DOI: 10.1111/2041-210X.13073

RESEARCH ARTICLE

Methods in Ecology and Evolution

Boundary strength analysis: Combining colour pattern geometry and coloured patch visual properties for use in predicting behaviour and fitness

John A. Endler | **Gemma L. Col[e](http://orcid.org/0000-0002-3365-3580)** | **Alexandrea M. Kran[z](http://orcid.org/0000-0002-7980-1374)**

Centre for Integrative Ecology, School of Life & Environmental Sciences, Deakin University, Waurn Ponds, Victoria, Australia

Correspondence John A. Endler Email: john.endler@deakin.edu.au

Funding information Australian Research Council, Grant/Award Number: DP110101421 and DP150102817

Handling Editor: Simon Blomberg

Abstract

- 1. Colour patterns are used by many species to make decisions that ultimately affect their Darwinian fitness. Colour patterns consist of a mosaic of patches that differ in geometry and visual properties. Although traditionally pattern geometry and colour patch visual properties are analysed separately, these components are likely to work together as a functional unit. Despite this, the combined effect of patch visual properties, patch geometry, and the effects of the patch boundaries on animal visual systems, behaviour and fitness are relatively unexplored.
- 2. Here, we describe boundary strength analysis (BSA), a novel way to combine the geometry of the edges (boundaries among the patch classes) with the receptor noise estimate (Δ*S*) of the intensity of the edges. The method is based upon known properties of vertebrate and invertebrate retinas. The mean and *SD* of Δ*S* (m_0 _{*S*}, s_0) of a colour pattern can be obtained by weighting each edge class Δ*S* by its length, separately for chromatic and achromatic Δ*S*. This assumes those colour patterns, or parts of the patterns used in signalling, with larger $m_{\text{A}s}$ and $s_{\text{A}s}$, are more stimulating and hence more salient to the viewers. BSA can be used to examine both colour patterns and visual backgrounds.
- 3. Boundary strength analysis was successful in assessing the estimated conspicuousness of colour pattern variants in two species, guppies *Poecilia reticulata* and Gouldian finches *Erythrura gouldiae*, both polymorphic for patch colour, luminance and geometry. The 3D representations of the Δ*S* of patch edges (Fort Diagrams) of both species show that there is little or negative geometric correspondence between the chromatic and achromatic edges. All individuals have $m_{\Lambda S}$ > 1.5 for both chromatic and achromatic measures, indicating the high within-pattern contrast expected for display signals. In contrast from what one would expect from sexual selection, all guppies have $m_{\Delta S}$ less than expected from random contacts between all pairs of patch colour/luminance classes. The correlation between chromatic and luminance Δ*S* is negative in both species but zero when correlating all possible kinds of edges between the colours of each species and morph, indicating nonrandom colour geometry.
- 4. The pattern difference between chromatic and achromatic edges in both species reveals the possibility that chromatic and achromatic edges could function differently. The smaller than random expected $m_{\Delta S}$ values in guppies suggests an

anti-predator function because guppies are never found without predators. Moreover, $m_{\Lambda S}$ could vary with predation intensity within and among species. BSA can be applied to any colour pattern used in intraspecific and interspecific behaviour. Seven predictions and four questions about colour patterns are presented.

5. In species which are very convex in cross-section, both chromatic and luminance m_{AS} change with viewing angle; geometry of signalling is as important as signal geometry.

KEYWORDS

boundary strength analysis, colour pattern analysis, colour transitions, conspicuousness, fort diagrams, pattern edges, receptor noise, visual signals

1 | **INTRODUCTION**

Colour patterns are important in survival and reproduction in diverse species because they affect mating success, contests, avoiding predators, luring prey or attracting pollinators. In general, the fitness of the sender (individual with the colour pattern) is affected because the receiver (viewer of the colour pattern) can make a behavioural or physiological decision about the sender, based upon reception and perception of the sender's colour pattern (e.g., receivers will mate, fight, attack, be lured close and eaten, pollinate or disperse seeds). Colour patterns offer an effective way of investigating the complex relationship between genes, morphology, performance, fitness and evolution (Arnold, 1983, 2003) because the functions of most colour patterns are relatively easy to identify (Endler, 1978, 1980). However, the links between visual properties, perception, receivers' decision-making processes and fitness are not well understood.

Decisions made by the receiver depend upon both the signal design of the colour pattern (the physical structure of the signal) and its signal content (information about the signaller, reviewed in Endler, 1993a). For both components, the first-stage affecting fitness is the stimulation of the receiver's retina by the colour pattern; all subsequent processes leading to perception and decision-making flow through this step (Lythgoe, 1979). Although all components of a colour pattern may affect the viewer's decision making, their relative importance in retinal and brain stimulation is not known. In particular, we do not know how colour, luminance, patch size and patch geometry work together to affect receiver behaviour, and so cannot yet make explicit predictions about colour pattern properties or the behavioural decisions based upon them.

Relating patterns to fitness has been successful for some species with cryptic colour pattern components (Troscianko, Skelhorn, & Stevens, 2017; Troscianko, Wilson-Aggarwal, Stevens, & Spottiswoode, 2016), but there is a tendency in the literature to study only pattern or one or two colour pattern components. Previous attempts to quantify colour patterns have included mapping the pattern components (Van Belleghem et al., 2018), mapping pattern component boundaries (Stevens & Cuthill, 2006) and estimating the distributions of relative pattern component edge lengths (Endler, 2012). Other analyses have calculated colour patch discriminability

(Siddiqi, Cronin, Loew, Vorobyev, & Summers, 2004). However, all of these methods ignore whether or not the colour patches share common boundaries. Colour patch boundaries are important because adjacent colour patches will influence the visual perception of a given patch as well as the contrast across the boundary.

Here, we present boundary strength analysis (BSA), a way to combine the effects of both patch properties and the intensity of patch edges (transitions between patches) based upon how they are processed by the visual system in the retina. BSA estimates the effects of both colour and patch edges by combining two existing methods for the first time, one for discriminability between adjacent patches (Δ*S*, Vorobyev & Osorio, 1998) and one for the geometric arrangement of patches (Endler, 2012). Unlike all previous methods, BSA includes the estimated visual intensity of the boundaries (estimated by Δ*S*) and their length, rather than just recording which boundaries are present, and calculates Δ*S* statistics only between patches which come in contact. This is consistent with the opponent visual processes that detect colour and colour patch edges, and the fact that these processes sample small parts of the visual field (Dowling, 2012; Kelber, 2016). This allows us to begin to examine colour patterns less arbitrarily, by incorporating estimates of how strongly patch boundaries stimulate the retina as a proxy for conspicuousness.

Boundary strength analysis can be used for animal and plant colour patterns as well as visual backgrounds, and allows investigation of both within pattern and pattern-background contrast. For brevity, we will describe and give examples of BSA in terms of within-pattern contrast but the resulting statistics can be calculated for visual backgrounds as well as patterns and the two compared to estimate pattern-background contrast.

