Imperfect transparency and camouflage in glass frogs.

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## Classification

## Keywords

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#### Abstract

Camouflage patterns prevent detection and/or recognition by matching the background, disrupting edges, or mimicking particular background features. In variable habitats, however, a single pattern cannot match all available sites all of the time and efficacy may therefore be reduced. Active color change provides an alternative where coloration can be altered to match local conditions, but again efficacy may be limited by the speed of change and range of patterns available. Transparency, on the other hand, creates high fidelity camouflage that changes instantaneously to match any substrate but is potentially compromised in terrestrial environments where image distortion may be more obvious than in water. Glass frogs are one example of terrestrial transparency, and are well known for their transparent ventral skin, through which their bones, intestines, and beating hearts can be seen. However, sparse dorsal pigmentation means these frogs are better described as translucent. To investigate whether this imperfect transparency acts as camouflage, we used in situ behavioral trials, visual modelling, and laboratory psychophysics. We found that the perceived luminance of the frogs changed depending on the immediate background, lowering detectability, and increasing survival when compared to opaque frogs. Moreover, this change was greatest for the legs, which surround the body at rest, and create a diffuse transition from background to frog luminance rather than a sharp, highly salient, edge. This passive change in luminance, without significant modification of hue, suggests a camouflage strategy, 'edge diffusion', distinct from both transparency and active color change.


## Significance

Transparency is one of the most intuitive forms of camouflage, where predators see straight through their prey as if it were not there. Glass frogs are a classic example of animal transparency and are well known for their transparent ventral skin, through which the internal organs can be clearly seen. The efficacy of transparency on land has, however, been controversial. We found that under natural conditions these frogs are better described as translucent, and that translucency acts as modifiable camouflage. Differences in the degree of translucency over the frog act to disguise the frog's outline and highlight the potential of
"edge diffusion" as a novel form of camouflage. Making glass frog camouflage distinct from both transparency and active color change.

## Introduction

Camouflage renders the visible invisible, or unrecognizable, with color patterns that blend into the background and/or disguise distinctive features (1,2). Matching visually complex or heterogeneous backgrounds, however, demands compromise, and so, many animals have developed ways of adapting their coloring to match the local conditions (3). Yet, changing color is still limited by the range of pigments that can be produced, the pattern resolution (i.e. chromatophore density), and the speed with which pigments can be moved or synthesized $(3,4)$. Thus, rather than attempting to mirror the background, it may be better to act as a window through which the background itself can be seen $(5,6)$.

Transparency is one of the most intuitive mechanisms of camouflage, allowing for instantaneous, high fidelity, background-matching regardless of the visual environment (6-8). Achieving transparency, however, requires the modification of a suite of tissues and structures to maximize light transmission, reduce scattering (9), avoid surface reflection, and minimize image distortion $(6,10)$.

Among aquatic species, transparency is relatively common and is facilitated by a variety of environmental factors, including the similarity in refractive indices between water and animal tissue, and a limited need for UV protective pigments $(6,9)$. Despite direct evidence of survival benefits being rare, indirect effects suggest that transparency does indeed act as camouflage. In transparent Daphnia spp. increasing the size of the eye (an obligate opaque structure) increases the risk of predation, and in areas where predatory fish are present eye size is significantly smaller than in fish-free environments (11, 12). Similarly, aquatic copepods reduce their expression of pigments when presented with predator cues (13), and the concentration of pigmentation is significantly lower in environments containing fish predators (14).

In contrast, on land camouflage via transparency has only been suggested in a few instances: namely certain lepidopterans, such as the glasswing butterflies (Nymphalidae), $(15,16)$ and the glass frogs (Centrolenidae) (17). Evidence is emerging that the transparent wings of these lepidopterans can decrease detectability compared to opaque wings, and that avian predators are less likely to capture transparent-
winged species $(15,16)$, although the pigmented wing edges and bright flash marks of certain species may reduce the efficacy of transparency (18). Conversely, despite glass frogs being well known for their transparent skin, through which the internal organs can be observed $(7,17)$, the camouflage hypothesis has been controversial, with doubts over i) the efficacy of transparency in terrestrial environments, and ii) the degree to which glass frogs can be described as transparent $(6,7)$.

