Wing transparency in butterflies and moths: structural diversity, optical properties, and ecological relevance

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Abstract. In water, transparency seems an ideal concealment strategy, as testified by the variety of transparent aquatic organisms. By contrast, transparency is nearly absent on land, with the exception of insect wings, and knowledge is scarce about its functions and evolution, with fragmentary studies and no comparative perspective. Lepidoptera (butterflies and moths) represent an outstanding group to investigate transparency on land, as species typically harbor opaque wings covered with colored scales, a key multifunctional innovation. Yet, many Lepidoptera species have evolved partially or fully transparent wings. At the interface between physics and biology, the present study investigates wing transparency in 123 Lepidoptera species (from 31 families) for its structural basis, optical properties, and biological relevance in relation to visual detection (concealment), thermoregulation, and protection against UV. Our results suggest that transparency has likely evolved multiple times independently. Efficiency at transmitting light is largely determined by clearwing microstructure (scale shape, insertion, coloration, dimensions, and density) and macrostructure (clearwing area, species size, or wing area). Microstructural traits, scale density and dimensions, are tightly linked in their evolution, with different constraints according to scale shape, insertion, and coloration. Transparency appears highly relevant for concealment, with size-dependent variations. Links between transparency and latitude are consistent with an ecological relevance of transparency in thermoregulation, and not so for protection against UV radiation. Altogether, our results shed new light on the physical and ecological processes driving the evolution of transparency on land and underline that transparency is a more complex coloration strategy than previously thought.

Key words: Lepidoptera; microstructure; structural strategy; thermoregulation; transparency; UV protection; vision.

INTRODUCTION

Following the invisibility myth, transparency seems an ideal camouflage strategy: being "hidden in plain sight" works whatever the background, from all viewpoints and irrespective of behavior (Cuthill 2019). The large "success story" of transparency in water as a protection against predators (especially in pelagic habitats where there is nowhere to hide) is attested by its broad phylogenetic distribution since transparency spans seven phyla

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including Arthropoda, Mollusca, Annelida, Chordata, and Cnidaria (Johnsen 2001). By contrast, transparency is nearly absent on land, and it is almost entirely confined to insect wings (one phylum). This contrast can be explained by physical factors: compared to water, larger refractive index mismatch between air and biological tissues produces higher light reflection that ruins invisibility (Johnsen 2001). In addition, greater ultraviolet (UV) radiation on land imposes greater selection on UV protection often through light absorption by pigments.

Research in transparent aquatic organisms has shown a role for concealment from visually hunting predators, and predators' ability to break this camouflage (e.g., Tuthill and Johnsen 2006). As underlined by Johnsen (2014), many questions are left unanswered about

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transparency: the structural bases of transparency (Bagge 2019), its genetic and developmental pathways, its functional roles in vital functions like thermoregulation and potential trade-offs with optics, and the selective pressures driving its evolution and its design. Comparative studies at a broad interspecific level are absent, yet they are essential to better understand the links between structural diversity and adaptive functions of transparency. The lack of studies is even more crucial for transparency on land, where knowledge is scarce, with fragmentary monographic studies by physicists using bioinspired approaches, based on transparent wing antireflective, hydrophobic and antifouling properties (e.g., Deparis et al. 2014, Liu et al. 2016, Elbourne et al. 2017).

Within insects, Lepidoptera represent an outstanding group to explore these questions. While most insects harbor transparent wings, Lepidoptera are characterized by wings covered with scales. Scales are chitin extensions that are usually long and large and that often contain pigments or structures that interact with light, thereby producing opaque color patterns (e.g., Stavenga et al. 2014). Wings covered by scales represent an evolutionary innovation involved in many functions such as antipredator defenses (camouflage, deflection, aposematism, e.g., Stevens et al. 2008), communication (Kemp 2007), thermoregulation (Miaoulis and Heilman 1998, Berthier 2005, Krishna et al. 2020), and water repellency (Wagner et al. 1996, Wanasekara and Chalivendra 2011). In this opaque world, many species from different families have evolved partially or totally transparent wings. The handful of relevant biophysical studies in Lepidoptera suggests that an important aspect of structural diversity-having a nude membrane (completely lacking scales) or with scales of various morphologies, insertion angle on the wing membrane, and colorationmay underlie the occurrence of transparency in many Lepidoptera lineages (Yoshida et al. 1997, Hernandez-Chavarria et al. 2004, Berthier 2007, Goodwyn et al. 2009, Wanasekara and Chalivendra 2011, Stavenga et al. 2012, Siddique et al. 2015). This opens questions regarding the phylogenetic extent of transparency, the diversity and evolution of optical properties and of the underlying structures, the existence, if any, of structural constraints on transparency, and the ecological relevance of transparency on land.