1.1 | **Visual modelling of colour discrimination**

We use the receptor noise model or RN (Vorobyev & Osorio, 1998; reviewed in Kelber, Vorobyev, & Osorio, 2003) to estimate detection thresholds for colour discrimination. The input to the model consists of the relative light (photon) captures for each photoreceptor class in the viewer's retina for two colour patches. The output of the RN model is Δ*S*, which is similar to a multivariate equivalent to *t* in statistics in that it compares the difference between the two sets of cone captures to the standard error of the difference; it is the Mahalanobis distance of multivariate statistics (Clark, Santer, & Brebner, 2017). Like other signal/noise measures Δ*S* = 1 is regarded as the difference required for two colours to be noticeable, or one just noticeable difference (JND). RN predictions have been tested using behaviour of several species, and work reasonably well (e.g., Fleishman et al., 2016; Kelber et al., 2003; Olsson, Lind, & Kelber, 2015). However, RN modelling must be used with caution for four reasons: (a) RN was designed to predict discrimination when Δ*S* is near one (near the threshold), and may be inaccurate for colours that are very different (Δ*S* > 1). This arises because the relationship between predicted difference and perceived difference is nonlinear. For example, consider three colours A, B and C. Let the difference between A and B be Δ*S* = 2, and between A and C Δ*S* = 8; the frequent implicit assumption is that Δ*S* = 6 between B and C. Although the JND scale suggests that A and B are almost as far apart as A and C, if the perception response to Δ*S* is logarithmic then B and C may not be perceived as very different from each other and both perceived as very different from A. (b) Behaviour observations often show that some colours are discriminated as predicted by RN while others are not (unpublished observations; Cheney, pers. comm 2017; Fleishman et al., 2016; Olsson et al., 2015). This may arise from preexisting colour preferences. Different RN models need to be used at higher and lower light intensities to make good predictions (Olsson et al., 2015; Vorobyev & Osorio, 1998). (c) Data on actual receptor noise values are scarce yet they underpin all Δ*S* calculations (Olsson, Lind, & Kelber, 2017). (d) The model is limited; it is designed to capture what happens during early processing in the retina and does not include downstream processing in the brain, including decision making as well as perception. Estimates of detection and discrimination depend upon animals making decisions. Consequently, the RN could be correct in the retina, but later neural processes may mean that behaviour-tests may not match all RN predictions (for example Dyer, Spaethe, & Prack, 2008). Despite these limitations, what happens at the early retinal level is important because all visual processing starts there (Lythgoe, 1979). The RN model must be treated simply as a starting point analogous to the Hardy–Weinberg equilibrium in population genetics. In addition to providing a foundation, RN model estimates of Δ*S* can be used to explore the visual effect of the entire colour pattern, not just differences between colour pairs.

1.2 | **Assessment of patch edges**

Previous work with colour discrimination and Δ*S* has not accounted for whether compared patches were in contact or separated by other colours. Here, we explore Δ*S* explicitly for patches which come into contact because what happens at the patch edges may be important. The neurobiological justification for assessing the effects of edges (transitions between patches) is described in detail in Elder and Sachs (2004), Stevens and Cuthill (2006), Troscianko et al. (2017) and Endler (2012). Briefly, the photoreceptors in both vertebrate and invertebrate visual systems are

connected to neurons that calculate the differences between the photoreceptor outputs over a small visual field. Groups of photoreceptors involved in opponency are called units and can not only detect colour but also serve as edge detectors. Units consist of two adjacent groups (zones) of photoreceptors covering a small part of the visual field, and a ganglion cell calculates the difference in outputs between the two groups' opponency (Dowling, 2012; Dyer, Paulk, & Reser, 2011; Kelber, 2016; Sanes & Zipursky, 2010). If the photoreceptors in the two zones are sensitive to *different* wavelengths, then the unit outputs are colour signals because colour is based upon intensity differences among different parts of the visible spectrum. Edges between patches of different colours are detected if the edge cuts across the boundary between the unit zones. If the photoreceptors in the unit are sensitive to the *same* wavelengths then the outputs result from patch edges at the zone boundary regardless of chroma if they differ in luminance. Both edge types are detected depending upon the physical size of the retinal unit relative to the image and/or how rapidly the eye scans the colour pattern (Dowling, 2012; Elder & Sachs, 2004; Gegenfurtner & Sharpe, 1999; Kelber, 2016; Sanes & Zipursky, 2010). The stronger the edges (steeper gradients and greater differences between the patches, yielding larger Δ*S* between the two patches), the stronger the signal they produce in the units. The longer the edges the more units that they will stimulate. Consequently, both the geometry and reflectance spectra of patches in colour patterns affect edge intensity and conspicuousness. Both chromatic and achromatic opponent units operate over small parts of the visual field, suggesting that local colour pattern properties may be more important than global properties.

The effects of edges also depend upon the visual acuity (resolution angle) of the viewer as well as the distance between the viewer and the colour pattern. Acuity effects may eliminate or modify visual contrast, particularly if the visual fields of the opponent units are larger than the patches. Although opponent units are known to cover a small part of the visual field, their actual sizes are unknown in most species. Moreover, there may be higherorder units in the brain which will not be accounted for by the retinal estimations. For these reasons, calculations of edge effects must be done with good data on acuity and viewing distance, and results treated as a first approximation, even if the unit field sizes are known.

2 | **MATERIALS AND METHODS**

Let *C* be the number of colour and luminance classes in a given colour pattern. The challenge of this, and any other colour pattern analysis, is identifying the *C* classes and making identification repeatable. This is a classic image analysis problem known as image segmentation, and is particularly problematic where there are colour or luminance gradients. One could identify the classes by (human) eye, but for almost all diurnal nonprimate animals their vision is sufficiently different from humans

that human-based classifications may range from unreliable to misleading, particularly if there are UV reflecting patches present. Another method is to move a portable reflectance spectrometer sensor over the animal's body to determine how patch reflectance spectra vary. If any of the spectra vary more than is visible to the human eye then samples must be taken from both the invisible and visible patches and labelled accordingly. A third method which is less likely to miss patches invisible to humans is to scan the entire body evenly in a grid with a spectrometer and use various clustering methods to classify the colour/luminance patches by spectral clusters. This can be refined by doing clustering of calibrated photographic pixels (Van Belleghem et al., 2018), spectra or cone stimulations and clustering based upon Δ*S* (van den Berg et al., in preparation). A final stage is ensuring that all patches in the segmented image are visible with the viewer's visual acuity and viewing distance. In what follows, we will assume that the patch classification into *C* classes has been completed along with a matching list of cone captures estimated from patch spectra (Endler & Mielke, 2005) or from calibrated photographs (Troscianko & Stevens, 2015).

All cone capture estimates should be made under the normal viewing conditions in the wild. This includes the distances between signals and receivers as well as light intensity because visual acuity declines with declining light and the combination of the visual acuity of the viewer and the viewing distance affects the smallest patch which can be resolved. If two patches are not resolved at the ordinary distance and light intensity, then the two patches should be combined into a single patch and the patch spectrum should be an average of the two spectra, weighted at each wavelength by the relative areas of the two indistinguishable patches. The geometry of patches should be relevant to the viewer's vision and visual conditions during viewing.

2.1 | **Relative frequency of each patch edge class**

The first stage of analysis of a colour pattern is to estimate the lengths or relative frequencies of the *C* edges between adjacent colour/luminance patch classes. A *C* × *C* matrix should be made to organize the list of all possible edge or colour/luminance transition classes (example in Table S1). For *C* classes there are at most *E* = *C*(*C* − 1)/2 different edge or transition classes (Endler, 2012). Note that in any one colour pattern, it is likely that not all patch classes will contact all other classes, especially for larger *C*. Consequently, the number of observed kinds of different transitions (edges) among patches, *n*, will be less than the maximum possible number of edge classes, *E*. A simple example is found in the North American coral snakes *Micrurus fulvius* and *Micrurus euryxanthus*, where there are colloquial phrases to distinguish them from the Batesian mimetic king snakes *Lampropeltis* species, such as "red on yellow, beware the fellow, red on black, it's all right Jack." There are three possible transitions in these snakes: redyellow, yellow-black and red-black, but red-black is a missing transition in these coral snakes, while and red-yellow is missing from the mimics (this is not true for other coral snake species). Once the edge classes are determined, they need to be mapped onto the outline of the animal. An example using a male guppy *Poecilia reticulata* is shown in Figure 1a–c.