In truth, whereas the ventral skin of glass frogs is often transparent, the dorsum is pigmented, albeit sparsely $(7,17)$. Thus, under natural conditions, unlike the truly transparent wing membranes of glasswing butterflies $(15,16,19)$, these frogs would be better described as translucent. This raises the question: what good is the glass frog's imperfect glass?

Whereas the role of transparency in camouflage is much discussed, translucency has received comparatively little attention. Indeed, the few published studies into translucency show that it can be used to increase, rather than decrease, conspicuousness in lizards. These lizards (Anolis sp. Dactyloiodae \& Draco sp. Agamidae) orientate their translucent dewlaps to be backlit by the sun so that the diffuse transmission of sunlight brightens their sexually selected signals $(20,21)$.

Here, we describe an alternative: where translucency facilitates a novel form of camouflage distinct from transparency and active color change. We examined the extent of glass frog translucency using models of predator vision, and assessed whether translucency affected the efficacy of camouflage through a computer-based detection experiment with human participants and an in situ predation study with the frog's wild predators (22).

## Results

Visual modeling. We took calibrated photographs of 25 adult Teratohyla midas in French Guiana and 30 Espadarana prosoblepon in Ecuador. All frogs were photographed under natural daylight conditions, on two backgrounds that differed in luminance and hue: dark green (fresh leaves) and bright white (Rite-in-the-Rain White All-Weather Copier Paper 8512-M, JL Darling LLC. Tacoma, WA, USA). The
photographs were taken with a Nikon DSLR camera (T. midas - D3200, E. prosoblepon - D7200) and AF-S DX NIKKOR 35 mm prime lens (Nikon Corporation, Tokyo, Japan), and each contained a ColorChecker Passport (X-Rite Inc. 2009. MI, USA) to enable color calibration and scaling. The frog's color on each background was then assessed using four visual models: tetrachromatic avian (23), trichromatic snake (24), dichromatic mammal (25), and trichromatic human (26, 27).

Firstly, we quantified perceived color change by comparing the accuracy with which a generalized linear mixed effects model (R package lme4 (28) in R 3.5.1 (29)) could discriminate between the frogs on the two backgrounds. The higher the classification accuracy, the greater the change between the backgrounds. Classification accuracy was characterized using the Area Under the Curve (AUC) of Receiver Operated Characteristic (ROC) curves ( R package $p R O C(30)$ ), following a common grading system: 1.0-0.9 $=$ excellent, $0.9-0.8=$ good, $0.8-0.7=$ fair, $0.7-0.6=$ poor, and $0.6-0.5=$ fail $(31)$.

We found that, overall, each visual model was fair (T. midas) or poor (E. prosoblepon) at discriminating the frogs on the white background from the frogs on the leaves (Fig. 1; see SI Appendix, Tables S1, S2). However, accuracy was not equal across the frog: each model was either poor or failed to detect any change in luminance or hue from the body but was good at distinguishing the legs. This indicates that although the frog's body largely retained its appearance atop the two different backgrounds, the legs changed significantly. This change was largely the result of a shift in achromatic contrast rather than hue and, although the frogs' base color did not exactly match this sample of leaves, the perceived luminance shifted towards that of the immediate background (Fig. 1; see SI Appendix, Tables S1, S2).

Detection. Secondly, to evaluate whether the changes described in the visual modelling could enhance camouflage, without the potential confound of neophobia, we ran an ex situ computer-based detection experiment with human participants. It is important to note the existence of differences in visual perception between species and to be cautious when applying human-derived data to other animals. There are, however, deep underlying similarities in visual processing between species (32) and many studies
show a strong correspondence in what makes prey detectable between humans and wild avian predators (e.g. 16, 26, 33, 34).

We created frog-shaped stimuli using a standardized template of $T$. midas in its diurnal resting posture. We extracted the mean luminance and hue of the body and the degree of translucency from the legs for each of the 25 T. midas used for the visual modelling. We then used these data to create treatments A-D where we manipulated the distribution of the translucent elements. Treatment E used the mean colors from randomly selected green leaves photographed in situ. This created five treatments: A - natural pattern, translucent legs and opaque body; B - reversed pattern, opaque legs and translucent body; C fully translucent; D - fully opaque; and E - opaque leaf color. Twenty human participants were tasked with finding the frogs as quickly as possible against a background made up of calibrated photographs of green leaves. Reaction time (RT) was log transformed and analyzed with a general linear mixed effects model and detection accuracy (DA) was analyzed with a binomial generalized mixed-effects model ( R package lme4 (28) in R 3.5.1 (29)).

