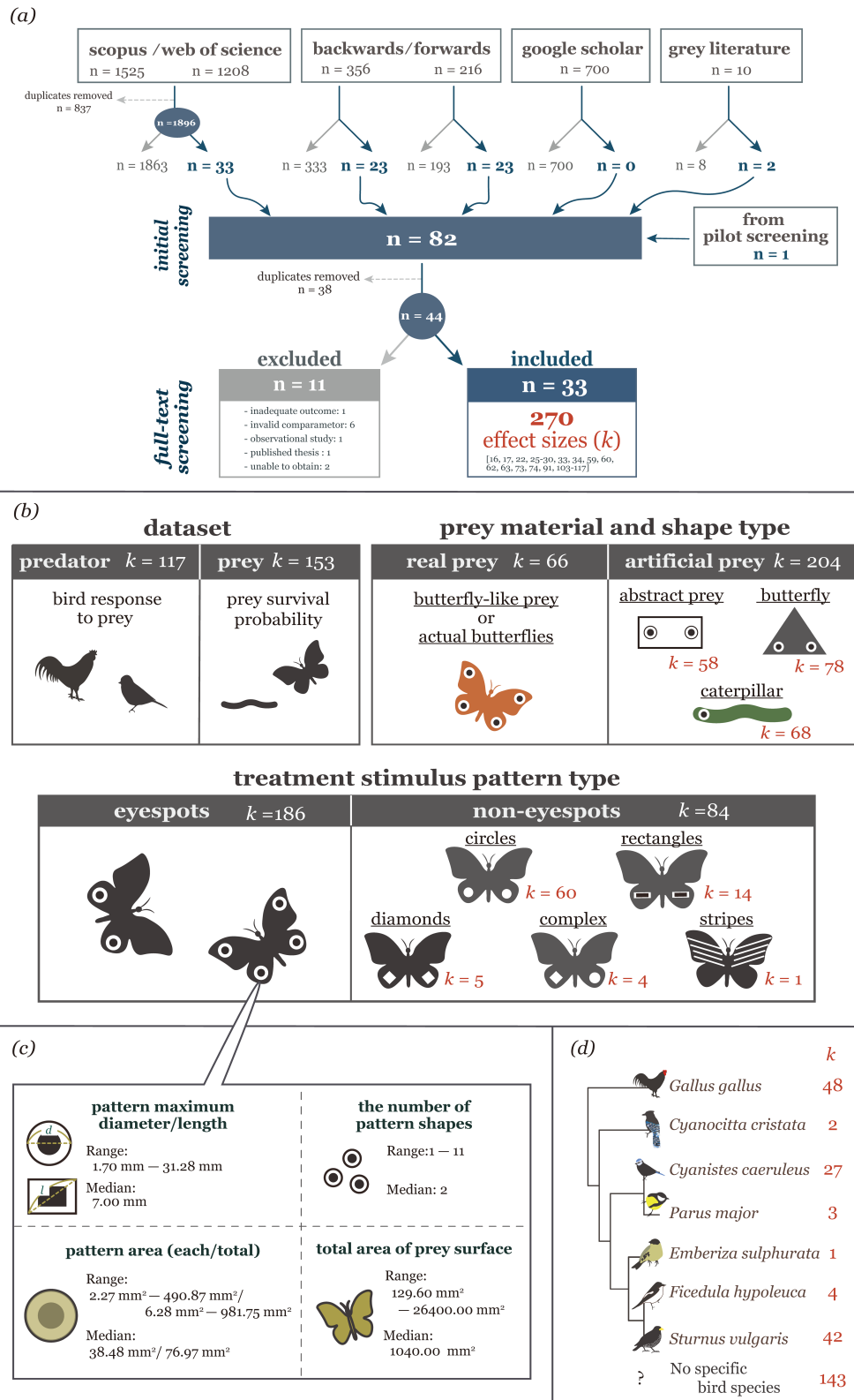
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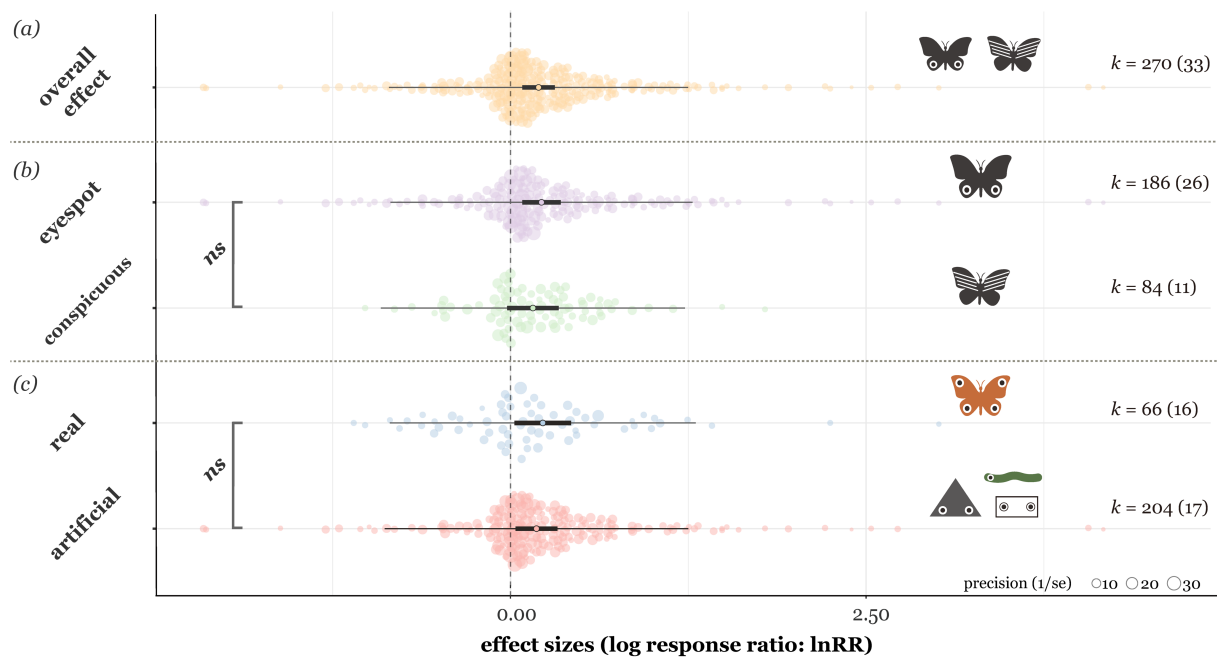
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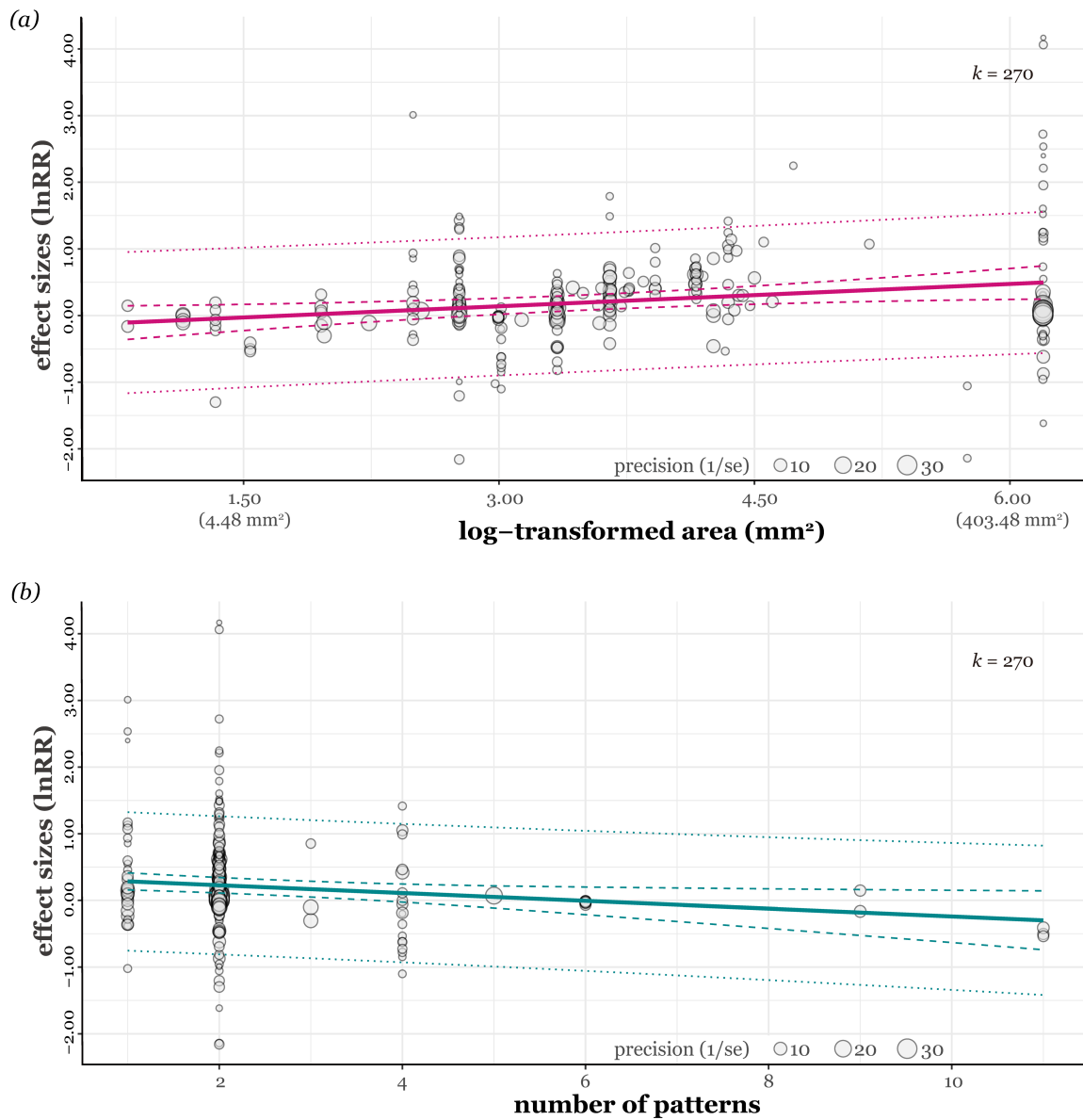
857 **Figure 1.** A visual summary of three hypotheses that explain the predation avoidance
 858 function of eyespot patterns and the predictions that can be derived from these two
 859 hypotheses.