The relative frequency or length of each transition class can be obtained from one of two methods. Measure the length for each edge directly from the edge map (Figure 1c) or extract edges from the zone map of the patch pattern. A zone map is simply a digital mosaic diagram of the same size as the original image where each pixel contains a label for the colour/luminance class in which it is found (Figure 1b); this is also known as a label matrix. The zone map also allows additional parameters to be extracted (Endler, 2012). Because pixels are in a square array, diagonal distances as well as horizontal or vertical distances will have to be used for slanted edges, but this should produce minor errors if the pixel spacing is small enough. Accumulating the colour/luminance class transitions over all adjacent pixels in the zone map yields a transition or adjacency matrix, where rows and columns correspond to the colour classes (as in Table S1). The transition matrix diagonal entries are proportional to each colour's relative area. The off-diagonals yield the relative frequency of each transition class or edge (Endler, 2012). This matrix is symmetric with separate estimates of a particular transition class in both the upper and lower off-diagonals (Table S1). For further analysis, add the equivalent upper and lower off-diagonals together in order to obtain frequencies of each patch edge type (Table S2); these numbers are equivalent to lengths of edges extracted directly from the image (Figure 1c), and, like lengths, can be divided by their grand total to yield relative edge lengths. The result of either method is a *C* × *C* lower off-diagonal transition or edge matrix, T_F (Table S2), where the lower off-diagonal numbers are the lengths or frequencies of the edge class defined by the intersection of the corresponding row and column. For example, if a particular cell (row *i* and column *j*) has the value *fij*, then *fij* is the frequency of the transition between colours *i* and *j* in both directions*.* Potential transitions between colours which are not observed because the appropriate patches do not come in contact will be represented by f_{ii} = 0. A given f_{ii} in $\mathbf{T_F}$ estimates how commonly two colour/luminance classes share a common edge or the size of each patch type boundary.

2.2 | **Magnitude hence salience of patch boundaries**

The second and novel stage of analysis is an estimate of how conspicuousness the edge is likely to be to a given viewer under given environmental conditions. The receptor noise Δ*S* estimate for any pair of colours is an estimate of edge conspicuousness or strength because colour and/or luminance differences are easier to detect for larger Δ*S*. We can obtain photon captures for each patch using the irradiance spectrum illuminating the pattern in nature, the reflectance spectrum of the patch in the direction of the viewer, the transmission spectrum of the air or water between the pattern and viewer in nature, the transmission spectrum of the eye optics, and the absorption spectra of the visual pigments in each photoreceptor class (Endler, Cole, & Kranz, 2005; Endler & Mielke, 2005; Kelber et al., 2003; Lythgoe, 1979). We obtain the Δ*S* for all possible pairs of patches in the colour pattern (as did Siddiqi et al., 2004) based upon the photoreceptor captures, the relative abundance of each photoreceptor, and an assumption about the level of receptor noise (the Weber fraction, Kelber et al., 2003). Methods for obtaining Δ*S* are well established, including in the r package pavo (Maia, Eliason, Bitton, Doucet, & Shawkey, 2013). The Δ*S* for each kind of colour class comparison is then placed in the appropriate row and column in a second matrix with *C* rows and *C* columns (same format as Table S2). It is only necessary to fill in the lower off-diagonal because the upper off diagonal should be identical, and the diagonals will be zero (no difference in a comparison of the same colour). This yields a *C* × *C* transition or Δ*S* matrix **T**_{*S*} with data in the lower off-diagonal, where each entry *s_{ii}* is the Δ*S* for patch colour/luminance classes indicated by row *i* and column *j*. Two different **T^s** should be calculated by: (a) using all the photoreceptors used in colour vision (e.g., cones in vertebrates) to obtain chromatic Δ*S* and (b) using the specific photoreceptor(s) used in luminance to get luminance or achromatic Δ*S*. Consequently the result will be two Δ*S* transition matrices, **T**_{SC} from the chromatic ΔS calculations and **T_{SL}** from the luminance or

achromatic ΔS calculations. The rows and columns of T_{SC} and T_{SL} must correspond exactly in both length (*C*) and row order to the rows and columns of T_F .

The matrix T_F should contain the relative frequencies of each kind of transition and the matrices T_{sc} and T_{SL} should contain the RN estimate of how differently (Δ*S*) the two adjacent colours in the corresponding T_F entry stimulate the retina with respect to chromaticity or luminance, respectively. They should have the same form as Table S2. The lower off-diagonal values of these three matrices should be converted into vectors (one-dimensional lists) of length *E* = *C*(*C* − 1)/2, and placed together in a *E* × 3 data matrix for convenience in further calculations (see Table S3). This data matrix has the edge length, the chromatic Δ*S*, and the luminance Δ*S* for the transition (edge) class *k* in row *k*; call these f_k , sc_k and sl_k where $k = 1, 2, ..., n$ patch classes. Table S3 shows an example where *k* = *a*, *b*, …, *f* and *n* = 9.

The data matrix provides a correspondence between edge lengths and their estimated visual magnitudes or salience*.* This,

FIGURE 1 Example analysis of a male guppy colour pattern. (a) Photograph of a guppy (scale not shown). (b) Part of the resulting zone map indicated by the circle in panels (a) and (c). Each pixel has a code indicating which colour/luminance class overlaps that pixel (see Endler, 2012 for details). (c) Edge map; this can either be derived directly from the photograph (a) or from the zone map (b). (d) Diagram in which the *x*,*y* (horizontal) coordinates correspond to the edge map in (c) and the vertical axis corresponds to the chromatic Δ*S* between adjacent patches under specific ambient light conditions. (e) as in (d) but for luminance Δ*S*. Note the lack of topographic correspondence between the chromatic and luminance diagrams. For brevity we will refer to (d) and (e) as "Fort Diagrams" because they resemble old fashioned fortresses)

along with an annotated map of the patch boundaries (Figure 1c), allows plotting the geometry of estimated patch boundary strengths for both chromatic and luminance Δ*S*. In these diagrams, the x and y axes are as in Figure 1c and the *z*-axis is proportional to Δ*S*. Figure 1d,e show 3D plots of chromatic and luminance edge Δ*S* for the guppy shown in Figure 1a. We will call these diagrams "fort diagrams" because they resemble forts and "fort" means strong in French and Latin, so also refers to boundary strength. Note the very different geometric patterns of chromaticity and luminance boundaries in Figure 1d,e; the guppy shows high edge contrasts in different places for chromaticity and luminance. More specifically, luminance contrast is dominated by the black patch edges almost independently of the patch class they contact. Note the very high luminance Δ*S* (height) where a black patch contacts the very highly reflective silver patch towards the front of the guppy in (compare Figure 1a,e).