Using a large data set comprising 123 clearwing Lepidoptera species encompassing 31 families, we examine the phylogenetic distribution of transparency in Lepidoptera. Theoretical and empirical studies on aquatic organisms suggest there exist several structural and optical routes to transparency (Johnsen 2001), highlighting the interest of quantifying structural diversity in terrestrial organisms.

First, we aim to quantify the extent of structural diversity for transparency in Lepidoptera at various scales: macrostructure (wing size, wing area, clearwing area, and proportion of clearwing area) and microstructure (presence of scales, type, insertion, and coloration). We assess to what extent structural features are conserved across the phylogeny, and whether some of the features are involved in correlated evolution, which can help us reveal evolutionary constraints. We quantify the investment in transparency, relative to what is invested in the opaque scales.

Second, we aim to examine the potential impact of macro- and microstructure on optical properties. We assess to what extent scale type, insertion, coloration, and density influence optical transparency, and we hypothesize that lower wing membrane coverage by scales (smaller surface and/or in lower density) should result in higher levels of optical transparency.

Third, we test the ecological relevance of transparency for visual detection, and especially for concealment. The prominent role of transparency in camouflage shown so far (Johnsen 2014, Arias et al. 2019, 2020, McClure et al. 2019) suggests that visually hunting predators may exert an important selective pressure on the evolution of transparency on land as they can prey on flying or resting diurnal prey as well as on nocturnal resting prey. We hypothesize that higher levels of optical transparency should yield lower visual contrasts with the background as perceived by natural predators. Given the principle of transparency (giving visual access to what is behind, under any light), this should apply to both visual systems found in birds and in various visual conditions of ambient light and background.

Fourth, we test the ecological relevance of transparency for thermoregulation. The thermal melanism hypothesis states that individuals from colder places (at higher latitudes or altitudes) gain thermal benefits from being more strongly pigmented as radiation absorption helps heating (Bogert 1949). This hypothesis has received support from comparative analyses at large taxonomical and geographical scales (e.g., Zeuss et al. 2014, Xing et al. 2018, Stelbrink et al. 2019) and from analyses at a species level in Lepidoptera (e.g., Colias species in Ellers and Boggs [2004]) showing that thermal benefits come from melanization of the proximal wing, close to the body. Recent large-scale comparative analyses in opaque butterflies have shown that body and proximal wing coloration correlates to climate in the nearinfrared (700-1,100 nm) range but not so below 700 nm where not only thermoregulation but also vision operates (Munro et al. 2019). Hence, wavelength absorption should be higher at high latitudes compared to low latitudes, especially in the infrared range where most of the thermal radiative exchanges take place.

Fifth, we finally test the ecological relevance of transparency for UV protection. Exposure to highly energetic and penetrating UVB (280–315 nm) and UVA (315– 400 nm) radiations has detrimental effects on physiology, fecundity, and survival in terrestrial living organisms (e.g., in insects Zhang et al. 2011). The absorption of UV radiation by pigments is efficient to prevent UV harmful effects, as shown in transparent fish larvae, where exposure to UV light rapidly induces melanization as a UV protection (Mueller and Neuhauss 2014). Levels of UV radiation are higher at low latitudes than at higher latitudes (Beckmann et al. 2014). Hence, if UV protection is a strong selective force, absorption of UV radiation is expected to be more important at low latitudes. Moreover, pathogens are more important in warmer and more humid regions, hence in lower latitudes. Stronger melanization at lower latitudes has been shown to provide protection against pathogens (True 2003). Hence, a higher UV absorption at lower latitudes would also potentially protect against pathogens.

Testing the potential role of transparency in thermoregulation or UV protection generates two mutually exclusive hypotheses. In optics, the light received by an object can be either transmitted, reflected, or absorbed. Variation in light transmission can indicate variation in absorption if reflection levels are maintained at similar levels. We hypothesize that species with fully or partially transparent wings living at increasing distance from the equator should transmit less light (absorb more) through their wings (at least in the near-infrared range) if transparency plays a role in thermoregulation, as light absorption by pigments is a way of gaining energy. Conversely, if transparency plays a role in protection against UV, species with fully or partially transparent wings living at increasing distance from the equator are expected to absorb less in the UV range, and thus to transmit more. The present study thus addresses for the first time the links between structure and optics at a broad phylogenetic scale to understand the "small success story of transparency on land," its evolution, and putative functions in relation to camouflage and thermoregulation.