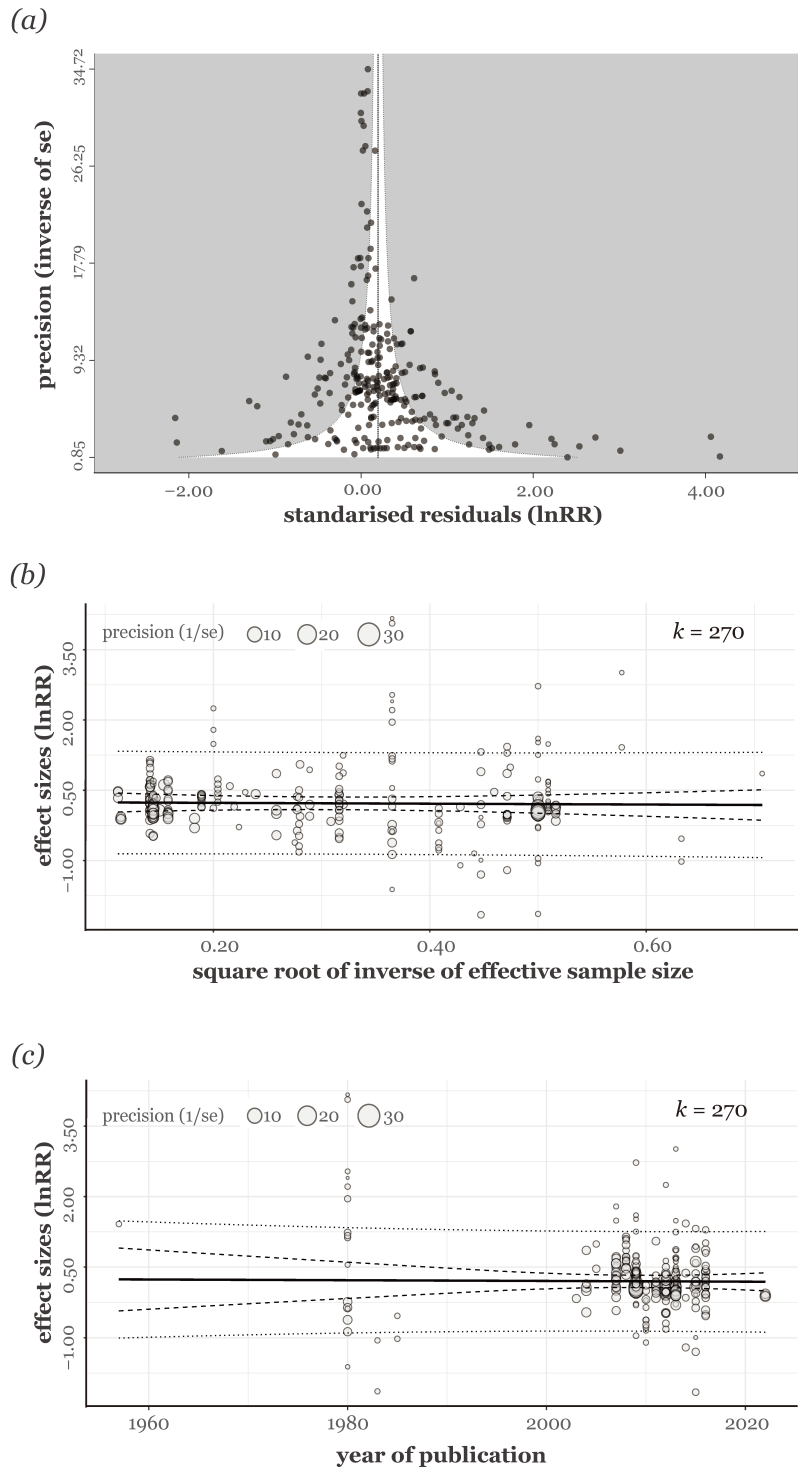
860 **Figure 2.** Overview of the dataset. (a) shows a PRISMA-like flowchart of the systematic
861 literature search for the meta-analysis. (b) and (c) give details of the main moderators
862 examined in the meta-analysis. (d) provides the phylogenetic tree of bird species included in
863 the meta-analysis, together with the sample sizes and number of effect sizes per species.