2.3 | **Combining patch properties and edges**

If edges contribute significantly to the conspicuousness of the entire colour pattern, then we may be able to capture at least part of what makes a colour pattern conspicuous by obtaining an aggregate measure of the edge magnitudes. We suggest the mean, standard deviation and CV of the edges' Δ*S*, weighted by their corresponding lengths or frequencies. These are calculated from either the sc_{k} (chromatic ΔS) or sl_k (luminance ΔS) as s_k from T_{sc} or T_{SL} , and using the f_k (from $\mathbf{T}_\mathbf{F}$) as weights in the formulae:

weighted mean:
$$
m_{\Delta S} = \frac{\sum_{k=1}^{E} f_k S_k}{\sum_{k=1}^{E} f_k}
$$
 (1)
\nweighted standard deviation: $S_{\Delta S} = \sqrt{\frac{n \sum_{k=1}^{E} f_k (S_k - m_{\Delta S})^2}{(n-1) \sum_{k=1}^{E} f_k}}$ (2)

weighted coefficient of variation:
$$
CV = \frac{S_{\Delta S}}{m_{\Delta S}},
$$
 (3)

where *E* is the number of all possible different kinds of edges and *n* is the number of observed transitions (S_k) or those with nonzero *fk* (Filliben, Heckert, & Lipman, 1996); *n* ≤ *E*. The Supplemental Appendix provides a MATLAB function to calculate the weighted mean and standard deviation; the equivalent functions in R are wt.mean and wt.sd within the r package *SDMTools* (VanDerWal, Falconi, Januchowski, Shoo, & Storlie, 2014). Formulae 1–3 are the same formulae used to calculate the mean, *SD* and CV of chroma and luminance for overall within-contrast measurements, substituting chroma or luminance for s_k and mean chroma or luminance for $m_{\Lambda S}$; but circular statistics have to be used for hue angles (Endler & Mielke, 2005).

The weighted mean $m_{\Lambda S}$ is an estimate of the average conspicuousness of the whole pattern but weighting longer edges more than shorter ones. Similarly, the weighted standard deviation $s_{\Delta S}$ measures how variable the edge magnitudes are over the entire pattern weighted by their lengths. The coefficient of variation CV is the standard deviation relative to the mean. If it is known that the viewer attends only to part of the pattern then $m_{\Delta S}$ and $s_{\Delta S}$ should be calculated over the relevant part of the colour pattern. The assumption here is that a longer edge will stimulate more opponency units in the retina, and when the pattern is moving, a longer edge will sweep out more of the retinal area than a smaller edge. It is not known or obvious whether the mean, standard deviation, or even the CV would be a better predictor of salience. For example, a larger $m_{\overline{\Delta S}}$ might be more stimulating, but it is unknown whether this should be accompanied by a smaller $s_{\Delta S}$ for consistently high stimulation over the entire pattern, or a larger *s*_{ΔS} and hence less predictable edge magnitude to prevent sensory adaptation. Using CV instead of the standard deviation might be important if a given degree of variation is not more important for small versus larger means. These conjectures can only be answered by extensive behavioural studies with different $m_{\Delta S}$ and $s_{\Delta S}$, measured under the appropriate conditions and appropriate parts of the body.

Boundary strength analysis can be applied to an animal colour pattern in order to estimate within-pattern visual contrast. They can also be applied to visual backgrounds to estimate within-background contrast, and if so estimates of signal-background contrast can be made by comparing parameters of animal and background. For simplicity the examples will concentrate on within-signal contrast.

3 | **EXAMPLES AND THEIR IMPLICATIONS**

To illustrate and explore the biological significance of BSA, we chose two species that are polymorphic in their patch colour, luminance and geometry, male guppies *P. reticulata* and Gouldian finches *Erythrura gouldiae,* because they have very different signal and signalling geometry. This allows us to showcase the power of the method in colour pattern research and the important effects of local patterns and viewing angle between the sender and receiver.

3.1 | **Guppy examples and implications**

Male guppies are extremely polymorphic in patch geometry and properties (Endler, 1978, 1980). Figure 2 shows Fort diagrams of six male guppies in the same format as Figure 1c,d, ordered by decreasing chromatic $m_{\overline{\Delta S}}$ and calculated in open/cloudy light conditions (Endler, 1993b). The numbers are $m_{\Delta S}$ and CV from Equations (1) and (3). These six randomly selected guppies yield five observations: (a) Each guppy has edges with unique geometry. This goes with the considerable polymorphism of male guppy colour patterns (photos in Endler, 1978). (b) There is little geometric correspondence between the strength and positions of chromatic and achromatic (luminance)

Luminance JND

FIGURE 2 Examples of Fort Diagrams for 6 different guppy colour patterns, arranged in order of decreasing chromatic *m*_{ΔS}. Rows correspond to the same individual guppy and columns refer to the guppy's chromatic or luminance Fort diagram, respectively. Numbers under the diagrams for each row are chromatic *m*_{Λ S} and CV (left column) and luminance *m*_{Λ S} and CV (right column) for the same guppy. Note the lack of topographic correspondence between the chromatic and luminance diagrams, and the variation among individuals

edges; the peaks in chromaticity do not correspond with peaks in luminance, and both depend upon which pair of patches form the edge. (c) The spatial correlation between chromatic and luminance Δ*S* is always negative within a guppy although not always significantly so (Figure 3a,b). (d) The negative correlation between the two Δ*S* is not present when we consider all possible patch combinations (Figure 3c); patch contacts and hence boundary strengths are clearly nonrandom. (e) Guppies differ in how variable their Δ*S* heights are, indicating variation in which patches form common edges.

Maximum chroma and luminance should be negatively correlated because the only way to increase chroma is to remove parts of a spectrum. Removing part of the spectral radiance reduces luminance. At the same time, it increases the differences in stimulation among different photoreceptor classes, increasing chroma (Endler & Mielke, 2005; Endler & Théry, 1996; Endler et al., 2005). However, $m_{\Delta S}$ and $s_{\Delta S}$ depend upon geometry as well as

patch properties and consequently predictions based upon patch properties alone may be invalid. For example, chromatic and luminance $m_{\Lambda S}$ might even be positively correlated if sexual selection jointly increases both luminance and chromatic *m*_{ΔS}, which would make males more conspicuous. We tested for a possible chromatic-luminance relationship by analysing 200 male guppies. The two $m_{\Delta S}$ are positively correlated (Figure 3d). This is not what one would expect from random patch geometry, where every patch class has an equal probability of contacting the others (see also Figure 3a,c). It suggests that particular colours are adjacent and adjacency has evolved to set particular levels of overall conspicuousness, as estimated by $m_{\Delta S}$. Random associations yield different *m*_{ΔS}. The relationship for s _{ΔS} is also positive (Figure 3e), but the 200 points are widely scattered and appear in 3 clumps. This suggests partially discontinuous variation among fish boundary Δ*S*, and could result from polymorphic colour pattern genes that

FIGURE 3 Relationships between chromatic and luminance edge statistics in guppies. (a) Significant negative correlation between chromatic and luminance Δ*S* within a guppy having an average correlation value. (b) Distribution of the correlations among 11 guppies; all are negative but two are not significantly negative. (c) Lack of correlation between all possible chromatic and luminance edges; note the larger rage and higher joint values compared to a. (d) The relationship between chromatic and luminance $m_{\Lambda S}$ of 200 guppies. (e) Relationship for *s*_{ΔS}. (f) relationship for $CV_{\Delta S}$

control particular sets of spots (review in Endler, 1978). The correlation and clumping for CV (Figure 3f) is lower than for $m_{\Delta S}$ and *s*_{AS}. Patterns of variation in boundary strength could predict fitness in any species because they affect pattern conspicuousness and hence colour pattern function and fitness.