We found a significant effect of treatment on the time taken to detect the frogs $(\chi 2=7310.30, \mathrm{df}=4, \mathrm{p}<$ 0.001; Fig. 2 Left \& Middle) and for detection accuracy ( $\chi 2=430.41, \mathrm{df}=4, \mathrm{p}<0.001$ ). Pairwise tests of a priori interest (R package multcomp (35)) showed that the natural pattern was significantly harder to detect than the entirely opaque frog ( $\mathrm{A}-\mathrm{D}: \mathrm{RT} \mathrm{z}=-2.95, \mathrm{p}=0.012, \mathrm{DA} \mathrm{z}=0.34, \mathrm{p}=0.991$ ), and that there was no significant difference between the three patterns that included translucent elements $(A-B$ : $R T z=-1.26, p=0.542, D A z=0.11, p=1.00 ; A-C: R T z=0.98, p=0.733, D A z=0.99, p=0.718)$. In line with the visual modelling, we also found that the frog's opaque base color was detected significantly more quickly and accurately than the opaque background sample ( $\mathrm{D}-\mathrm{E}: \mathrm{RT} \mathrm{z}=26.75, \mathrm{p}<0.001$, $\mathrm{DA} \mathrm{z}=$ 13.72, $\mathrm{p}<0.001$ ).

Survival. Finally, to test whether these effects corresponded to survival advantages under natural conditions, we ran an in situ predation study with model frogs and wild avian predators in Ecuador. We
used green gelatin to create two treatments: translucent and opaque ( $\mathrm{n}=180$ per treatment). These models were placed on vegetation, in 19 independent blocks, along riparian transects. Over the course of 72 h , we recorded how quickly each treatment was predated by wild predators (mainly birds, based on the frequent presence of beak marks) and we analyzed mortality with a mixed effects Cox model (R package coxme (36)). We found a significant effect of treatment $\left(\chi^{2}=14.81, \mathrm{df}=1, \mathrm{P}<0.001\right)$ and that translucent models had a significantly lower mortality rate than opaque frogs $(z=-3.70, p<0.001$; Fig. 2 Right $)$.

## Discussion

Taken together our data suggest that glass frog translucency does act to provide ecologically relevant camouflage. We found that, without any changes in pigmentation, the perceived luminance of the frogs changed depending on the background, making each frog a closer match to its immediate surroundings. Unlike true transparency, however, color change was largely restricted to luminance rather than hue. The frogs therefore appeared green at all times but seemed to brighten and darken depending on the substrate against which they were viewed. We further found that this translucency resulted in glass frogs being more difficult for human participants to detect and less likely to be attacked by wild predators than opaque frogs.

Being transparent, like the membranous wings of glasswing butterflies, allows whole background structures to be visible through the organism $(15,16)$. Translucent glass frogs, on the other hand, retain their green hue regardless of the background and background patterning does not show through the frog. The glass frogs' natural history does, however, provide a means to bridge the potential efficacy gap between translucency and transparency. Under natural conditions glass frogs remain stationary during daylight hours, resting flat against leaves, with their legs tucked to the sides of their body. By predominantly resting on living green leaves, the frogs' natural background does not include complex visual textures and will vary much more in luminance than hue.

This limits the necessity for more dramatic changes in hue, reduces the need for dynamic pattern matching, and allows the frogs' green pigmentation to act as generalist background-matching camouflage. This generalist base color is then calibrated by diffuse light transmission to passively increase camouflage against the immediate background. This camouflaging effect would apply to both diurnal and nocturnal predators, and was consistent across a diverse range of potential predator visual systems that differ greatly in color perception. Translucency, therefore, allows perceived luminance to change, whilst continuing to allow pigmentation to screen any obligate opaque structures (e.g. thick or complex body tissues) from view and, potentially, from harmful UV radiation.

Moreover, the change in perceived luminance was greater on the frogs' legs than on the body, that remained principally the same brightness regardless of the background. As such, when at rest and the relatively opaque body is surrounded by the more translucent legs, the frog becomes a diffuse gradient from substrate to dorsal color rather than producing a sharp, high contrast, outline. Visual systems are particularly sensitive to high contrast edges, especially in the luminance domain (32,37), and low intensity, graduated boundaries have been demonstrated to make artificial camouflaged targets more difficult to detect (38).