864 **Figure 3.** Mean effect sizes of (a) overall for all highly salient patterns, (b) effects split by
 865 experiments with eyespots versus conspicuous patterns, and (c) two prey types used in the
 866 experiments. Thick horizontal lines represent 95% confidence intervals, and thin horizontal
 867 lines represent 95% prediction intervals. The points in the centre of each thick line indicate
 868 the average effect size. *k* is the number of effect sizes used to estimate the statistics, followed
 869 by the number of studies in the brackets.



870 **Figure 4.** The relationships between (a) prey salient pattern area (log-transformed) and effect
 871 sizes and (b) number of prey salient patterns and effect sizes. Circle sizes are scaled
 872 according to precision, k represents the number of effect sizes. Each fitted regression line is
 873 shown as a coloured straight line, and 95% confidence and prediction intervals are shown as
 874 dashed and dotted coloured lines, respectively.

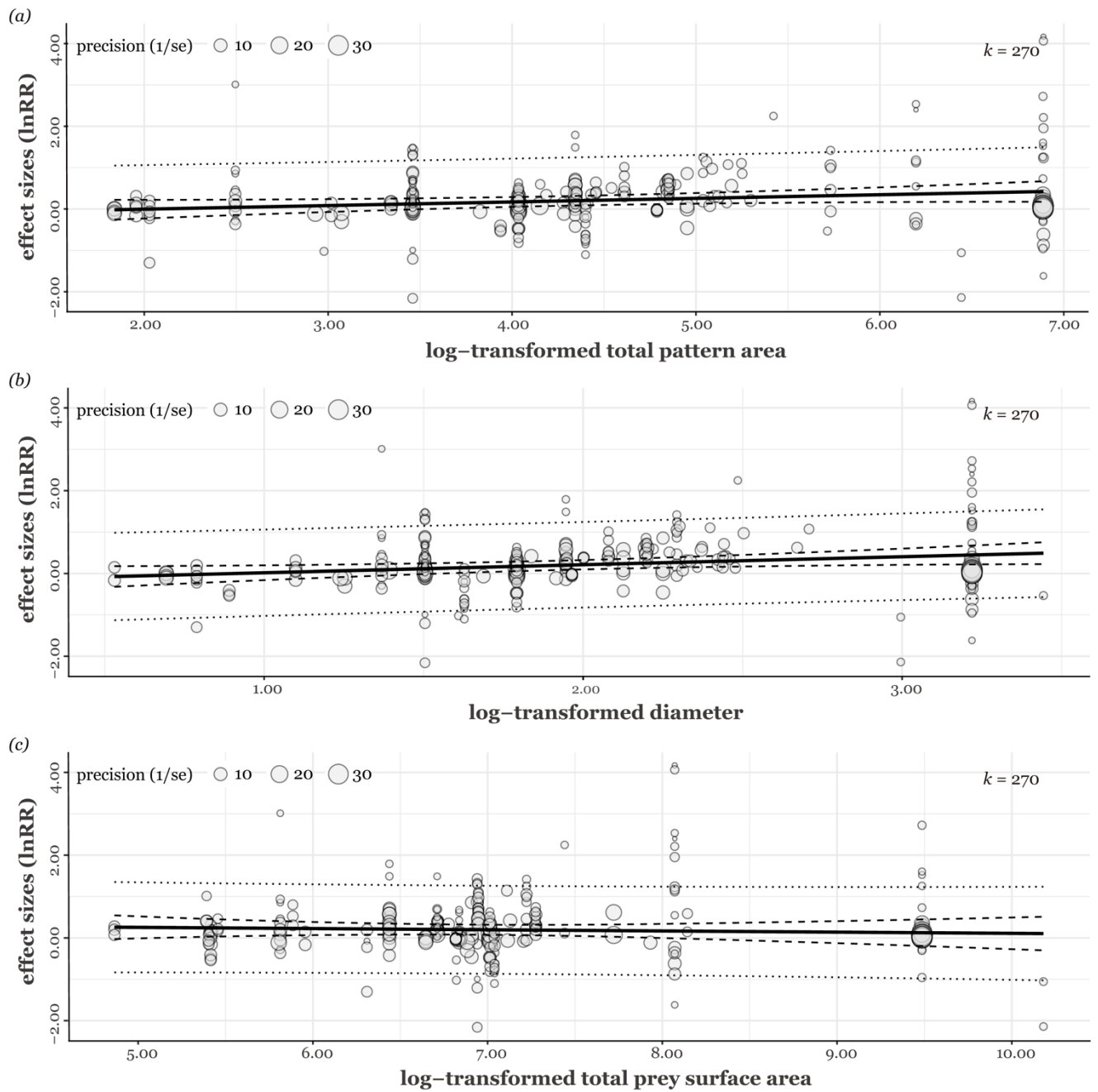


875 **Figure 5.** (a) Funnel plot using effect size and its inverse standard error. The relationship
 876 between effect sizes and (b) the square root of the inverse of effective sample size and (c)
 877 publication year. In (b) and (c), circle sizes are scaled accordingly to precision, and k
 878 represents the number of effect sizes. Each fitted regression line is shown as a straight line,
 879 and 95% confidence and prediction intervals are shown as dashed and dotted lines,
 880 respectively.