Figure 4 shows chromatic and luminance $m_{\Delta S}$ and $s_{\Delta S}$ distributions for the 200 guppies analysed. The means are moderately symmetrically and unimodally distributed but the standard deviations are multimodal, as in Figure 3e,f. Note that $m_{\Lambda S}$ > 1.5 indicates that, on average, the boundaries are detectable by females, but some may not be ($m_{\Delta S}$ = 1 is one JND, the threshold for distinguishing patches). Patches with similar colours or luminance which would lead to smaller ΔS and $m_{\Delta S}$ tend not to be adjacent. In general, we hypothesise that having adjacent patches with larger Δ*S* would be advantageous in conspicuous signalling, but disadvantageous for crypsis. If most boundaries are not detectable and a few were, this might be a previously unrecognised form of disruptive colouration.

The thick black line in Figure 4 is the estimate for randomly arranged patch classes, as opposed to their observed geometry. This was calculated by letting every patch class contact every other patch class as in Figure 3c. For $m_{\Lambda S}$, it is larger than actually found in any fish, and for *s*_{ΔS}, it is larger than all fish except for chromatic *s*_{ΔS} where it is larger than 98% of the fish. This suggests that the observed colour patterns are less conspicuous than they would be if the patches were arranged at random. One would at first think that this is contrary to that expected because we assume that females should mate with males with larger $m_{\Delta S}$ because they are more conspicuous than those with smaller $m_{\Delta S}$. However, visually hunting predators are always present in natural guppy populations, resulting in variation in the trade-off between sexual selection and predation (Endler, 1978, 1980). We speculate that guppies have been selected over millions

FIGURE 4 The distributions of chromatic and luminance edge statistics $m_{\Delta S}$ and $s_{\Delta S}$ of the 200 guppies in Figures 3 and 4. (a) Chromatic $m_{\Lambda S}$, (b) chromatic *s*_{Δ*S*}, (c) luminance *m*_{Δ*S*}, (d) luminance *s*_{ΔS}. All guppies have $m_{ΔS}$ > 1 indicating that adjacent patches are always discriminable to guppies under the environmental conditions. The thick vertical lines show the same statistics if the colour patches were distributed at random over each guppy's body; every patch class had an equal probability of contacting the others. Almost all guppies show smaller values than expected from random patch locations

of generations for optimal edge strengths balancing sexual selection and predation. We predict that samples taken from high predation populations would have distributions of $m_{\Delta S}$ and $s_{\Delta S}$ that extensively overlap Δ*S* = 1, indicating less conspicuous coloration representing the local balance between sexual selection and predation. This may apply to any species where there is a shifting balance between sexual selection and predation.

3.2 | **Gouldian finch examples and implications**

Gouldian finches provide examples of additional insights that can be gained from BSA. Gouldian finches have three polymorphs differing in head colour: black, yellow (golden) or red. Both males and females are coloured with females having less chromatic colours and a mauve rather than a purple chest. Unlike guppies, which have a relatively flat surface that is displayed towards females, Gouldian finches have a 3D colour pattern in which the relative proportion of patches and edges changes with viewing angle. Consequently, we present Fort diagrams from Gouldian finches seen at two viewing angles: a ¾ view and a side view (Figure 5a,b). The analysis of the ¾ view is shown in Figures 5 and 6 and the side view in Figure 7. More details are shown in the Supplemental Appendix.

Like guppies, there is a divergence between chromatic and luminance Δ*S* (Figure 5c–h) and the spatial correlation between them is negative (except in the golden female morph). However, with fewer points than in the guppy data, none of the correlations are significant. Nevertheless, each correlation is smaller than the correlation between all possible pairs of colours for that morph and gender (see

Supplemental Appendix) suggesting that the negative correlation has some function in both species.

Given that the chromatic and achromatic patterns are different and almost complementary we suggest that the chromatic and achromatic components of colour patterns could be used for different functions, such as sexual selection, species recognition, or defense. Chromaticity and luminance are processed independently, and there is variation in their relative importance in stimulus choice and discrimination, among many species including crabs, psyllids, honeybees, bumblebees, flies, hawkmoths, birds and humans (Baldwin & Johnsen, 2012; Dyer et al., 2008; Farnier, Dyer, & Steinbauer, 2014; Giurfa, Vorobyev, Brandt, Posner, & Menzel, 1997; Keil, Miskovic, Gray, & Martinovic, 2013; Kelber, 2005, 2016; Osorio & Vorobyev, 2005; White & Kemp, 2016, 2017; White, Rojas, Mappes, Rautiala, & Kemp, 2017; Zhou, Ji, Gong, Gong, & Liu, 2012). This suggests that chromatic and achromatic channels could have different functions in any taxa. There are also distance effects, probably due to the fact that in many animals, visual acuity is greater for achromatic than chromatic stimuli. For example, bees use chromatic cues when they subtend larger angles on their retina and achromatic cues when the visual angles are smaller (Giurfa et al., 1997). This means that achromatic cues may be more useful at greater distances than chromatic cues, especially at lower light levels when acuity decreases, and colour vision stops working at still lower irradiances. Moreover, chromatic and luminance components are roughly independent in natural scenes (Hansen & Gegenfurtner, 2009), suggesting that crypsis may be possible independently of signalling. The functional differences between chromatic and achromatic edges are worth further investigation.

Gouldian finches also illustrate that: (a) The viewing angle (b) Edge map traced from a side view photograph. (c–h) Fort diagrams of the three male morphs (rows) showing the difference in pattern for chromatic and luminance Δ*S* (columns) in the 3/4 view

significantly affects the perceived relative area of each patch, significantly affecting $m_{\Delta S}$ and $s_{\Delta S}$; the ³/₄ view having higher $m_{\Delta S}$ and often higher s_{AS} than the side view (Table 1). This highlights the importance of recording the viewing angle during visual signalling. (b) Sexual dimorphism within each morph is associated with reduced edge intensities, $m_{\Delta S}$ and s_{ΔS}, in females of all morphs for both chromatic and achromatic Δ*S* (Figure 6, Table 1), with less reduction in achromatic Δ*S* (Table 1). This illustrates the utility of BSA in estimating sexual dimorphism. (c) Within males or females, the three morphs differ in chromatic *m*_{Λ S} with the golden and red morphs similar but different from the black morph (Table 1). They differ less in achromatic $m_{\Delta S}$, and there is surprisingly little variation in $s_{\Delta S}$ among morphs; perhaps this is the sign of a species-specific signal. (d) There is a clear difference in pattern between the head and the rest of the body, with the head values larger than the body. The difference in location-specific edge intensities is stronger in the side view. This reiterates the importance of calculations using the same view angle as used by the viewers, but it also shows a weakness of using $m_{\Delta S}$ and $s_{\Delta S}$ calculated over the entire body. It may be reasonable in guppies or other species that present the

Female, Black, 3/4 view, Chromatic ΔS (b)

(d) Female, Golden, 3/4 view, Chromatic ΔS

 $\Delta S \ge 10$

 $SS \times 10$

 (f)

 Ω

(c) Male, Golden, 3/4 view, Chromatic ΔS

Male, Black, 3/4 view, Luminance ΔS (e)

(g) Male, Golden, 3/4 view, Luminance ΔS

Female, Black, 3/4 view, Luminance ΔS

(h) Female, Golden, 3/4 view, Luminance ΔS

FIGURE 6 Fort diagrams showing sexual dimorphism in the black (a,b,e,f) and golden-headed morphs (c,d,g,h) with respect to both chromatic (a–d) and luminance (e–h) Δ*S* in the 3/4 view. The red-headed morph does not differ very much from the golden-headed morph (see Supplemental Appendix for all fort diagrams)

entire side of a relatively flat surface to the viewer, but it will be inaccurate if the viewer attends more to some parts of the body than the others. The stronger edges in the Gouldian finch heads may be associated with, and even selected by, conspecifics paying more attention to the heads than the rest of the body. The rest of the body may be used in species recognition and, or, reduction of predator risk. Consequently, $m_{\Delta S}$ and $s_{\Delta S}$ should be calculated on the parts of the colour pattern used in social interactions for signal design assessment whereas they should be calculated separately on the parts of the body seen by predators (using predator vision parameters). These two functions may be spatially separated. Clearly we need to know about the geometry of signalling as much as the geometry of the signals for accurate use of BSA.