Camouflage strategies are defined by the perceptual processes that the pattern exploits rather than the features of the pattern per se (2). Transparency is most frequently considered to produce a form of camouflage akin to background matching, with the transparent animal blending into its immediate surroundings (6). Transparency and translucency may, however, also facilitate other defensive strategies. Many species of clearwing moths (e.g. Sesiidae, Lepidoptera) are Batesian mimics and their transparent wings match those of venomous wasps (e.g. Vespidae, Hymenoptera). Similarly, many species of katydid (Tettigonidae, Orthoptera) and leaf insect (Phyllidae, Phasmatodea) masquerade as leaves with incredibly precise mimicry that often includes translucent elements representative of leaf damage.

Furthermore, the irregular translucent elements found on the otherwise opaque wings of many moths may act as disruptive camouflage, with translucent components blending into the background to leave unrecognizable false edges $(33,39)$. The 'edge diffusion' observed in glass frogs represents a form of camouflage conceptually linked with disruptive coloration (40), but whereas disruption reduces the relative contrast of the target's outline, edge diffusion reduces the absolute intensity of the edge.

As such, the translucent appearance of many glass frogs acts to provide camouflage in a manner conceptually distinct from both true transparency and active color change. Rather than allowing the background to be directly seen, diffuse light transmission through the frog adjusts a generalist camouflage pattern to more closely match the immediate background. This change in perceived luminance then transforms the frogs' salient high-intensity outline into a less conspicuous graduated boundary. Thus, the imperfect glass of the glass frog provides effective camouflage, disguising the frogs' outline, and blending the frog and the leaf more smoothly together.

## Materials and Methods

Image collection. Teratohyla midas $(\mathrm{n}=25)$ were encountered in the rainforest surrounding the SautPararé camp of the Nouragues Natural Reserve, French Guiana (December 2014 - January 2015), and Espadarana prosoblepon $(\mathrm{n}=30)$ were sampled along streams near the town of Mindo, Pichincha Province, Ecuador (May - July 2019). In order to assess the degree to which translucency produces color change, each frog was photographed on two backgrounds in a random order and in quick succession (to remove the influence of any change in pigmentation): a sheet of white waterproof paper (Rite-in-the-Rain White All-Weather Copier Paper 8512-M, JL Darling LLC. Tacoma, WA, USA), and fresh dark green leaves collected from the same trees on which glass frogs were seen calling (Fig. 1). In addition, for the detection experiment we photographed 10 leaves from each of 10 tree species $(\mathrm{n}=100)$ found in the same location as $T$. midas were observed to be calling.

Each frog, and each leaf, was photographed from directly above, under natural daylight conditions, with a Nikon DSLR camera (T. midas - D3200, E. prosoblepon - D7200) and AF-S DX NIKKOR 35 mm prime lens (Nikon Corporation, Tokyo, Japan). To enable color calibration and scaling, all images contained a ColorChecker Passport (X-Rite Inc. 2009. MI, USA). Frogs were then released at the precise location from where they were collected.

Visual modelling. Each photograph was linearized, size-scaled, and calibrated in accordance with the ColorChecker Passport (41). The pixels corresponding to the frog (both legs and body) and the background (white background vs green leaf) were specified and saved separately in MATLAB 2018b (The MathWorks Inc., Natick, MA, USA).

Image analysis used four visual models corresponding to three potential visual predators as well as human vision: tetrachromatic avian LMS, trichromatic snake LMS, dichromatic mammalian LS, and trichromatic human $L * a * b *(26,42,43)$. Human vision was included to allow intuitive interpretation of visual and detection data. Humans are trichromatic, with peak absorption ( $\lambda_{\max }$ ) of 564 nm (LWS), 534 nm (MWS), and 420 nm (SWS) (27). Human $L^{*} a * b *$ is an international standard color space derived from psychological testing (CIELAB, 1976); however, as an equivalent visual model for non-human species is not available avian LMS, snake LMS, and mammalian LS models used relative cone capture rates.