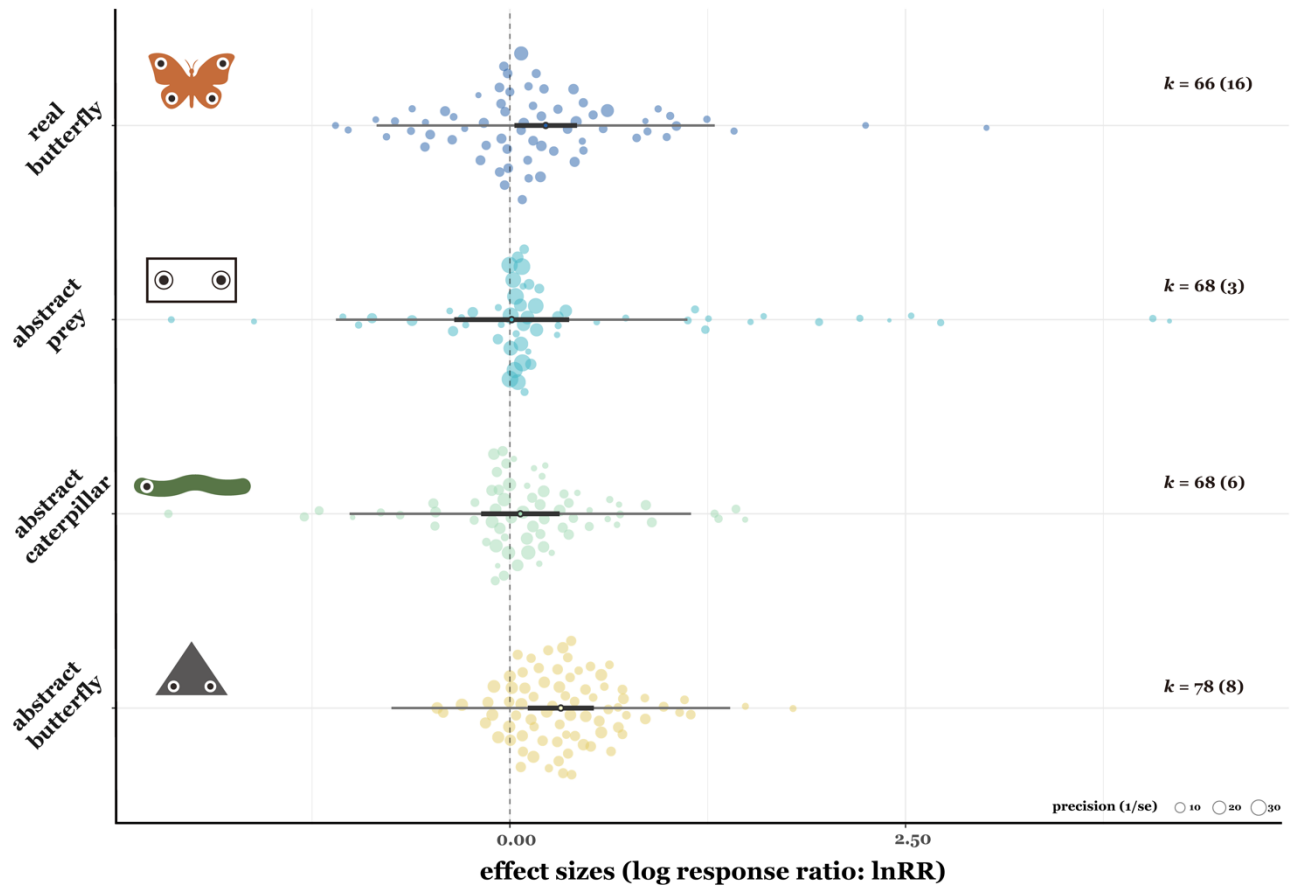
881 **Table 1.** Descriptions of the population, Intervention, comparator, and outcome (PICO) used
882 to define the scope of this study.

PICO	Description
Population	Birds as predators and butterflies, moths, caterpillars, and their models as prey
Intervention	Presenting eyespot or conspicuous pattern stimulus to birds
Comparator	Presenting stimulus that is neither eyespot nor conspicuous patterns
Outcome	Avian behavioural responses to eyespot or conspicuous pattern stimuli The probability of prey surviving or being attacked (for the stimuli)

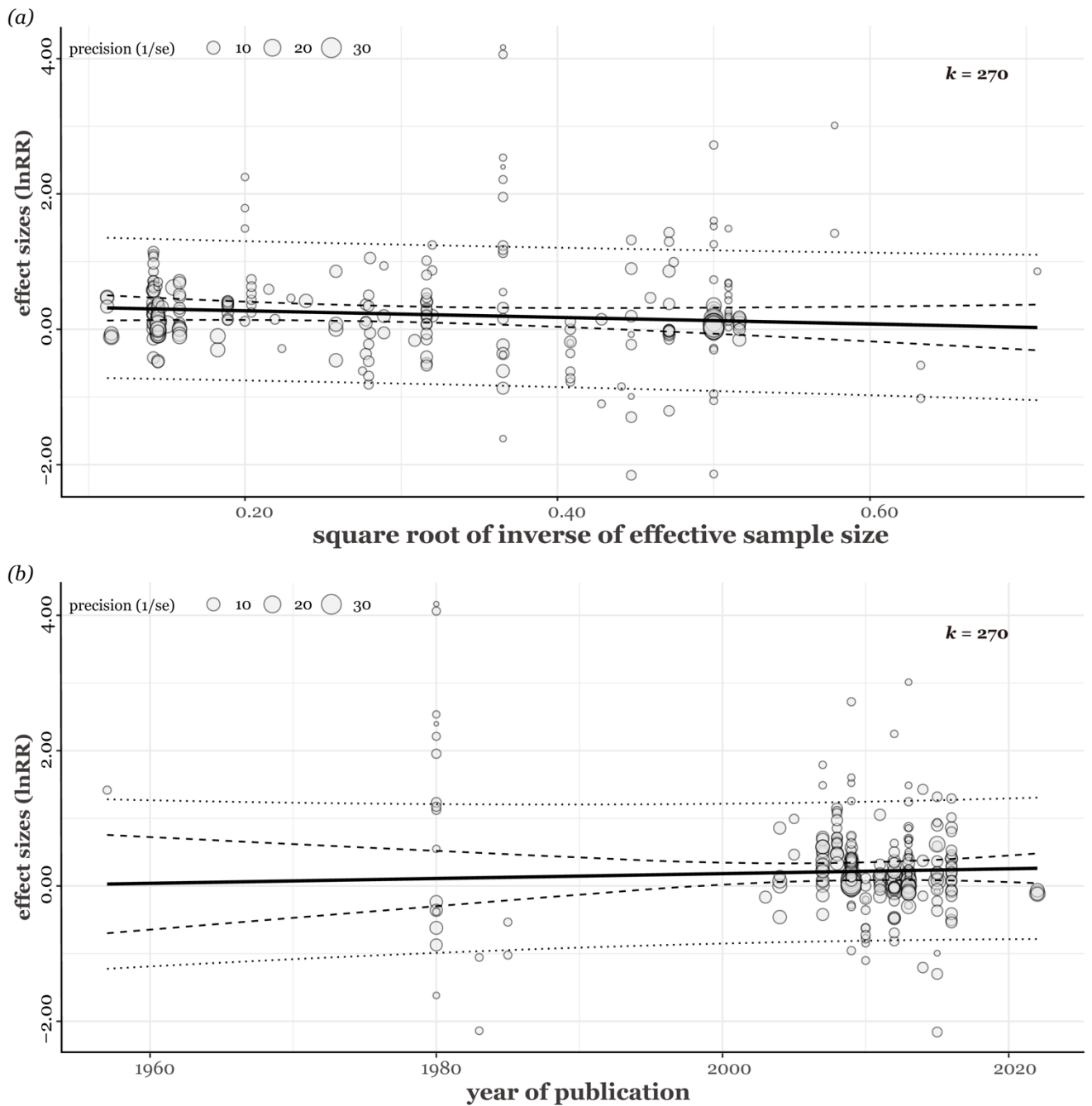
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886 **Figure S1.** The relationships between (a) total pattern area, (b) pattern maximum
 887 diameter/length, and (c) total prey surface area and effect sizes. k shows the number of effect
 888 sizes. Each fitted regression line is shown as a solid straight line, and 95% confidence and
 889 prediction intervals are shown as dashed and dotted lines, respectively.