FIGURE 7 Fort diagrams of side views of the black (a,b,e,f) and golden-headed (c,d,g,h) morphs. See Supplemental Appendix for all fort diagrams

4 | **GENERAL PREDICTIONS**

Because BSA can be used to analyse any animal or plant colour pattern, it is useful to make some general predictions, based upon the assumption that edges are important in colour pattern detection and perception (Dowling, 2012; Gegenfurtner & Sharpe, 1999; Stevens & Cuthill, 2006), and that stronger edges (larger Δ*S* and greater length) are more effective than weaker edges.

1. If $m_{\Delta S}$ is important in intraspecific signalling then it should predict behaviours such as mate choice or any other visually based choice behaviour. The relative importance of chromatic and luminance $m_{\Delta S}$ is unknown, and this may vary among higher taxonomic groups. Consequently, we predict that the relationship between *m*_{ΔS}, pattern conspicuousness, decision-making, and fitness will be context, habitat and species specific. Restriction of $m_{\Lambda S}$ to calculations just over the part of the colour pattern tracked by viewers should be limited to species with well-studied signalling geometry, or will have to wait for more advances in eye-tracking methodologies

2. If s_{AS} is important in colour pattern conspicuousness then it should predict visually-based choices. However, it is not clear whether larger or smaller $s_{\Delta S}$ increases the overall conspicuousness. Small $s_{\Delta S}$ (or CV) could give a consistently higher stimulation to the retina. However, larger s _{ΔS} might be more effective if spatially similar Δ*S* (low *s*_{Δ*S*}) leads to sensory adaptation and hence inefficient reception. This could be particularly true for fast moving patterns.

TABLE 1 Gouldian finch mean ($m_{\Lambda S}$) and *SD* ($s_{\Lambda S}$) of patch edge chromatic (Cr) and luminance (Lm) Δ*S*, weighted by edge lengths

Cr $m_{\overline{\Delta S}}$	Cr s _{AS}	Lm $m_{\Lambda S}$	$\mathsf{Lm}\,s_{\Lambda S}$	Morph-gender- view
7.56	4.97	11.07	10.43	Black, Male, 3/4 view
5.71	4.25	7.84	9.29	Black, Male, Side view
4.49	2.64	8.55	6.53	Black, Female, 3/4 view
3.19	2.21	5.78	6.50	Black, Female, Side view
12.30	5.46	11.84	11.08	Golden, Male, $3/4$ view
8.58	5.55	9.75	10.91	Golden, Male, Side view
6.70	3.43	9.90	10.41	Golden, Female, 3/4 view
4.77	3.57	8.33	9.91	Golden, Female, Side view
11.44	4.94	12.95	9.56	Red, Male, 3/4 view
7.96	4.95	9.68	10.10	Red, Male, Side view
5.75	2.98	11.80	9.30	Red, Female, 3/4 view
440	3.25	9.76	9.24	Red, Female, Side view

- **3.** For colour patterns, or components used in signalling, edges should have $m_{\Lambda S}$ > 1 with respect to chromatic and luminance ΔS ; edges with Δ*S* ≤ 1 are unlikely to be detected. Patterns with small *m*_{ΔS} have fewer detectable edges, leading to inefficient visual signalling. For crypsis, having mostly undetectable edges ($m_{ΛS}$ ≤ 1) is an advantage. However, if the background has many Δ*S* > 1 and the animal has many Δ*S* ≤ 1 the animal's shape will be conspicuous. If both have many Δ*S* > 1 then the pattern may be cryptic (Endler, 1978) or disruptively coloured (Endler, 2006).
- **4.** For colour patterns or pattern parts used in signalling, the distribution of both $m_{\Delta S}$ and $s_{\Delta S}$ should be different from those of the visual background with respect to either chromatic or luminance Δ*S* or both. The animal-background colour pattern component distributions should be similar for cryptic species, or parts of the colour patters that are seen more often by predators than conspecifics.
- **5.** The animal-background match or mismatch of both $m_{\Lambda S}$ and s_{ΛS} should differ in different parts of the animal's body for species that are usually seen by predators from one viewing angle (e.g., above or behind) and by conspecifics from another viewing angle (e.g., frontal; e.g., Salticid spiders); parts viewed by predators should be more cryptic than parts viewed by conspecifics. Colour pattern functions could not only differ in regions of the body viewed from different angles, but may also differ when viewed

from different distances because this may cause some adjacent patches to blend (Endler, 1978).

- **6.** For prey species living in areas over a range of predation intensities, the fraction of edges with Δ*S* ≤ 1 should be relatively higher in areas with higher predation because Δ*S* ≤ 1 leads to poorer perception of separate patches, but the opposite is needed for disruptive colouration. The absolute fraction of edges with Δ*S* ≤ 1 should depend upon the background patch pattern. For example, in visual backgrounds with highly contrasting patches (most $\Delta S \gg 1$, large $m_{\Delta S}$) the $m_{\Delta S}$ and the distributions of Δ*S* in the animal and backgrounds should be more similar in areas of higher predation intensity than areas of lower predation. For prey species that use only parts of the pattern for signalling, the signalling components should be smaller, with shorter edges and lower Δ*S* in areas of greater predation risk.
- **7.** For species attending more to chromaticity than luminance in intraspecific signalling the chromatic *m*_{ΔS} and most or all chromatic Δ*S* should be larger than 1 with the opposite for luminance. This ensures that the pattern is maximally conspicuous to the receiver's visual system. A similar pattern should appear for luminance *m*_{ΔS} and Δ*S* in species using luminance more than chromaticity.

5 | **GENERAL QUESTIONS**

There is so little known about the implications of estimates of patch boundary strengths that predictions are limited, but there are several questions which are worth further investigation until we can make explicit predictions.

- **1.** Which is more important in intraspecific signalling, $m_{\Lambda S}$ or $s_{\Lambda S}$? If both are important, does their relative importance change with the complexity of the visual background or the mixture of different intraspecific and interspecific viewers?
- **2.** *m*_{ΔS} and *s*_{ΔS} estimate the effects of patch boundaries on the overall colour pattern conspicuousness. It is also possible that within-pattern variation in hue, chroma and luminance of patches also affect overall conspicuousness, regardless of whether or not they come into contact (Endler & Mielke, 2005). What is the relative importance of overall variation in hue, chroma, luminance, and edge properties? Which measures successfully predict mate choice and survival under specific visual and ecological conditions?
- **3.** Do different aspects of salience allow for "private channels," allowing mitigation of the trade-off between being conspicuous to potential mates and inconspicuous to predators? This might be most likely if, for example, predators used different visual processing, different components of the colour patterns, or different viewing distances and angles than the prey use for intraspecific signalling.
- **4.** How do patch and patch edge properties communicate signal content? Do they constrain content enough to make predictions

about the kind and amount of information to be transmitted to conspecifics?

In sum, within the limitations outlined in Sections 1.1 and 1.2, BSA will enable these questions to be addressed in any species that use vision to make decisions based upon reception and perception of a sender's colour pattern.