Cone capture rates were used to generate a three-dimensional color space which categorized perceivable colors along orthogonal opponent channels in a similar manner to $\mathrm{L}^{*} \mathrm{a}^{*} \mathrm{~b}^{*}$ : luminance ( L ), and the opponent channels red-green (rg) and yellow-blue (yb). Although opponent processes have not been characterized for the non-human visual systems that we model, these orthogonal color channels represent an efficient way to encode visual information and, as such, a biologically plausible estimate of visual contrast (44).

To check for UV reflectance, we photographed E. prosoblepon under natural daylight conditions with a UV-sensitive, full-spectrum quartz converted, Canon EOS 7D (Canon Inc., Tokyo, Japan) and NIKKOR

EL 80 mm lens (Nikon Corporation, Tokyo, Japan), as well as appropriate filters for human visible and infrared light. Each image contained two luminance standards to enable calibration: $77 \%$ and $10 \%$ reflectance (Labsphere, North Sutton, NH, USA). UV irradiance is low below the forest canopy, and we found there to be little UV reflectance from either the frog or its background (see SI Appendix, Fig. S1). Consequently, the UV channel will not contain any information and was excluded from our visual models (26). The mammalian visual model lacks an MWS cone and so was modelled in two dimensions: L and yb (26, 34, 42).

Avian LMS used the violet-sensitive tetrachromatic vision of the Indian peafowl (Pavo cristatus Galliformes) with $\lambda_{\max }$ at 605 nm (LWS), 537 nm (MWS), 477 nm (SWS), 432 nm (VS), and luminance measuring double cones at 567 nm (D) (23). L was generated from the response of the D cone, rg was the relative stimulation of the LWS to MWS cone, and yb was the relative stimulation of the LWS and MWS to the SWS cone.

Snake LMS used the UV-sensitive trichromatic vision of the whip snake (Masticophis flagellum Colubridae) with $\lambda_{\max }$ at 561 nm (LWS), 458 nm (MWS), and 362 nm (UV) (24). L was calculated from the stimulation of the LWS cone, rg from the relative stimulation of the LWS to MWS cone, and yb from the combined stimulation of the LWS and MWS to the UV cone.

The mammalian LS model used the dichromatic vision of the domestic ferret (Mustela putorius Mustelidae) with $\lambda_{\max }$ of 558 nm (LWS) and 430 nm (SWS) (25). L was calculated from the LWS cone, and yb was the ratio of the LWS to the SWS.

Image analysis. Chromatic information was analyzed using a generalized linear mixed effects model ( R package lme4 (28)) in R 3.5.1 (29). Our approach has parallels with the Vorobyev and Osorio receptor-noise-limited model (45), but rather than comparing pairs of pixels, we are interested in determining the separability of two clouds of points within the color space. To avoid overfitting, we used leave-one-out cross validation (31), the unit of omission being the frog rather than the pixel (26). The degree of color
change was quantified as the discrimination accuracy between the frog on the white background and the frog on the leaf using the Area Under the Curve (AUC) of Receiver Operated Characteristic (ROC) curves (R package $p R O C(30))$. The AUC represents the probability of correct identification with 1.0 indicating perfect discrimination and 0.5 indicating chance. We followed a common grading system: 1.0$0.9=$ excellent, $0.9-0.8=$ good, $0.8-0.7=$ fair, $0.7-0.6=$ poor, and $0.6-0.5=$ fail $(26,31)$. The higher the AUC the greater the change between the two backgrounds.

Detection stimuli. To create the experimental backgrounds, each leaf photograph $(\mathrm{n}=100)$ was calibrated in accordance with the ColorChecker Passport (41), and the whole leaf was cropped from the image and saved separately using MATLAB 2018b. The experimental backgrounds were generated by layering all 100 leaves in a random sequence, with leaf orientation and position randomly selected.

For the frog stimuli we used the colors of the 25 T. midas photographed for the visual modelling and a standardized template in the shape of $T$. midas in a diurnal resting posture, flat to the leaf with its legs to the side of its body. As the color of the frogs' bodies did not change significantly between backgrounds, we calculated the mean color of each frog's body and used these as the base colors for the experimental stimuli.

We manipulated frog color to generate five treatments. Treatments A, B, C, and D used the mean colors of the frogs: A was a frog with translucent legs and an opaque body (as found in the natural condition); B had a translucent body and opaque legs (reverse of the natural condition); C was a frog with a translucent body and translucent legs; D was an opaque frog; and E represented random sample background matching where each replicate was an opaque frog with the mean color of a randomly selected leaf.