Methods

Specimens and ecological data

We looked for clearwing species in the Lepidoptera collection of the French Museum of Natural History, by sampling as many families as possible to increase the power of comparative analyses. We found clearwing species in 31 out of the 124 existing Lepidoptera families (Supplementary file File1_SpeciesList.txt, see families in Appendix S1: Fig. S1) and gathered a total of 123 species. We included transparent and translucent species (i.e., when a text can be read when placed just behind but not from a distance, like a tracing paper) in this data set as they were likely on a continuum of light transmission and it was important to see whether this continuum entailed qualitative or quantitative changes. We took one specimen per species (see Fig. 1 for some examples; sex information was unavailable for most specimens and thus discarded). Those specimens were often unique or precious, which prevented us from conducting destructive measurements. There were 77 specimens for which labels specified exact collect location that could be tracked down to GPS coordinates (see Appendix S1: Fig. S2).

Structure measurements

Museum specimens were photographed using a D800E Nikon camera (Nikon) equipped with a 60 mm lens, placed on a stand with an annular light. Photos were then analysed using ImageJ (Schneider et al. 2012) to extract descriptors of wing macrostructure: wing length (expressed in mm, see Appendix S1: Fig. S2 for its phylogenetic distribution in the forewing), wing area (mm²), and clearwing area (the surface of transparent area in mm²), for the forewing and hindwing separately. We defined and computed proportion of clearwing area as the ratio clearwing area/wing area, i.e., the proportion of the total wing area occupied by transparency (see Appendix S1: Fig. S2 for its phylogenetic distribution in the forewing).

Museum specimens were also photographed with microscopes (Zeiss Stereo Discovery V20 [Zeiss] and a high-resolution Keyence VHX-5000 [Keyence]) to get detailed images of the dorsal side of transparent and opaque zones of each wing. These images were analysed using the built-in measurement software to describe wing microstructure. For each wing separately, we measured scale density (number of scales per mm², as zero for nude membrane, whatever the density of scale sockets, see Appendix S1: Fig. S2 for its phylogenetic distribution in the forewing), length (μ m), and width (μ m). We computed scale surface (in μ m²) as the product of length by width, and scale coverage as the product of scale surface (expressed in mm²) by scale density.

As shown from examples in Fig. 1, the transparent zone showed diversity in scale morphological type, in scale insertion, and in scale coloration. (1) In scale morphological type (hereafter referred to as type): no scale (N), bifid or monofid "hair-like" piliform scales (P), scales with any shape other than piliform, hereafter called lamellar scales (L) as in Kristensen et al. (2003), presence of both lamellar scales and piliform scales (PL). (2) In scale insertion on the membrane: erect (E) or flat (F) scale, or unknown insertion (U, when scales were absent). (3) In scale coloration: colored (C) or transparent (T, when scales were partly or totally transparent), or unknown coloration (U, when scales were absent).

We define as *structural strategy* the combination of the following traits: scale type, insertion, and coloration (examples in Fig. 1). The name of a structural strategy is thus composed of the abbreviated letters for scale type, insertion, and coloration, in this order. The NUU strategy (hereafter called N strategy; e.g., *Senecauxia coraliae* or *Chalioides ferevitrea*) has no scales, unknown insertion, and unknown coloration. PEC (e.g., *Pseudohaetera hypaesia*) is based on colored erect piliform scales. PFC (e.g., *Attacus atlas*) is based on colored flat piliform scales. LEC (e.g., *Isostola flavicollaris*) is based on erect colored lamellar scales while LET (e.g., *Oxynetra semihyalina*) is based on erect transparent lamellar scales. LFC (e.g., *Cressida*



FIG. 1. Examples of structural strategies in Lepidoptera. A structural strategy is defined as the combination of scale type, insertion, and coloration. Type is P, piliform scales, or L, lamellar scales (the combination of piliform and lamellar scales was assimilated to lamellar scales); insertion is E, erect, or F, flat; and coloration is C, colored, or T, transparent. The N strategy has no scales, no insertion, and no coloration. Bar scales are 1 cm for entire specimens (left columns), 100 µm for microscopic imaging (right columns and scale details). Species are the erebid *Senecauxia coraliae* (SE), the psychid *Chalioides ferevitrea* (CH), the nymphalid *Pseudohaetera hypaesia* (PS), the saturniid *Attacus atlas* (AT), the erebid *Isostola flavicollaris* (IS), the hesperid *Oxynetra semihyalina* (OX), the papilionid *Cressida cressida* (CR), and the crambid *Diaphania unionalis* (DI). Notice that we chose not to illustrate the PL category (combination of both scales and piliform scales) because, due to their rarity in the data set, they were pooled with the L category in subsequent analyses. Symbols for lamellar and piliform are meant as illustrative and do not represent exact scale morphologies encountered in the documented species.