ACKNOWLEDGEMENTS

We thank three reviewers for excellent and useful comments on the manuscript, Adrian Dyer for useful comments about the receptor noise model and Adelaide Sibeaux for comments on the manuscript and being willing to try it out as a way to predict guppy mating success (in progress). We thank the Australian Research Council for two discovery grants which supported this research (DP110101421 and DP150102817).

AUTHORS' CONTRIBUTIONS

J.A.E. devised the method, tested it and wrote the first draft of the paper; G.L.C. and A.M.K. prepared the guppy photographs, extracted the colour patch geometry from photographs, and helped revise the paper; J.A.E. did the Gouldian Finches.

DATA ACCESSIBILITY

A MATLAB script for calculating weighted means and standard deviations is found in the online supplemental material and in Dryad. Data and MATLAB script available from the Dryad Digital Repository:<https://doi.org/10.5061/dryad.g66247g>(Endler, Cole, & Kranz, 2018).

ORCID

John A. Endler <http://orcid.org/0000-0002-7557-7627> *Gemma L. Col[e](http://orcid.org/0000-0002-3365-3580)* <http://orcid.org/0000-0002-3365-3580> *Alexandrea M. Kranz* <http://orcid.org/0000-0002-7980-1374>

REFERENCES

- Arnold, S. J. (1983). Morphology, performance and fitness. *American Zoologist*, *23*, 347–361. <https://doi.org/10.1093/icb/23.2.347>
- Arnold, S. J. (2003). Performance surfaces and adaptive landscapes. *Integrative and Comparative Biology*, *43*, 367–375. [https://doi.](https://doi.org/10.1093/icb/43.3.367) [org/10.1093/icb/43.3.367](https://doi.org/10.1093/icb/43.3.367)
- Baldwin, J., & Johnsen, S. (2012). The male blue crab, *Callinectes sapidus*, uses both chromatic and achromatic cues during mate choice. *Journal of Experimental Biology*, *215*, 1184–1191. [https://doi.org/10.1242/](https://doi.org/10.1242/jeb.067512) [jeb.067512](https://doi.org/10.1242/jeb.067512)
- Clark, R. C., Santer, R. D., & Brebner, J. S. (2017). A generalized equation for the calculation of receptor noise limited colour distances in *n*-chromatic visual systems. *Royal Society Open Science*, *4*, 170712. <https://doi.org/10.1098/rsos.170712>
- Dowling, J. E. (2012). *The retina, an approachable part of the brain*. Cambridge, MA: Harvard/Belknap Press.
- Dyer, A. G., Paulk, A. C., & Reser, D. H. (2011). Colour processing in complex environments: Insights from the visual system of bees. *Proceedings of the Royal Society B: Biological Sciences*, *278*, 952–959. <https://doi.org/10.1098/rspb.2010.2412>
- Dyer, A. G., Spaethe, J., & Prack, S. (2008). Comparative psychophysics of bumblebee and honeybee colour discrimination and object detection. *Journal of Comparative Physiology A*, *194*, 617–627. [https://doi.](https://doi.org/10.1007/s00359-008-0335-1) [org/10.1007/s00359-008-0335-1](https://doi.org/10.1007/s00359-008-0335-1)
- Elder, J. H., & Sachs, A. J. (2004). Psychophysical receptive fields of edge detection mechanisms. *Vision Research*, *44*, 795–813. [https://doi.](https://doi.org/10.1016/j.visres.2003.11.021) [org/10.1016/j.visres.2003.11.021](https://doi.org/10.1016/j.visres.2003.11.021)
- Endler, J. A. (1978). A predator's view of animal color patterns. *Evolutionary Biology*, *11*, 319–364.
- Endler, J. A. (1980). Natural selection on color patterns in *Poecilia reticulata*. *Evolution*, *34*, 76–91. [https://doi.org/10.1111/j.1558-5646.1980.](https://doi.org/10.1111/j.1558-5646.1980.tb04790.x) [tb04790.x](https://doi.org/10.1111/j.1558-5646.1980.tb04790.x)
- Endler, J. A. (1993a). Some general comments on the evolution and design of animal communication systems. *Philosophical Transactions of the Royal Society, London B*, *340*, 215–225. [https://doi.org/10.1098/](https://doi.org/10.1098/rstb.1993.0060) [rstb.1993.0060](https://doi.org/10.1098/rstb.1993.0060)
- Endler, J. A. (1993b). The color of light in forests and its implications. *Ecological Monographs*, *63*, 1–27.<https://doi.org/10.2307/2937121>
- Endler, J. A. (2006). Disruptive and cryptic coloration. *Proceedings of the Royal Society B*, *273*, 2425–2426. [https://doi.org/10.1098/](https://doi.org/10.1098/rspb.2006.3650) [rspb.2006.3650](https://doi.org/10.1098/rspb.2006.3650)
- Endler, J. A. (2012). A framework for analysing colour pattern geometry: Adjacent colours. *Biological Journal of the Linnean Society, London*, *107*, 233–253. [https://doi.org/10.1111/j.1095-8312.2012.](https://doi.org/10.1111/j.1095-8312.2012.01937.x) [01937.x](https://doi.org/10.1111/j.1095-8312.2012.01937.x)
- Endler, J. A., Cole, G. L., & Kranz, A. (2018). Data from: Boundary strength analysis: Combining colour pattern geometry and coloured patch visual properties for use in predicting behaviour and fitness. *Dryad Digital Repository*,<https://doi.org/10.5061/dryad.g66247g>
- Endler, J. A., & Mielke Jr, P. W. (2005). Comparing entire colour patterns as birds see them. *Biological Journal of the Linnean Society, London*, *86*, 405–431. <https://doi.org/10.1111/j.1095-8312.2005.00540.x>
- Endler, J. A., & Théry, M. (1996). Interacting effects of lek placement, display behaviour, ambient light and colour patterns in three neotropical forest-dwelling birds. *American Naturalist*, *148*, 421–452. [https://doi.](https://doi.org/10.1086/285934) [org/10.1086/285934](https://doi.org/10.1086/285934)
- Endler, J. A., Westcott, D. A., Madden, J. R., & Robson, T. (2005). Animal visual systems and the evolution of colour patterns; sensory processing illuminates signal evolution. *Evolution*, *50*, 1795–1818. [https://](https://doi.org/10.1111/j.0014-3820.2005.tb01827.x) doi.org/10.1111/j.0014-3820.2005.tb01827.x
- Farnier, K., Dyer, A. G., & Steinbauer, M. J. (2014). Related but not alike: Not all Hemiptera are attracted to yellow. *Frontiers in Ecology and Evolution*, *2*(67), 1–12.<https://doi.org/10.3389/fevo.2014.00067>
- Filliben, J. J., Heckert, A., & Lipman, R. R. (1996). NIST Dataplot software reference manual, pp 2-66 to 2-67. US national Institute of Standards and Technology. Retrieved from [http://www.itl.nist.gov/div898/](http://www.itl.nist.gov/div898/software/dataplot/refman2/ch2/weightsd.pdf) [software/dataplot/refman2/ch2/weightsd.pdf](http://www.itl.nist.gov/div898/software/dataplot/refman2/ch2/weightsd.pdf)
- Fleishman, L. J., Perez, C. W., Yo, A. I., Cummings, K. J., Dick, S., & Almonte, E. (2016). Perceptual distance between colored stimuli in the lizard *Anolis sagrei*: Comparing visual system models to empirical results. *Behavoural Ecology and Sociobiology*, *70*, 541–555. [https://doi.](https://doi.org/10.1007/s00265-016-2072-8) [org/10.1007/s00265-016-2072-8](https://doi.org/10.1007/s00265-016-2072-8)
