Avoidance of an aposematically coloured butterfly by wild birds in a tropical forest

DENISE D. DELL’AGLIO, 1,2 MARTIN STEVENS 3 and CHRIS D. JIGGINS 1,2 1Butterfly Genetics Group, Department of Zoology, University of Cambridge, Cambridge, U.K., 2Smithsonian Tropical Research Institute, Panama City, Panama and 3Centre for Ecology & Conservation, College of Life & Environmental Sciences, University of Exeter, Penryn, U.K.

Abstract. 1. Birds are considered to be the primary selective agents for warning colouration in butterflies, and select for aposematic mimicry by learning to avoid brightly coloured prey after unpleasant experiences. It has long been thought that bright colouration plays an important role in promoting the avoidance of distasteful prey by birds.

2. The hypothesis that warning colouration facilitates memorability and promotes predator avoidance was tested by means of a field experiment using distasteful model butterflies. Artificial butterflies with a Heliconius colour pattern unknown to local birds were generated using bird vision models, either coloured or achromatic, and hung in tree branches in a tropical forest. Two sequential trials were conducted at each site to test avoidance by naïve and experienced predators.

3. There was a significant reduction in predation in the second trial. Also, coloured models were attacked less than achromatic models. Specifically, coloured butterflies were attacked significantly less in the second trial, but there was no significant decrease in predation on achromatic models.

4. The present results imply an important role for colour in enhancing aversion of aposematic butterflies. It has also been demonstrated that previous experience of distasteful prey can lead to enhanced avoidance in subsequent trials, supporting mimicry theory.

Key words. Aposematism, artificial models, Heliconius, memory, predation.

Introduction

The aposematic signals of unpalatable prey are a defence against visually hunting predators. In particular, conspicuous colouration is strongly favoured in defended prey as it can increase detection efficiency and lead to rapid decision-making (Endler, 1988). Colours such as red, yellow, and orange are normally highly contrasting with the background and are commonly used to advertise unpalatability (Stevens & Ruxton, 2012; Arenas et al., 2014). Therefore, these brightly coloured signals support rapid discrimination from cryptic prey and have long been considered to facilitate avoidance learning when compared with less visible colouration (Guilford & Dawkins, 1991; Speed, 2000).

Birds are widely considered to be the primary selective agent for the aposematic colouration of butterflies. After unpleasant experiences with an unpalatable prey, bird predators learn to avoid similar morphs (Ham et al., 2006; Lindström et al., 2006). This learning ability leads to selection favouring the most abundant colour patterns in a local area and generates aposematism and Müllerian mimicry in which predator attacks are reduced through aversion learning of locally common aposematic patterns (Müller, 1879; Mallet & Joron, 1999).

Learning and forgetting are essential for the maintenance of Müllerian mimicry (Speed & Turner, 1999). Memory is linked to recognition, and if predators forget about experiences with prey, then recognition of an aposematic signal is not possible (Speed, 2000). Warning signals should, therefore, be selected to be memorable, to provoke low rates of forgetting and enhance predator aversion (Servedio, 2000; Speed, 2000). Among mimetic butterflies, long-term memorability of learned avoidance of the model is vital for the protection of the co-mimic. There is a large body of evidence supporting the role of colour in avoidance learning and memory, but this primarily

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comes from cage experiments (Sillén-Tullberg, 1981; Osorio et al., 1999; Ham et al., 2006; Sandre et al., 2010).

Experiments in the wild with natural predators can better estimate the overall response of a local population and can complement cage studies. Responses from captive birds might be influenced by their appetite (Sandre et al., 2010), food deprivation, and artificial environments with constrained viewing, whereas natural environments are heterogeneous and offer a wider variety of alternative food, which might alter decision-making strategies. For example, a study with natural bird populations using artificial models of the wood tiger moth [Parasemia planctagina (Linnaeus)] suggested that spatial heterogeneity in a predator community creates a mosaic of selection facilitating polymorphism (Nokelainen et al., 2013). Also, another study with wild birds showed that achromatic (non-coloured) Heliconius models were attacked significantly more than coloured models of a local pattern, demonstrating the importance of aposematic signals in avoiding predation (Finkbeiner et al., 2014). Furthermore, an experiment with model poison frogs [Dendrobates tinctorius (Schneider)] showed varying attack rates of wild tropical predators in different light conditions (Rojas et al., 2014). Still, few studies, to date, have explored attack rates on different coloured models using wild birds and under natural conditions.