890 **Figure S2.** Mean effect sizes of total prey shape types. Thick horizontal lines represent 95%
 891 confidence intervals, and thin horizontal lines represent prediction intervals. The points in the
 892 centre of each thick line indicate the average effect size. k shows the number of effect sizes.



893 **Figure S3.** (a) relationship between effect size and the square root of the inverse of effective
 894 sample size, and (b) relationship between effect size and publication year. Both plots were
 895 based on the multi-moderator model. k shows the number of effect sizes. Each fitted
 896 regression line is shown as a solid straight line, and 95% confidence intervals and prediction
 897 intervals are shown as dashed and dotted lines, respectively.

Checklist item	Sub-item number	Sub-item	Reported by authors?	Notes
Title and abstract	1.1	Identify the review as a systematic review, meta-analysis, or both	Yes	
	1.2	Summarise the aims and scope of the review	Yes	
	1.3	Describe the data set	Yes	
	1.4	State the results of the primary outcome	Yes	
	1.5	State conclusions	Yes	
	1.6	State limitations	Yes	
Aims and questions	2.1	Provide a rationale for the review	Yes	
	2.2	Reference any previous reviews or meta-analyses on the topic	Yes	
	2.3	State the aims and scope of the review (including its generality)	Yes	
	2.4	State the primary questions the review addresses (e.g. which moderators were tested)	Yes	
	2.5	Describe whether effect sizes were derived from experimental and/or observational comparisons	Yes	
Review registration	3.1	Register review aims, hypotheses (if applicable), and methods in a time-stamped and publicly accessible archive and provide a link to the registration in the methods section of the manuscript. Ideally registration occurs before the search, but it can be done at any stage before data analysis.	Yes	
	3.2	Describe deviations from the registered aims and methods	Yes	
	3.3	Justify deviations from the registered aims and methods	Yes	
Eligibility criteria	4.1	Report the specific criteria used for including or excluding studies when screening titles and/or abstracts, and full texts, according to the aims of the systematic review (e.g. study design, taxa, data availability)	Yes	

	4.2	Justify criteria, if necessary (i.e. not obvious from aims and scope)	Yes	
Finding studies	5.1	Define the type of search (e.g. comprehensive search, representative sample)	Yes	
	5.2	State what sources of information were sought (e.g. published and unpublished studies, personal communications)	Yes	
	5.3	Include, for each database searched, the exact search strings used, with keyword combinations and Boolean operators	Yes	
	5.4	Provide enough information to repeat the equivalent search (if possible), including the timespan covered (start and end dates)	Yes	
Study selection	6.1	Describe how studies were selected for inclusion at each stage of the screening process (e.g. use of decision trees, screening software)	Yes	
	6.2	Report the number of people involved and how they contributed (e.g. independent parallel screening)	Yes	
Data collection process	7.1	Describe where in the reports data were collected from (e.g. text or figures)	Yes	
	7.2	Describe how data were collected (e.g. software used to digitize figures, external data sources)	Yes	
	7.3	Describe moderator variables that were constructed from collected data (e.g. number of generations calculated from years and average generation time)	Yes	
	7.4	Report how missing or ambiguous information was dealt with during data collection (e.g. authors of original studies were contacted for missing descriptive statistics, and/or effect sizes were calculated from test statistics)	Yes	
	7.5	Report who collected data	Yes	
	7.6	State the number of extractions that were checked for accuracy by co-authors	No	

Data items	8.1	Describe the key data sought from each study	Yes	
	8.2	Describe items that do not appear in the main results, or which could not be extracted due to insufficient information	Yes	
	8.3	Describe main assumptions or simplifications that were made (e.g. categorising both 'length' and 'mass' as 'morphology')	NA: no assumptions or simplifications needed to be made	
	8.4	Describe the type of replication unit (e.g. individuals, broods, study sites)	Yes	
Assessment of individual study quality	9.1	Describe whether the quality of studies included in the systematic review or meta-analysis was assessed (e.g. blinded data collection, reporting quality, experimental <i>versus</i> observational)	No	
	9.2	Describe how information about study quality was incorporated into analyses (e.g. meta-regression and/or sensitivity analysis)	No	
Effect size measures	10.1	Describe effect size(s) used	Yes	
	10.2	Provide a reference to the equation of each calculated effect size (e.g. standardised mean difference, log response ratio) and (if applicable) its sampling variance	Yes	
	10.3	If no reference exists, derive the equations for each effect size and state the assumed sampling distribution(s)	Yes	
Missing data	11.1	Describe any steps taken to deal with missing data during analysis (e.g. imputation, complete case, subset analysis)	NA: there were no missing data	
	11.2	Justify the decisions made to deal with missing data	NA: there were no missing data	
Meta-analytic model description	12.1	Describe the models used for synthesis of effect sizes	Yes	
	12.2	The most common approach in ecology and evolution will be a random-effects model, often with a hierarchical/multilevel structure. If other types of models are chosen (e.g. common/fixed effects model, unweighted model), provide justification for this choice	NA: only (weighted) random-effects models were used	