cressida) is based on lamellar scales flat on the wing membrane and colored, while LFT (e.g., Diaphania unionalis) is based on lamellar scales flat on the wing membrane and transparent. The structural strategies based on a combination of lamellar scales and piliform scales (starting with PL), are not illustrated because of their rarity. PL strategies were pooled with strategies based on lamellar scales (starting with L) of their corresponding insertion and coloration, for several reasons: (1) the exploration of structural changes in scale length and width could only address one type of scale, not two, (2) strategies involving the combination of lamellar scales and piliform scales were rare and their effects could not be easily investigated with statistical models. (3) In PLF strategies (PLFC and PLFT), both lamellar scales and piliform scales were in similar density and in PLE strategies (PLEC and PLET), lamellar scales were in greater density than piliform scales, which suggested that lamellar scales played a similar role or a greater role than piliform scales for aspects linked to scale density. Although defined for the transparent zone, this terminology also applied to the opaque zone, but fewer strategies existed (Appendix S1: Table S1).

Optical measurements and visual perception modeling

We measured direct (specular) transmittance from 300 to 1,100 nm, using a deuterium-halogen lamp (Avalight DHS, Avantes), separate lighting and collecting optic fibers (FC-UV200-2-1.5 \times 100, Avantes), and a spectrometer (Starline Avaspec-2048 L, Avantes). Fibers were aligned 5 mm apart and the wing sample was placed perpendicular between them at equal distance, guaranteeing that we measured direct (specular) transmittance and not diffuse transmittance. The incident light beam made a 1 mm diameter spot. Spectra were taken relative to a dark (light off) and to a white reference (no sample between the fibers) measurement. For each species and wing, we took five measurements in the transparent zone. We did not measure diffuse transmittance as it required using an integrating sphere and put specimens in contact with it, which would have damaged the specimens.

We analyzed spectral shape using Avicol v6 (*available* online)⁹ and the pavo R package (Maia et al. 2019) to

⁹ http://sites.google.com/site/avicolprogram/

extract physically and biologically relevant parameters. We computed the mean transmittance over (300-700 nm), which described the level of transparency. In addition, we computed the proportion of UV transmittance as the ratio (total transmittance over (300-400 nm)/total transmittance over (300-700 nm)), i.e., the proportion of the total amount of transmitted light that occurred in the ultraviolet range. Notice that perfect optical transparency consists in transmitting 100% of the incident light at all wavelengths. It yields a proportion of UV transmittance of 0.25. Chitin, of which wings and wing scales are made, absorbs negligibly above 500 nm, but more as wavelength decreases, especially in the ultraviolet range (Azofeifa et al. 2012, Stavenga et al. 2014). Given that absorption + reflection + transmission = 1, a loss in transmission indicates an increase in absorption if reflection is maintained at similar levels. We also computed the mean transmittance separately for the ultraviolet range (300-400 nm), the humanvisible range (400-700 nm) and the near infrared range (700-1,100 nm). We could thus disentangle the nearinfrared range where only thermoregulation can act as a selective pressure from shorter wavelengths where vision also acts on coloration (as in Munro et al. 2019), and we could isolate the UV range to test the potential role of transparency as a parasol against UV radiation.