Neotropical Heliconius butterflies are one of the best-studied mimicry systems (Mallet & Joron, 1999), in which unpalatable sympatric species form mimicry rings. Many Heliconius species are highly variable in colouration and patterns (Mallet & Gilbert, 1995). Several studies have investigated predator behaviour towards Heliconius butterflies in cages using wild-caught rufous-tailed jacamars (Galbula ruficauda Cuvier), which are specialist predators of fast-flying insects and exhibit specific butterfly handling strategies. Jacamars readily reject Heliconius by sight or by taste and discriminate them from other butterfly species (Chai, 1986; Langham, 2004). Field experiments, using other butterfly predators, kingbirds, and flycatchers, also showed taste-rejection of Heliconius butterflies (Pinheiro, 2003, 2011). Previous field studies have demonstrated mimicry selection by releasing live butterflies (Benson, 1972; Mallet & Barton, 1989) and monitoring recapture rates.

Therefore, to better understand the dynamics of Heliconius mimicry, more information from the predators’ perspective in the wild is required. Here we investigate the role of colouration in attack rates, testing the ability of bird predators to avoid an unpalatable Heliconius warning signal in a tropical forest. The assumption is that wild birds would have a bias against aposematic colouration, which would facilitate the memory of novel butterfly colour pattern. We performed a field test of the hypothesis that aposematism facilitates avoidance of novel distasteful prey using artificial distasteful butterflies with a colour pattern unknown to local bird predators.

**Material and methods**

**Production of artificial butterflies**

Artificial butterflies were produced based on wings of Heliconius erato lativitta (Linnaeus) which is found only in the Amazon basin, not in Panama (Brown, 1979; Hines et al., 2011). We calibrated the appearance of the artificial wings to account for bird colour and luminance vision. Photographs of real wings and a printer colour palette were taken with a Fuji-calibrated UV SLR camera (Fujifilm, Düsseldorf, Germany) with an ultraviolet (UV)-transmitting quartz lens (Jenoptic) with a UV pass filter (transmitting between 300 and 400 nm; Baader U filter, David Hinds Ltd., Bedfordshire, U.K.) and a UV/IR-Cut pass filter (blocking UV below 400 nm and IR above 700 nm; Baader UV/IR Cut Filter, David Hinds Ltd.), representing the UV and human visible spectrum, respectively. After this, predicted photon catch values of four single cones (used in colour vision) and double cones (probably used in achromatic vision) were calculated, based on the sensitivity of a UV vision bird receptors, Blue tit [Cyanistes caeruleus (Linnaeus)] (Hart et al., 2000; Endler & Mielke, 2005), according to the methodology created by Troscianko and Stevens (2015). Our criteria for selecting appropriate colours were that the ‘just-noticeable-differences’ (JND) values (Vorobyev & Osorio, 1998) of the printer colours against real butterfly colours (Finkbeiner et al., 2012; Merrill et al., 2012) should be as close as possible to the threshold of discrimination of three JND (Siddiqi et al., 2004) (Table S1 in Supporting information). For achromatic models, only achromatic contrast was used. Colours were closely reproduced as demonstrated in avian colour space vision (Figure S1 in Supporting information). Afterwards, two types of artificial butterflies were designed, coloured, and achromatic (Fig. 1). These were printed on Whatman filter paper (GE Healthcare Life Sciences, Buckinghamshire, U.K.), which produces reflectance spectra close in brightness to actual wings (Finkbeiner et al., 2012), using an HP Colour Laser Jet 4700dn printer (HP Inc., London, U.K.). A 3-hydroxy-DL-kynurenine (3-OHK, Sigma-Aldrich Company Ltd., Dorset, U.K.) pigment was applied to the yellow bands of the foregoing to provide accurate UV reflectance (Finkbeiner et al., 2012).

The artificial wings were attached with a nylon line to an edible pastry body (flour, lard, water, and black food dye). To provide an unpleasant taste, quinine monohydrochloride dihydrate (4% solution) was sprayed on the body and wings of both model types. This concentration is aversive and has a similar effect to sampling a toxic prey (Rowe & Skelhorn, 2005). Finally, Krylon matte finishing spray was applied lightly to coat the artificial butterflies with a waterproofing element 24 h before placing the models out, without altering the colour or smell.