Software	13.1	Describe the statistical platform used for inference (e.g. <i>R</i>)	Yes	
	13.2	Describe the packages used to run models	Yes	
	13.3	Describe the functions used to run models	Yes	
	13.4	Describe any arguments that differed from the default settings	Yes	
	13.5	Describe the version numbers of all software used	Yes	
Non-independence	14.1	Describe the types of non-independence encountered (e.g. phylogenetic, spatial, multiple measurements over time)	Yes	
	14.2	Describe how non-independence has been handled	Yes	
	14.3	Justify decisions made	Yes	
Meta-regression and model selection	15.1	Provide a rationale for the inclusion of moderators (covariates) that were evaluated in meta-regression models	Yes	
	15.2	Justify the number of parameters estimated in models, in relation to the number of effect sizes and studies (e.g. interaction terms were not included due to insufficient sample sizes)	Yes	
	15.3	Describe any process of model selection	Yes	
Publication bias and sensitivity analyses	16.1	Describe assessments of the risk of bias due to missing results (e.g. publication, time-lag, and taxonomic biases)	Yes	
	16.2	Describe any steps taken to investigate the effects of such biases (if present)	Yes	
	16.3	Describe any other analyses of robustness of the results, e.g. due to effect size choice, weighting or analytical model assumptions, inclusion or exclusion of subsets of the data, or the inclusion of alternative moderator variables in meta-regressions	Yes	
Clarification of <i>post hoc</i> analyses	17.1	When hypotheses were formulated after data analysis, this should be acknowledged.	NA: there are no hypotheses that were formed after data collection	
	18.1	Share metadata (i.e. data descriptions)	Yes	

Metadata, data, and code	18.2	Share data required to reproduce the results presented in the manuscript	Yes	
	18.3	Share additional data, including information that was not presented in the manuscript (e.g. raw data used to calculate effect sizes, descriptions of where data were located in papers)	Yes	
	18.4	Share analysis scripts (or, if a software package with graphical user interface (GUI) was used, then describe full model specification and fully specify choices)	Yes	
Results of study selection process	19.1	Report the number of studies screened	Yes	
	19.2	Report the number of studies excluded at each stage of screening	Yes	
	19.3	Report brief reasons for exclusion from the full text stage	Yes	
	19.4	Present a Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA)-like flowchart (www.prisma-statement.org).	Yes	
Sample sizes and study characteristics	20.1	Report the number of studies and effect sizes for data included in meta-analyses	Yes	
	20.2	Report the number of studies and effect sizes for subsets of data included in meta-regressions	Yes	
	20.3	Provide a summary of key characteristics for reported outcomes (either in text or figures; e.g. one quarter of effect sizes reported for vertebrates and the rest invertebrates)	Yes	
	20.4	Provide a summary of limitations of included moderators (e.g. collinearity and overlap between moderators)	Yes	
	20.5	Provide a summary of characteristics related to individual study quality (risk of bias)	Yes	
Meta-analysis	21.1	Provide a quantitative synthesis of results across studies, including estimates for the mean effect size, with confidence/credible intervals	Yes	

Heterogeneity	22.1	Report indicators of heterogeneity in the estimated effect (e.g. I^2 , τ^2 and other variance components)	Yes	
Meta-regression	23.1	Provide estimates of meta-regression slopes (i.e. regression coefficients) and confidence/credible intervals	Yes	
	23.2	Include estimates and confidence/credible intervals for all moderator variables that were assessed (i.e. complete reporting)	Yes	
	23.3	Report interactions, if they were included	NA: no interactions were included	
	23.4	Describe outcomes from model selection, if done (e.g. R^2 and AIC)	Yes	Please see the link provided in the Data Accessibility.
Outcomes of publication bias and sensitivity analyses	24.1	Provide results for the assessments of the risks of bias (e.g. Egger's regression, funnel plots)	Yes	
	24.2	Provide results for the robustness of the review's results (e.g. subgroup analyses, meta-regression of study quality, results from alternative methods of analysis, and temporal trends)	Yes	
Discussion	25.1	Summarise the main findings in terms of the magnitude of effect	Yes	
	25.2	Summarise the main findings in terms of the precision of effects (e.g. size of confidence intervals, statistical significance)	Yes	
	25.3	Summarise the main findings in terms of their heterogeneity	Yes	
	25.4	Summarise the main findings in terms of their biological/practical relevance	Yes	
	25.5	Compare results with previous reviews on the topic, if available	Yes	
	25.6	Consider limitations and their influence on the generality of conclusions, such as gaps in the available evidence (e.g. taxonomic and geographical research biases)	Yes	

Contributions and funding	26.1	Provide names, affiliations, and funding sources of all co-authors	Yes	
	26.2	List the contributions of each co-author	Yes	
	26.3	Provide contact details for the corresponding author	Yes	
	26.4	Disclose any conflicts of interest	NA: there were no conflicts of interest	
References	27.1	Provide a reference list of all studies included in the systematic review or meta-analysis	Yes	
	27.2	List included studies as referenced sources (e.g. rather than listing them in a table or supplement)	Yes	

900 **Table S2.** Search strings used for each database. We accessed Scopus, ISI Web of Science
901 core collection, Google Scholar (*Japanese, Polish, Portuguese, Russian, Spanish, Simplified*
902 *Chinese, and Traditional Chinese*) on 08/06/2023, and Bielefeld Academic Search Engine
903 (BASE) on 26/06/2023. BASE was used as a source of grey literature. We conducted
904 backward and forward reference searches for key review articles using Scopus on
905 19/06/2023. We modified search strings to collect studies to capture studies examining the
906 effects of eyespot patterns on birds using experimental methods. Search strings were adapted
907 to the structure of each database.

Database	Search strings
Scopus	TITLE-ABS-KEY (((eyespot* OR eye-spot* OR "eye spot*" OR eye-like* OR "eye like*" OR eye-mimic* OR "eye mimic*" OR "eye similari*" OR "predator* eye*" OR "eye similar*" OR concentric*) AND (attack* OR antipredator* OR anti-predator* OR aposematic* OR avoid* OR conspicuous* OR warn* OR fear* OR intimidat* OR predator-prey* OR butterfl* OR moth* OR bird* OR avian* OR caterpillar* OR prevent* OR aves OR passeri*)) AND NOT (fish* OR manti* OR lizard* OR bat* OR nano* OR health* OR patients OR women OR men OR children OR pediatric OR medic* OR hormon* OR genes OR magnet* OR valve* OR fluid* OR concrete OR beam* OR tissue* OR charge* OR energ* OR electro*)))
ISI Web of Science	TS = (((eyespot* OR eye-spot* OR "eye spot*" OR eye-like* OR "eye like*" OR eye-mimic* OR "eye mimic*" OR "eye similari*" OR "predator* eye*" OR "eye similar*" OR concentric*) AND (attack* OR antipredator* OR anti-predator* OR aposematic* OR avoid* OR conspicuous* OR warn* OR fear* OR intimidat* OR predator-prey* OR butterfl* OR moth* OR bird* OR avian* OR caterpillar* OR prevent* OR aves OR passeri*)) NOT (fish* OR manti* OR lizard* OR bat* OR nano* OR health* OR patients OR women OR men OR children OR pediatric OR medic* OR hormon* OR genes OR magnet* OR valve* OR fluid* OR concrete OR beam* OR tissue* OR charge* OR energ* OR electro*)))
BASE	eyespot* AND (avoid* predator* prevent* intimidat* mimi*) AND (ave* bird* passerine* butterfl* moth* lepidoptera caterpillar*) AND (experiment* stud*)
Google scholar	eyespot avoid predator prevention intimidation mimic aves bird passerine butterfly moth lepidoptera caterpillar experiment study