- Gegenfurtner, K. R., & Sharpe, L. T. (1999). *Colour vision, from genes to perception*. Cambridge: Cambridge University Press.
- Giurfa, M., Vorobyev, M., Brandt, R., Posner, B., & Menzel, R. (1997). Discrimination of coloured stimuli by honeybees: Alternative use of achromatic and chromatic signals. *Journal of Comparative Physiology*, *180*, 235–243.<https://doi.org/10.1007/s003590050044>
- Hansen, T., & Gegenfurtner, K. R. (2009). Independence of colour and luminance edges in natural scenes. *Visual Neuroscience*, *26*, 35–49. <https://doi.org/10.1017/S0952523808080796>
- Keil, A., Miskovic, V., Gray, M. J., & Martinovic, J. (2013). Luminance, but not chromatic visual pathways, mediate amplification of conditioned danger signals in human visual cortex. *European Journal of Neuroscience*, *38*, 3356–3362.<https://doi.org/10.1111/ejn.12316>
- Kelber, A. (2005). Alternative use of chromatic and achromatic cues in a hawkmoth. *Proceedings of the Royal Society B: Biological Sciences*, *272*, 2143–2147. <https://doi.org/10.1098/rspb.2005.3207>
- Kelber, A. (2016). Colour in the eye of the beholder: Receptor sensitivities and neural circuits underlying colour opponency and colour perception. *Current Opinion in Neurobiology*, *41*, 106–112. [https://doi.](https://doi.org/10.1016/j.conb.2016.09.007) [org/10.1016/j.conb.2016.09.007](https://doi.org/10.1016/j.conb.2016.09.007)
- Kelber, A., Vorobyev, M., & Osorio, D. (2003). Animal colour vision Behavioural tests and physiological concepts. *Biological Reviews*, *78*, 81–118.<https://doi.org/10.1017/S1464793102005985>
- Lythgoe, J. N. (1979). *The ecology of vision*. Oxford: Oxford University Press.
- Maia, R., Eliason, C. M., Bitton, P.-P., Doucet, S. M., & Shawkey, M. D. (2013). pavo: An R Package for the analysis, visualization and organization of spectral data. *Methods in Ecology and Evolution*, *4*, 609–613. <https://doi.org/10.1111/2041-210X.12069>
- Olsson, P., Lind, O., & Kelber, A. (2015). Bird colour vision: Behavioural thresholds reveal receptor noise. *Journal of Experimental Biology*, *218*, 184–193.<https://doi.org/10.1242/jeb.111187>
- Olsson, P., Lind, O., & Kelber, A. (2017). Chromatic and achromatic vision: Parameter choice and limitations for reliable model predictions. *Behavioural Ecology*, *29*, 273–282. [https://doi.org/10.1093/beheco/](https://doi.org/10.1093/beheco/arx133) [arx133](https://doi.org/10.1093/beheco/arx133)
- Osorio, D., & Vorobyev, M. (2005). Photoreceptor spectral sensitivities in terrestrial animals; adaptation for luminance and colour vision. *Proceedings of the Royal Society B*, *272*, 1745–1752. [https://doi.](https://doi.org/10.1098/rspb.2005.3156) [org/10.1098/rspb.2005.3156](https://doi.org/10.1098/rspb.2005.3156)
- Sanes, J. R., & Zipursky, S. L. (2010). Design principles of insect and vertebrate visual systems. *Neuron*, *66*, 15–36. [https://doi.org/10.1016/](https://doi.org/10.1016/j.neuron.2010.01.018) [j.neuron.2010.01.018](https://doi.org/10.1016/j.neuron.2010.01.018)
- Siddiqi, 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*. *Journal of Experimental Biology*, *207*, 2471–2485. <https://doi.org/10.1242/jeb.01047>
- Stevens, M., & Cuthill, I. C. (2006). Disruptive coloration, crypsis and edge detection in early visual processing. *Proceedings of the Royal Society B*, *273*, 2141–2147. [https://doi.org/10.1098/](https://doi.org/10.1098/rspb.2006.3556) [rspb.2006.3556](https://doi.org/10.1098/rspb.2006.3556)
- Troscianko, J., Skelhorn, J., & Stevens, M. (2017). Quantifying camouflage: How to predict detectability from appearance. *BMC Evolutionary Biology*, *17*, 7. [https://doi.org/10.1186/s12862-016-](https://doi.org/10.1186/s12862-016-0854-2) [0854-2](https://doi.org/10.1186/s12862-016-0854-2)
- 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. <https://doi.org/10.1111/2041-210X.12439>
- Troscianko, J., Wilson-Aggarwal, J., Stevens, M., & Spottiswoode, C. N. (2016). Camouflage predicts survival in ground-nesting birds. *Scientific Reports*, *6*, 19966. <https://doi.org/10.1038/srep19966>
- Van Belleghem, S. M., Papa, R., Ortiz-Zuazaga, H., Hendrickx, F., Jiggins, C. D., McMillan, W. O., & Counterman, B. A. (2018). Patternize: An R package for quantifying colour pattern variation. *Methods in Ecology and Evolution*, *9*, 390–398.<https://doi.org/10.1111/2041-200X.12853>
- VanDerWal, J., Falconi, L., Januchowski, S., Shoo, L., & Storlie, C. (2014). SDMTools: Species Distribution Modelling Tools: Tools for processing data associated with species distribution modelling exercises. R package version 1.1-221. Retrieved from [https://CRAN.R-project.](https://CRAN.R-project.org/package=SDMTools) [org/package=SDMTools](https://CRAN.R-project.org/package=SDMTools)
- Vorobyev, M., & Osorio, D. (1998). Receptor noise as a determinant of colour thresholds. *Proceedings of the Royal Society B*, *265*, 351–358. <https://doi.org/10.1098/rspb.1998.0302>
- White, T. E., & Kemp, D. J. (2016). Colour polymorphic lures target different visual channels in prey. *Evolution*, *70*, 1398–1408. [https://doi.](https://doi.org/10.1111/evo.12948) [org/10.1111/evo.12948](https://doi.org/10.1111/evo.12948)
- White, T. E., & Kemp, D. J. (2017). Colour polymorphic lures exploit innate preferences for spectral versus luminance cues in dipteran prey. *BMC Evolutionary Biology*, *17*(191), 1–10. [https://doi.org/10.1186/](https://doi.org/10.1186/s12862-017-1043-7) [s12862-017-1043-7](https://doi.org/10.1186/s12862-017-1043-7)
- White, T. E., Rojas, B., Mappes, J., Rautiala, P., & Kemp, D. J. (2017). Colour and luminance contrasts predict the human detection of natural stimuli in complex visual enviornments. *Biology Letters*, *13*, 20170375.<https://doi.org/10.1098/rsbl.2017.0375>
- Zhou, Y., Ji, X., Gong, H., Gong, Z., & Liu, L. (2012). Edge detection depends upon an achromatic channel in *Drosophila melanogaster*. *Journal of Experimental Biolology*, *215*, 3478–3487. [https://doi.org/10.1242/](https://doi.org/10.1242/jeb.070839) [jeb.070839](https://doi.org/10.1242/jeb.070839)

SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.

How to cite this article: Endler JA, Cole GL, Kranz AM. Boundary strength analysis: Combining colour pattern geometry and coloured patch visual properties for use in predicting behaviour and fitness. *Methods Ecol Evol*. 2018;9:2334–2348. <https://doi.org/10.1111/2041-210X.13073>