**Field experiment**

The trials were conducted along three forest trails in Parque Nacional Soberanía, Panama. Models were hung by nylon line (10 cm long) on tree branches (1.7 m high) to swing freely similar to a live butterfly. We aimed to maximise attack rates by butterfly predators that catch insects during flight and detect movement. Models were hung every 10 m in pairs, one coloured and one achromatic on opposite sides of the trail, with the assignment, randomised.

To test memorability, the experiment had two trials. In the first trial, 152 models of each type were placed for 4 days,
followed by a second identical trial started 5 days after the first trial finished. In the second trial, the same procedure was repeated at the same location with new 152 new models of each type. The models were checked for attack marks after 48 and 96 h. An artificial model was considered attacked if the body or wings included clearly visible beak marks, or part or all of the body was missing. If a model had more than one beak mark on it, this was counted as a single attack. Evidence of attack by animals other than birds, notably insects such as ants, was readily distinguished and was not counted as an attack (Salazar Carrión et al., 2014).

Statistical analyses

We used the binomial response of attack (presence or absence) of two treatments (chromatic and achromatic) in two trials (1 and 2) across three localities. To test the homogeneity of variance between localities, a Bartlett test and Fligner–Killeen test were used. We used generalised linear mixed models (GLMM) with a binomial distribution, to test for the effect of trial, treatment, and locality (as a random factor), as well as their interaction terms, on predation. Tests used the R packages stats and lme4 in R statistical software (Bates et al., 2015; R Core Team, 2015).

Results

In total, 608 artificial butterflies were placed in the wild (152 chromatic and 152 achromatic on trial 1 and 152 chromatic and 152 achromatic on trial 2). The use of a nylon line allowed us to recover fully the models, 117 (19%) of which were attacked. Tests of homogeneity revealed no evidence that the three localities differed in predation events (Bartlett test: $\chi^2 = 0.85$, d.f. = 2, $P = 0.651$, Fligner–Killeen test: $\chi^2 = 2.71$, d.f. = 2, $P = 0.257$). The ‘locality’ term did not explain much variation in our model ($\hat{\sigma}^2 = 0.033$, SD = 0.18). There were clear differences in the number of predation events in the models between the two trials (Fig. 2). We observed no difference in predation of the achromatic butterfly between the two trials (37 on trial 1 and 31 on trial 2). A greater proportion of attacks occurred during the first trial (69 on trial 1 and 48 on trial 2, trial: $z_{604,608} = -2.35$, $P = 0.018$). Also, aposematic colour models were attacked less overall (colour: $z_{604,608} = -2.15$, $P = 0.031$). This was mainly as a result of a reduction in attacks in the second trial (32 on trial 1 and 17 on trial 2), but also compared with the achromatic pattern of the second trial (31 achromatic and 17 chromatic). However, although the GLMM showed a significant effect of both trial and colour alone, the interaction between trial and colour was not significant (trial*colour: $z_{604,608} = -1.06$, $P = 0.28$).

Discussion

We evaluated the influence of aposematic colouration on attack rate by bird predators in a tropical forest. We observed a reduction in attack rates on coloured models as compared to achromatic models, demonstrating a role for colour in enhancing the avoidance of a novel distasteful prey. Many previous experiments have demonstrated the protective value of Heliconius warning colour patterns alone (Benson, 1972; Chai, 1986; Mallet & Barton, 1989; Kapan, 2001; Langham, 2004; Merrill et al., 2012; Finkbeiner et al., 2014), including one study which compared chromatic and achromatic prey (Finkbeiner et al., 2014). Our results, therefore, support previous work showing that bright colours enhance the avoidance of aposematic prey, and contributes to an explanation of why aposematic insects in general and Heliconius in particular, often evolve bright colouration.

There was a significantly reduced attack rate in the second trial, suggesting that the bad experience of the distasteful model in the first trial may have induced later aversion. Prey palatability is known to influence predator learning and memory of warning colours (Lindström et al., 2006; Skelhorn & Rowe, 2006; Svádová et al., 2009). Having both warning colouration...
and distastefulness can change predator decision-making and increase avoidance (Servedio, 2000). However, the short period between trials means that we cannot distinguish between true ‘memory’ and a short-term aversion reaction to explain these results. It would be interesting to repeat similar experiments over different periods of time to test for long-term memory. Predation field studies in tropical forests are challenging, and it was not possible to identify predators to demonstrate that the same individual that had a bad experience later avoided the same prey type, so there may be other ecological explanations for our results. Nonetheless, whatever the cause, our experiment supports the prediction of mimicry theory that attack rates on aposematic prey should decline with predator experience.