We translated the above English search string into *Japanese, Polish, Portuguese, Russian, Spanish, Simplified Chinese, and Traditional Chinese* for searching on Google Scholar.

Japanese:

目玉模様|眼状紋 忌避|捕食|防除|威嚇|擬態 鳥|鳴禽|蝶|蛾|鱗翅目
|芋虫|幼虫 実験|研究

Polish:

oko|oczy skrzydla|wzor|plama
ochrona|unikanie|drapieźnik|zapobieganie|zastraszenie
ptak|motyl|gasiénica|owad eksperyment|badania

Portuguese:

ocelo|"mancha ocelar"|"olhos falsos"|"falsos olhos"
evitar|predador|prevenção|intimidação
ave|pássaro|borboleta|mariposa|lagarta experimento|estudo

Russian:

глаз|глаза избегать|хищник|профилактика|запугивание
птица|бабочка|мотылек|Воробьинообразные|Чешуекрылые|Гусеница
эксперимент|изучать

Spanish:

ocelo|"ojos falsos"|"falsos ojos"
evitar|depredador|prevención|intimidación
ave|pájaro|mariposa|polilla|oruga experimento|estudio

Simplified chinese:

眼点 避免|捕食者|预防|恐吓|模仿 鸟类|鸟|雀|蝴蝶|蛾|鳞翅目|
毛毛虫 实验|试验|学习

Traditional chinese:

眼點 避免|捕食者|預防|恐嚇|模仿 鳥類|鳥|雀|蝴蝶|蛾|鱗翅目|
毛毛蟲 實驗|試驗|學習

909 **Table S3.** List of (a)included and (b) excluded studies at the full-text screening stage with

910 exclusion reasons.

911 (a) included studies

title	year	authors	journal	doi
The Function of Eyespot Patterns in the Lepidoptera	1957	Blest, AD.	Behaviour	10.1163/156853956X00048
Reactions of male domestic chicks to two-dimensional eye-like shapes	1980	Jones, RB.	Animal Behaviour	10.1016/S0003-3472(80)80025-X
The Feeding Behaviour of Starlings (<i>Sturnus vulgaris</i>) in the Presence of 'Eyes'	1983	Inglis, IR., Huson, LW., Marshall, MB. and Neville, PA.	Zeitschrift für Tierpsychologie	10.1111/j.1439-0310.1983.tb02151.x
Butterfly wing markings are more advantageous during handling than during the initial strike of an avian predator	1985	Wourms, MK. and Wasserman, FE.	Evolution	10.1111/j.1558-5646.1985.tb00426.x
Significance of butterfly eyespots as an anti-predator device in ground-based and aerial attacks	2003	Lyytinen, A., Brakefield, PM. and Mappes, J.	Oikos	10.1034/j.1600-0706.2003.11935.x
Does predation maintain eyespot plasticity in <i>Bicyclus anynana</i> ?	2004	Lyytinen, A., Brakefield, PM., Lindström, L., and Mappes, J.	Proceedings of the Royal Society B: Biological Sciences	10.1098/rspb.2003.2571
Asymmetry in size, shape, and color impairs the protective value of conspicuous color patterns	2004	Forsman, A. and Herretröm, J.	Behavioral Ecology	10.1093/beheco/arg092
Prey survival by predator intimidation: an experimental study of peacock butterfly defence against blue tits	2005	Vallin, A, Jakobsson, S., Lind, J. and Wiklund, C.	Proceedings of the Royal Society B: Biological Sciences	10.1098/rspb.2004.3034

Field experiments on the effectiveness of 'eyespot' as predator deterrents	2007	Stevens, M., Hopkins, E., Hinde, W., Adcock, A., Connolly, Y., Troscianko, T. and Cuthill, IC.	Animal Behaviour	10.1016/j.anbehav.2007.01.031
The anti-predator function of 'eyespot' on camouflaged and conspicuous prey	2008	Stevens, M., Stubbins, CL. and Hardman, CJ.	Behavioral Ecology and Sociobiology	10.1007/s00265-008-0607-3
Conspicuousness, not eye mimicry, makes "eyespot" effective antipredator signals	2008	Stevens, M., Hardman, CJ. and Stubbins, CL.	Behavioral Ecology	10.1093/beheco/arm162
The protective value of conspicuous signals is not impaired by shape, size, or position asymmetry	2009	Stevens, M., Castor-Perry, SA. and Price, JRF.	Behavioral Ecology	10.1093/beheco/arm119
The function of animal 'eyespot': Conspicuousness but not eye mimicry is key	2009	Stevens, M., Cantor, A., Graham, J. and Winney, IS.	Current Zoology	10.1093/czoolo/55.5.319
Fixed eyespot display in a butterfly thwarts attacking birds	2009	Kodandaramaiah, U., Vallin, A. and Wiklund, C.	Animal Behaviour	10.1016/j.anbehav.2009.02.018
Can we use starlings' aversion to eyespot as the basis for a novel 'cognitive bias' task?	2009	Brilot, BO., Normandale, CL., Parkin, A. and Bateson, M.	Applied Animal Behaviour Science	10.1016/j.applanim.2009.02.015
Constant eyespot display as a primary defence-survival of male and female emperor moths attacked by blue tits	2010	Vallin, A., Sven J. and Christer W.	The Journal of Research on the Lepidoptera	10.5962/p.266504
Deflective effect and the effect of prey detectability on anti-predator function of eyespot	2011	Vallin, A. and Dimitrova, M., Kodandaramaiah, U. and Merilaita, S.	Behavioral Ecology and Sociobiology	10.1007/s00265-011-1173-7
Number of eyespot and their intimidating effect on naïve	2011	Merilaita, S., Vallin, A., Kodandaramaiah, U., Dimitrova, M.,	Behavioral Ecology	10.1093/beheco/arm135