We also aimed to test whether the physical transparency, mean transmittance or proportional UV transmittance, translated into a biologically meaningful transparency, namely a visual contrast with a background as perceived by a potential bird predator. Exploring a large variety of visual configurations ensured robust conclusions. We thus analysed spectra using Vorobyev and Osorio's discriminability model (Vorobyev and Osorio 1998), which quantifies color and brightness contrasts between (1) an object and a background, seen (2) under an illuminant and (3) by a visual system. (1) We considered three visual conditions (see details in Appendix S1: Fig. S3): a clearwing patch could be seen against a sky, a leaf, or a trunk background (see McClure et al. (2019) and Arias et al. (2020) for average leaf and trunk spectra respectively). (2) We considered three natural illuminants that covered most of diurnal conditions encountered in terrestrial environments, i.e., in forests and open areas, as defined by Endler (1993): woodland shade (found in forest canopy and forest edge), forest shade (found in forest understorey), and large gap (found in open areas and in forest canopy). (3) We modeled both vision systems found in birds, the UVS vision and the VS vision, as both types are found in insectivorous birds (Odeen et al. 2011). For the UVS vision system, we used the spectral data of the Blue Tit (Cyanistes caeruleus) and its relative cone densities of 1:1.9:2.7:2.7 for UVS:S:M:L (Hart et al. 2000). For the VS vision system, we used the spectral data for the Shearwater (Puffinus pacificus) and its relative cone densities of 1:0.7:1:1.4 for VS:S:M:L (Hart 2004). For both models, we assumed quantum noise, a Weber fraction of 0.1 for chromatic vision (Maier and Bowmaker 1993, Lind et al. 2014) and 0.2 for achromatic vision (average of the two species studied in Lind et al. 2013). We treated brightness detection as performed by double cones (Campenhausen and Kirshfeld 1998). We used the Blue Tit and the Shearwater as their sensitivity peaks are similar to most species of their groups, making them good representative of their respective UVS and VS vision type. Moreover, they are among the very few species for which we know other parameters needed to accurately model bird visual perception, such as relative cone densities, oil droplet transmission spectra and ocular media transmission spectra (Hart et al. 2000, Hart 2004, Hart and Vorobyev 2005).

Phylogeny reconstruction

We built the phylogeny for the 123 clearwing species representing the 31 families included in our data set, as follows. First, for each of the 123 clearwing species in our data set, we searched for DNA sequences in Gen-Bank and BOLD (Ratnasingham and Hebert 2007), and if none was available, we took a species from the same genus, tribe, subfamily, or family as a substitute, choosing the lowest taxonomic level for which we had a substitute (Supplementary file File1_SpeciesList.txt for the list of analysed species and substitute species). Second, we incorporated 60 additional species from families where we only had a few species (e.g., Bombycidae, Cossidae, Drepanidae, Limacodidae, Megalopygidae, Noctuidae, Notodontidae, and Pyralidae, among others), in order to infer tree topology with more certainty. These species were chosen of the same subfamily or, if not possible, in the same family to get three to five species per family. We thus obtained a set of 183 species to build the tree. We used DNA sequences for the mitochondrial COI and COII genes, and for the nuclear CAD, EF1, GADPH, IDH, MDH, RpS5, and wg genes (Supplementary file File2 SpeciesSequences.txt). We aligned the sequences with CodonCodeAligner (version 4.2.7, CodonCode Corporation, available online)¹⁰ and concatenated them with PhyUtility (version 2.2, Smith and Dunn 2008). The data set was then partitioned by gene and codon positions and the best models of substitution were selected over all models implemented in BEAST 1.8.3 (Drummond and Rambaut 2007), using the "greedy" algorithm and linked rates implemented in Partition Finder 2.1.1 (Lanfear et al. 2017, Supplementary file File5_PartitionFinder_BestScheme.txt). We constrained the topology of all families to follow Figure S12.tre in Supplementary Archive 8 from Kawahara et al. (2019), and we used the following secondary calibrations from that figure: node joining Bombycoidea and Lasiocampoidea at 84.05 Myr with 95% credibility interval [74.15, 94.4]), Noctuidea ancestor at 77.6 Myr [66.97, 88.57]), node joining Gelechioidea to Bombycoidea at

¹⁰ http://www.codoncode.com/

105.23 Myr [93.77, 117.3]), Papilionoidea ancestor at 98.34 Myr [86.85, 110.33]); node joining Sesioidea to Cossoidea at 105.66 Myr [93.4, 118.32]), and tree root at 145.03 Myr [128.96, 161.64]) on Cipres Science Gateway (Miller et al. 2010). We constrained monophyly at genus, tribe, subfamily, and family level. Specifically, species belonging to the same genus (or tribe, subfamily, and family) were considered to form a monophyletic clade. Four independent analyses were run for 10 million generations, with one Markov Chain Monte Carlo each and a sampling frequency of one out of 10,000 generations. We examined the trace of each run, and defined a burnin period for each run independently, using Tracer 1.6 (available online).¹¹ We retained the 2 runs that had a stable trace and combined the trees using LogCombiner 1.8.4 (available online).¹² We then computed the maximum clade credibility (MCC) tree with median node ages using TreeAnnotator 1.8.4 (Drummond et al. 2012). Additional species were then pruned from the tree and we used the resulting MCC tree with 123 species in subsequent comparative analyses (Supplementary file File6_PhylogeneticTree_123species.tre).