The avoidance of aposematic patterns is often considered not only as a learned trait but also as an innate response to conspicuous colours, whereby predators are unwilling to eat prey with a novel appearance (Marples et al., 1998; Lee et al., 2010). In addition, a study comparing predation rates on aposematic and cryptic prey, also in field conditions, showed that aposematic prey were completely consumed less often than cryptic prey but partially consumed more often suggesting ‘go-slow’ predation, in which predators are more cautious with aposematic prey (Carroll & Sherratt, 2013). However, there was no strong support for this in our data, with the two novel patterns equally attacked in the first trial of the experiment. Similar results were found for another Heliconius predation experiment in which the ‘nonlocal’ phenotype had higher attack rates (Finkbeiner et al., 2014). Different predators are likely to have different aversion responses to colour, and so the heterogeneity of predators in the wild might explain this result (Endler, 1988; Servedio, 2000; Speed et al., 2000; Endler & Mappes, 2004).

The least attacked prey were the coloured models in the second trial. This suggests that chromatic prey would have triggered a stronger aversion response than the achromatic prey, implying a role for colour in reducing attack rates. However, a test for the interaction between trial and pattern was not significant, so we cannot definitively conclude that colour influenced the reduced response in the second trial, although this seems likely. A power analysis suggested that we would need to quadruple the size of our experiment approximately to detect a significant interaction between colour and trial. The results are nonetheless consistent with the idea that colour enhances learning of aversion (Speed, 2000).

Predator psychology models assume that the rate of predation is dependent on learning and forgetting rates, and the absence of reinforcing experiences might lead to forgetfulness (Turner & Speed, 1996; Speed & Turner, 1999; Servedio, 2000; Speed, 2000). For instance, Jacamars have been shown to forget novel colour morphs after an interval of 2 years (Langham, 2004), which might have been as a result of a lack of reinforcing encounters with the artificial prey. Our artificial butterflies were in the sight of predators for 4 days during the trials, which may have led them to be seen several times and which could have stimulated memory. Occasional sampling in nature also might reinforce memory provided that butterflies can be rejected by sight or by taste, which is a common behaviour among butterfly predators (Chai, 1986; Pinheiro, 2003). Further experiments would be needed to determine whether distasteful models or repeated exposure could trigger long-term memory and faster learning rates.

In this experiment, there were no detectable effects of the pattern itself as a warning signal, as the distasteful achromatic pattern was equally attacked in both trials. Previous experiments with chicks indicate that colour differences are more memorable than luminance contrast, whereas pattern attracts attention (Osorio et al., 1999). Nonetheless, previous studies have shown avoidance learning using different patterns (Rowe et al., 2004; Aronsson & Gamberale-Stille, 2008; Rowland et al., 2010), and benefits of pattern mimicry may emerge at a later stage in the learning process (Rowe et al., 2004). Given the precise mimicry is seen in Heliconius, both pattern and colour seem to be vitally important for predator avoidance (Finkbeiner et al., 2014).

The attack frequency of this study was significantly higher than in previous work using artificial Heliconius patterns (Merrill et al., 2012; Finkbeiner et al., 2014; Salazar Carrión et al., 2014). This may be partly because the models represented a novel morph that birds had not experienced before. However, our methodology using suspended butterflies that could move in the wind might also have attracted more predators. This method may, therefore, be useful for future experiments studying selection on butterfly models.

This experiment indicates that attack rates on novel aposematic butterflies are reduced over time, consistent with experiments on caged birds showing learning of warning colours. Furthermore, we have also shown a role for colour in enhancing aversion towards aposematic prey. This experiment has shown avoidance of an aposematic butterfly in a tropical forest and contributes to a better understanding of the dynamics of Heliconius aposematic mimicry in the wild.

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**Supporting Information**

Additional Supporting Information may be found in the online version of this article under the DOI reference: 10.1111/een.12335

**Table S1.** Chromatic and achromatic contrast (JND) between the real wing and printed wing perceived by Blue tit (Cyanistes caeruleus) vision. Values >3 JND denote an increasing ability of discrimination, whereas values ≤3 JND denote colours generally indistinguishable from each other. Notice that yellow colour could be closely reproduced in the models and the orange was close but not possible to reproduce accurately.
**Figure S1.** Distribution of colours perceived by Blue tit (Cyanistes caeruleus) vision in a tetrahedral colour space. Each point is determined by the relative stimulation of the four cone colour channels and each axis represents a channel: ultraviolet (UV), short (SW), medium (MW), and long (LW) wavelength sensitive cones.