predators in the peacock butterfly		Ruuskanen, S. and Laaksonen, T.		
The 'sparkle' in fake eyes - the protective effect of mimic eyespots in lepidoptera	2012	Blut, C., Wilbrandt, J., Fels, D., Girgel, E.I. and Lunau, K.	Entomologia Experimentalis et Applicata	10.1111/j.1570-7458.2012.01260.x
Eyespots interact with body colour to protect caterpillar-like prey from avian predators	2012	Hossie, T.J. and Sherratt, T.N.	Animal Behaviour	10.1016/j.anbehav.2012.04.027
Anti-predator adaptations and strategies in the Lepidoptera	2012	de Wert, L.	Doctoral thesis	none
Bird attacks on a butterfly with marginal eyespots and the role of prey concealment against the background	2013	Olofsson, M., Jakobsson, S. and Wiklund, C.	Biological Journal of the Linnean Society	10.1111/bij.12063
Defensive posture and eyespots deter avian predators from attacking caterpillar models	2013	Hossie, TJ and Sherratt, TN	Animal Behaviour	10.1016/j.anbehav.2013.05.029
Revealed by conspicuousness: distractive markings reduce camouflage	2013	Stevens, M., Marshall, KLA, Troscianko, J., Finlay, S., Burnand, D. and Chadwick, SL.	Behavioral Ecology	10.1093/beheco/ars156
Eyespot display in the peacock butterfly triggers antipredator behaviors in naïve adult fowl	2013	Olofsson, M., Lovlie, H., Tibblin, J., Jakobsson, S. and Wiklund, C.	Behavioral Ecology	10.1093/beheco/ars167
The position of eyespots and thickened segments influence their protective value to caterpillars	2014	Skelhorn, J., Dorrington, G., Hossie, TJ. and Sherratt, TN.	Behavioral Ecology	10.1093/beheco/aru154
Predator mimicry, not conspicuousness, explains the efficacy of butterfly eyespots	2015	De Bona, S., Valkonen, JK., López-Sepulcre, A. and Mappes, J.	Proceedings of the Royal Society B: Biological Sciences	10.1098/rspb.2015.0202
Body size affects the evolution of eyespots in caterpillars	2015	Hossie, TJ., Skelhorn, J., Breinholt, JW.,	Proceedings of the National Academy	10.1073/pnas.1415121112

		Kawahara, AY. and Sherratt, TN.	of Sciences of the United States of America	
What makes eyespots intimidating- the importance of pairedness Evolutionary ecology and behaviour	2015	Mukherjee, R. and Kodandaramaiah, U.	BMC Evolutionary Biology	10.1186/s12862-015-0307-3
On the deterring effect of a butterfly's eyespot in juvenile and sub-adult chickens	2015	Olofsson, M., Wiklund, C. and Favati, A	Current Zoology	10.1093/czoolo/61.4.749
Multicomponent deceptive signals reduce the speed at which predators learn that prey are profitable	2016	Skelhorn, J., Holmes, GG., Hossie, T.J. and Sherratt, TN.	Behavioral Ecology	10.1093/beheco/arv135
Attack risk for butterflies changes with eyespot number and size	2016	Ho, S., Schachat, SR., Piel, WH. and Monteiro, A.	Royal Society Open Science	10.1098/rsos.150614
The effectiveness of eyespots and masquerade in protecting artificial prey across ontogenetic and seasonal shifts	2022	Postema, EG.	Current Zoology	10.1093/cz/zoab082

912

913 (b) excluded studies

title	year	authors	journal	doi	reason
The effects of a tranquilliser on the reactions of domestic chicks to an aversive eye-like shape	1979	Jones, RB.	IRCS Medical Science	none	No full-text
Young domestic chicks avoid eye-like shapes	1980	JONES, RB	Applied Animal Ethology	10.1016/0304-3762(80)90037-1	No full-text
The startle responses of blue jays to Catocala	1985	Schlenoff, DH.	Animal Behaviour	10.1016/S0003-3472(85)80164-0	Wrong outcome

<i>(Lepidoptera: Noctuidae)</i> prey models					
Fearful symmetry: Pattern size and asymmetry affects aposematic signal efficacy	1999	Forsman, A. and Merilaita, S.	Evolutionary Ecology	10.1023/A:1006630911975	Invalid comparator
"An eye for an eye?" - On the generality of the intimidating quality of eyespots in a butterfly and a hawkmoth	2007	Vallin, A., Jakobsson, S. and Wiklund, C.	Behavioral Ecology and Sociobiology	10.1007/s00265-007-0374-6	Invalid comparator
Coincident disruptive coloration	2009	Cuthill, IC and Székely, A	Philosophical Transactions of the Royal Society B-Biological Science	10.1098/rstb.2008.0266	Invalid comparator
Marginal eyespots on butterfly wings deflect bird attacks under low light intensities with UV wavelengths	2010	Olofsson, M., Vallin, A., Jakobsson, S. and Wiklund, C.	PLoS ONE	10.1371/journal.pone.0010798	Invalid comparator
Insect coloration as a defence mechanism against visually hunting predators	2011	Lyytinen, A.	Doctoral thesis	none	Published thesis
Effects of lepidopteran eyespot components on the deterrence of predatory birds	2015	Blut, C. and Lunau, K.	Behaviour	10.1163/1568539X-00003288	Invalid comparator
Antipredator behavior by a nesting hummingbird in response to a caterpillar with eyespots	2019	Marden, JH. and Pérez Carrillo, JF.	Ecology	10.1002/ecy.2582	Observational study

The Influence of the eyespots of peacock butterfly (<i>Aglais io</i>) and caterpillar on predator recognition	2020	Park, J. and Heo D	Open Science Journal	10.23954/osj.v5i2.2455	Invalid comparator
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915 **Table S4.** Summary of a multi-moderator model including all moderators. The bold typeface
 916 is used when a 95% confidence interval (CI) does not contain zero; thus, it can be interpreted
 917 as an existing significant effect in predator avoidance.

	Estimate	95%CI
intercept	-0.06	(-0.50, 0.34)
Treatment stimulus	-0.02	(-0.19, 0.23)
Log-transformed area	0.09	(0.009, 0.18)
Number pattern	-0.05	(-0.11, 0.004)
Material type of prey: real	0.18	(-0.09, 0.45)

918

919 **Table S5.** Average maximum diameter of Eyespots on *Bicyclus anynana*. AM obtained the
 920 pictures from lepdata.org/photos/animals/ and <https://data.nhm.ac.uk/> and measured the
 921 eyespot diameters. Raw data is <https://ayumi-495.github.io/eyespot/>.

Median	Range
3.41	1.82 – 5.04

922