Translucency was simulated by modifying the alpha data of the frog image file (opaque $\alpha=1.00$; translucent $\alpha=0.85$ ) in MATLAB 2018b. The ecological validity of the approach was assessed by repeating the visual modelling protocol on the experimental stimuli. We generated stimuli using the mean colors of each of the 25 frogs and translucency ( $\alpha$ ) values set at $5 \%$ increments between $100 \%$ (opaque)
and $75 \%$ (translucent). Each stimulus was digitally rendered both against a white background and on the experimental (leaf) backgrounds. One thousand pixels corresponding to the target area were then randomly selected, extracted, and converted into $L^{*} a^{*} b^{*}$ color space. We ran a generalized linear mixed effects model with leave-one-out cross validation using the mean human $L * a * b *$ model responses for each frog. We compared the frog color between the two backgrounds for each level of transparency, as well as for the mean color for the real frogs' legs on the white and leaf backgrounds. We found that an $\alpha$ of 0.85 resulted in a degree of color change (AUC) that most closely matched the legs of $T$. midas (AUC: Frog legs $=0.95 ; \alpha 100 \%=0.53 ; \alpha 95 \%=0.74 ; \alpha 90 \%=0.85 ; \alpha 85 \%=0.97 ; \alpha 80 \%=1.00 ; \alpha 75 \%=$ 1.00 ), and so this was used for the detection experiment.

Detection protocol. Each participant ( $\mathrm{n}=20$, with 'normal or corrected to normal' vision) was presented the five stimuli generated from each of the 25 frogs $(\mathrm{n}=125)$ on a 13 " MacBook Air (Apple Inc., Cupertino, CA, USA, 2018). Each frog was paired with a uniquely generated background and was placed at a random location with a randomly selected rotation. Each participant therefore saw a unique set of frog-background combinations and these stimuli were presented in a separately randomized sequence (see SI Appendix, Fig. S2). We recorded reaction time and detection accuracy, with all clicks on the frog, within a circle centered on the frog with a diameter equal to the frog's length, classified as correct.

Detection analysis. Reaction time was log-transformed and analyzed with a general linear mixed effects model and detection accuracy was analyzed with a binomial generalized mixed-effects model (R package Ime4 (28)). Pairwise comparisons, of a priori interest, were designed to test particular hypotheses ( R package multcomp (35)). We compared treatment A to treatments B, C, and D to assess whether the amount and distribution of translucency affected detectability, and treatment D to E to test whether the frogs' pigmented color was as successful as random sample background matching. As the number of comparisons is equal to the number of degrees of freedom $p$ values were not adjusted.

Survival protocol. The survival experiment was conducted along riparian transects near the town of Mindo (Pichincha Province, Ecuador), a site where the glass frog Espadarana prosoblepon was common. We used a randomized block design with two treatments: translucent and opaque. The models were placed out in 19 independent blocks, containing 10 of each treatment, aside from blocks 12 and 13 which, due to damaged models, each contained 5 of each treatment ( $n=180 /$ treatment). Models were tied to green leaves along each transect spaced apart as to be independent of one another. Evidence of predation was recorded at 24,48 , and 72 h after the block was placed out. Relative mortality was assessed with a mixed effects Cox model (R package coxme (36)), with evidence of predation recorded as full events, and missing, rain damaged, or surviving models recorded as censored values.

Model frogs were made from a mixture of water and unflavored sugar-free gelatin powder (Supermaxi, Corporación Favorita C.A., Quito, Ecuador). The solution naturally dried translucent with a slight yellow tint, and so we added a small amount of corn starch to modify opacity and green food coloring (La Reposterita, Leapan, Quito, Ecuador) to control color. To make 12 frogs, we first added the food coloring (translucent $=0.05 \mathrm{ml}$, opaque $=0.10 \mathrm{ml}$ ) to 380 ml of boiling water. We then mixed 56 ml of green boiling water into 15 g of gelatin powder plus a small amount of corn starch (translucent $=0.20 \mathrm{~g}$, opaque $=0.50 \mathrm{~g}$ ). The solution was then poured into a silicone mold (Alumilite ${ }^{\mathrm{TM}}$ Amazing Mold Rubber, Alumilite Corporation, Kalamazoo, MI, USA) created using a 3D print of a tree frog (Phyllomedusa tomopterna) in a resting posture (Digital Life Project, Department of Biology, University of Massachusetts Amherst, Amherst, MA, USA), which was scaled to match the body length ( 25 mm ) of $E$. prosoblepon. Models were left to solidify overnight and attached to leaves the following morning.