**References**

Arenas, L.M., Troschiano, J. & Stevens, M. (2014) Color contrast and stability as key elements for effective warning signals. *Frontiers in Ecology and Evolution*, 2, 1–12.

Aronsson, M. & Gamberale-Stille, G. (2008) Domestic chicks primarily attend to colour, not pattern, when learning an aposematic coloration. *Animal Behaviour*, 75, 417–423.

Bates, D., Mächler, M., Bolker, B. & Walker, S. (2015) Fitting linear mixed-effects models using lme4. *Journal of Statistical Software*, 67, 1–48.

Benson, W.W. (1972) Natural selection for Mullerian mimicry in Heliconius erato in Costa Rica. *Science*, 176, 936–939.

Brown, K. (1970) *Ecologia geográfica e evolução nas florestas neotrópicais*. Universidade Estadual de Campinas, Campinas, Brazil.

Carroll, J. & Sherratt, T.N. (2013) A direct comparison of the effectiveness of two anti-predator strategies under field conditions. *Journal of Zoology*, 291, 279–285.

Chai, P. (1986) Field observations and feeding experiments on the responses of rufous-tailed jacamars butterflies in a tropical rainforest. *Biological Journal of the Linnean Society*, 29, 161–189.

Endler, J.A. (1988) Frequency-dependent predation, crypsis and aposematic coloration. *Philosophical Transactions of the Royal Society, B: Biological Sciences*, 319, 505–523.

Endler, J.A. & Mappes, J. (2004) Predator mixes and the conspicuousness of aposematic signals. *The American Naturalist*, 163, 532–547.

Endler, J.A. & Mielke, P.W. (2005) Comparing entire colour patterns as birds see them. *Biological Journal of the Linnean Society*, 86, 405–431.

Finkbeiner, S.D., Briscoe, A.D. & Reed, R.D. (2012) The benefit of being a social butterfly: communal roosting deters predation. *Proceedings of the Royal Society of London. Series B*, 279, 2769–2776.

Finkbeiner, S.D., Briscoe, A.D. & Reed, R.D. (2014) Warning signals are seductive: relative contributions of color and pattern to predator avoidance and mate attraction in Heliconius butterflies. *Evolution*, 68, 3410–3420.

Guilford, T. & Dawkins, M.S. (1991) Receiver psychology and the evolution of animal signals. *Animal Behaviour*, 42, 1–14.

Ham, A.D., Ihalainen, E., Lindström, L. & Mappes, J. (2004) The importance of taste and visual appearance for predator education in Mullerian mimicry. *Frontiers in Ecology and Systematics*, 30, 201–233.

Mallet, J. & Barton, N.H. (1989) Strong natural selection in a warning-color hybrid zone. *Evolution*, 43, 421–431.

Mallet, J. & Gilbert, L.E. (1995) Why are there so many mimicry rings? Correlations between habitat, behaviour and mimicry in Heliconius butterflies. *Biological Journal of the Linnean Society*, 55, 159–180.

Mallet, J. & Joron, M. (1999) Evolution of diversity in warning color and mimicry: polymorphisms, shifting balance, and speciation. *Annual Review of Ecology and Systematics*, 30, 201–233.

Marpes, N.M., Roper, T.J. & Harper, D.G.C. (1998) Responses of wild birds to novel prey: evidence of dietary conservatism. *Oikos*, 83, 161–165.

Merrill, R.M., Wallbank, R.W.R., Bull, V., Salazar, P.C.A., Mallet, J., Stevens, M. et al. (2012) Disruptive ecological selection on a mating cue. *Proceedings of the Royal Society of London. Series B: Biological Sciences*, 279, 4907–4913.

Müller, F. (1879) Iuna and Thyridia; a caseful mimicry of butterflies. *Transactions of the Entomological Society of London*, 1879, xx–xxix.

Nokelainen, O., Valkonen, J., Lindstedt, C. & Mappes, J. (2013) Changes in predator community structure shifts the efficacy of two warning signals in Arctiid moths. *The Journal of Animal Ecology*, 83, 598–605.