To assess whether the two treatments differed in opacity to potential predators we replicated the image analysis protocol with the avian visual model (outlined above). We photographed the models ( $\mathrm{n}=5$ / treatment) on two backgrounds, white waterproof paper and a dark green leaf, under natural daylight conditions. Each image contained a ColorChecker Passport and was taken with a Nikon D7200 DSLR camera AF-S DX NIKKOR 35 mm prime lens. Plotting the model response shows that hue (red-green
and yellow-blue) was similar between the two models and was largely stable across the two backgrounds. Perceived luminance, however, shifted depending on background, in a similar manner to the real frogs, such that the translucent models were a closer match to the background luminance (see SI Appendix, Fig. S3).

Data deposition. Raw data are available in the University of Bristol Research Data Repository (doi: 10.5523/bris.3qnujjtjg4v4z2cztb4h882hh3).

Author Contributions. The experiments were conceived by JBB, CM, JY, JNP, NES-S and ICC. JBB, CM, HA, BLM and JY collected the data in the field, JBB performed the detection experiment, and JBB and ICC implemented the image analysis. JBB wrote the first draft of the manuscript with subsequent input from all authors.

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Figures


Fig. 1. Visual modelling of color change (avian visual model). A) Espadarana prosoblepon on the white background. B) E. prosoblepon on the leaf background. C \& D) Teratohyla midas and E. prosoblepon respectively: the response of each channel of the avian visual model to the leaf background (green), the whole frog on the leaf (orange), and the whole frog on the white background (black). The frogs' luminance, more so than hue, shifts towards that of the immediate background. E) ROC curves from the avian visual model for the whole frog (left), the frog's body (middle), and the frog's legs (right) - for both species color change is strongest on the frogs' legs. AUC: $0.5-0.6=$ fail, $0.6-0.7=$ poor, $0.7-0.8=$ fair, $0.8-0.9=$ good, and $0.9-1.0=$ excellent.


Fig. 2. Reaction time and relative mortality. Left: time taken for human participants to find frogs with different levels of translucency (means and $95 \%$ CI): stimuli using the base color of Teratohyla midas: A - natural pattern, translucent legs and opaque body; B - reverse pattern, opaque legs and translucent body; C - fully translucent; D - fully opaque. There was no significant difference between the natural pattern and frogs with some level of translucency (A-B \& A-C), but the frog's natural level of translucency was significantly harder to detect than a fully opaque frog (A-D). Middle: comparison between the mean color of each the frog's body (D) to random sample background matching (E). The frogs' mean colors did not match the background (see Fig. 1), and random sample background matching was significantly harder to detect than the mean color of each frog. Right: predation rate of model frogs (odds ratios with $95 \% \mathrm{CI}$ in relation to the opaque frog). The translucent frog had a significantly lower mortality rate (higher survival) than the opaque frog.

## Supplementary Information for

Imperfect transparency and camouflage in glass frogs.

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Fig. S1. Espadarana prosoblepon photographed in human visible RBG (left) and ultraviolet (right). There is minimal UV-reflectance from either the frog or the leaf, which appear dark in the UV image.


Fig. S2. Detection experiment example stimuli (frog shaped targets circled). Left - treatment A, natural pattern, translucent legs and opaque body, and Right - treatment B, reversed pattern, opaque legs and translucent body. Each frog was presented on a unique background of sympatric leaves. Backgrounds were generated by layering leaf photographs with sequence, location, and rotation independently randomised for each stimulus. Similarly, each frog was placed at a random location and with a randomly selected rotation.


Fig. S3. Survival experiment example stimuli and avian visual modeling. Example stimuli: A) translucent model on white background, B) translucent model on the leaf background, C) opaque model on white background, and D) opaque model on leaf background. Avian visual modeling: E) translucent model and F) opaque model (leaf = green, model on leaf = orange, and model on white = black). For both treatments there is a large overlap in hue (red-green and yellow-blue) between the model on each background and the leaf. However, the luminance of the translucent model shifts more dramatically between backgrounds than the opaque model, to become a better match to the background luminance.