Osorio, D., Jones, C.D. & Vorobyev, M. (1999) Accurate memory for colour but not pattern contrast in chicks. *Current Biology*, 9, 199–202.

Pinheiro, C.E.G. (2013) Does Mullerian mimicry work in nature? Experiments with butterflies and birds (Tyrannidae). *Biotropica*, 35, 356–364.

Pinheiro, C.E.G. (2011) On the evolution of warning colouration, Batesian and Mullerian mimicry in Neotropical butterflies: the role of jacamars (Galbulidae) and tyrant-flycatchers (Tyrannidae). *Journal of Avian Biology*, 42, 277–281.

R Core Team (2015) *R: A Language and Environment for Statistical Computing*. R Foundation for Statistical Computing, Vienna, Austria.

Rojas, B., Rautiala, P. & Mappes, J. (2014) Differential detectability of polymorphic warning signals under varying light environments. *Behavioural Processes*, 109, 164–172.

Rowe, C. & Skelhorn, J. (2005) Colour biases are a question of taste. *Animal Behaviour*, 69, 587–594.

Rowe, C., Lindström, L. & Lyytinen, A. (2004) The importance of pattern similarity between Mullerian mimics in predator avoidance learning. *Proceedings of the Royal Society of London. Series B: Biological Sciences*, 271, 407–413.

Rowland, H.M., Hoogesteger, T., Ruston, G.D., Speed, M.P. & Mappes, J. (2010) A tale of 2 signals: signal mimicry between aposematic species enhances predator avoidance learning. *Behavioural Ecology*, 21, 851–860.

Salazar Carrión, P.A., Stevens, M., Jones, R.T., Ogilvie, I. & Jiggins, C. (2014) A field test for frequency-dependent selection on mimic and colour patterns in Heliconius butterflies. *bioRxiv*, doi:10.1101/005249.

Sandre, S., Stevens, M. & Mappes, J. (2010) The effect of predator appetite, prey warning coloration and luminance on predator foraging decisions. *Behaviour*, 147, 1121–1143.

Servedio, M.R. (2000) The effects of predator learning, forgetting, and recognition errors on the evolution of warning coloration. *Evolution*, 54, 751–763.

Siddiqui, A., Cronin, T.W., Loew, E.R., Vorobyev, M. & Summers, K. (2004) Interspecific and intraspecific views of color signals in...
the strawberry poison frog Dendrobates pumilio. *The Journal of Experimental Biology*, 207, 2471–2485.

Sillén-Tullberg, B. (1981) The significance of coloration per se, independent of background, for predator avoidance of aposematic prey. *Animal Behaviour*, 33, 1382–1384.

Skelhorn, J. & Rowe, C. (2006) Prey palatability influences predator learning and memory. *Animal Behaviour*, 71, 1111–1118.

Speed, M.P. (2000) Warning signals, receiver psychology and predator memory. *Animal Behaviour*, 60, 269–278.

Speed, M.P. & Turner, J.R.G. (1999) Learning and memory in mimicry: II. Do we understand the mimicry spectrum? *Biological Journal of the Linnean Society*, 67, 281–312.

Speed, M.P., Alderson, N.J., Hardman, C. & Ruxton, G.D. (2000) Testing Mullerian mimicry: an experiment with wild birds. *Proceedings of the Royal Society of London. Series B*, 267, 725–731.

Stevens, M. & Ruxton, G.D. (2012) Linking the evolution and form of warning coloration in nature. *Proceedings of the Royal Society of London. Series B*, 279, 417–426.

Svádová, K., Exnerová, A., Štys, P., Landová, E., Valenta, J., Fučíková, A. *et al.* (2009) Role of different colours of aposematic insects in learning, memory and generalization of naïve bird predators. *Animal Behaviour*, 77, 327–336.

Troscianko, J. & Stevens, M. (2015) Image calibration and analysis toolbox – a free software suite for objectively measuring reflectance, colour and pattern. *Methods in Ecology and Evolution*, 6, 1320–1331.

Turner, J.R.G. & Speed, M.P. (1996) Learning and memory in mimicry. I. Simulations of laboratory experiments. *Philosophical Transactions of the Royal Society of London, Series B*, 351, 1157–1170.

Vorobyev, M. & Osorio, D. (1998) Receptor noise as a determinant of colour thresholds. *Proceedings of the Royal Society of London. Series B*, 265, 351–358.

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