Statistical analyses

All analyses were conducted using the R environment (R Development Core Team 2013). Data used for the analyses are available as supplementary files (File3_Da-ta_AllColors.txt and File4_Data_SpeciesMeans.txt).

Repeatability analysis.—We assessed the repeatability of optical and structural parameters, using the rptR package (Stoffel et al. 2017). For transparency measurements, we took the five measurements as repetitions of the same species and wing. For wing length, we measured each wing twice and thus obtained two repetitions of the same wing for each species. Regarding the repeatability of the measurements of scale density, length, and width, we measured a small number of scales of the same type, zone, and wing, and tested whether within-group variability was lower than between-group variability. All measurements were found highly repeatable (see sample sizes and results in Appendix S1: Table S2).

Phylogenetic signal and phylogenetic clustering.—We implemented two complementary approaches to estimate the statistical nonindependence among structural trait values due to species phylogenetic relatedness. First, we estimated the amount of phylogenetic signal in each structural and optical variable. For continuous variables, we used both Pagel's lambda (Pagel 1999) and Blomberg's K (Blomberg et al. 2003) implemented in the phytools R package (Revell 2012). For binary variables, we used Fritz and Purvis' D (Fritz and Purvis 2010) implemented in the caper R package (Orme et al.

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2018). Second, we assessed to what extent structural features and structural strategies were phylogenetically conserved, in other words we estimated the degree of phylogenetic clustering for structures. We calculated the mean pairwise phylogenetic distances (MPD) for each categorical structural parameter (Webb 2000), using the picante R package (Kembel et al. 2010). MPD measures the average phylogenetic distance that separates two species sharing a specific trait state. We computed MPD values for the forewing and the hindwing separately, or for both the forewing and the hindwing (in that case we considered all the species that presented the trait on at least one of the wings, the forewing or the hindwing). For a specific trait, we first computed the observed MPD, and then simulated MPD distribution by randomly shuffling trait values on the phylogeny. We then determined whether the observed value was lower than the 5% lower quantile of the distribution of simulated MPD values, in which case we concluded that the trait was found in species separated by fewer nodes than expected by chance.

Correlated evolution between structural traits.—To assess whether scale presence, type, insertion, and color evolved in a correlated fashion both within and across transparent and opaque zones, we computed Pagel's discrete model for those binary traits and compared the likelihood values from the dependent and independent models, using the FitPagel function, assuming a general formulation of dependency (ARD, i.e., all transition rates being different and the transition rate on character 1 depending on the state of character 2 and vice versa) from phytools (Revell 2012).

General statistical approach for structural constraints, structure-optics relationships, and ecological relevance.— We conducted (1) mixed models using the nlme R package (Pinheiro et al. 2020) and (2) Bayesian phylogenetic mixed models with Markov Chain Monte Carlo analyses using the mulTree R package (Guillerme and Healy 2019). Both approaches are suited to repeated observations but, unlike the former, the latter controls for phylogenetic relatedness. Using both allowed us to assess the influence, if any, of phylogeny on the observed relationships. (1) In the mixed model approach, we selected the best model based on minimization of the Akaike information criterion corrected for sample size (AIC_c) and based on the factors supposed to play a role in determining variation in the dependent variable. (2) In the Bayesian phylogenetic approach, we used the model formulated in the classic approach; uninformative priors were used with an inverse-gamma distribution with shape = 0.001 and scale = 0.001 for both random effect and residual variances (Hadfield 2010). We took species as random effects. Models were run using two chains of 500,000 iterations with a burn-in of 10,000 and thinning interval of 300. Fixed effects were considered statistically important and supporting the effect considered when the

¹¹ http://beast.bio.ed.ac.uk/tracer

95% credible intervals exclude zero, less so when only the 90% credible intervals exclude zero.

Structural constraints.—We aimed to investigate the variation in the proportion of clearwing area in relation to species macro and microstructural features to explore the macro and micro structural constraints on transparency. For this, we took the proportion of clearwing area as the dependent variable, and scale length, insertion, and coloration, as well as wing length, clearwing area, and wing as factors, with meaningful interactions.

We also quantified the relative investment in scales in the transparent zone as compared to the opaque zone. More specifically, we computed the relative investment in a given microstructural trait (scale density, scale length, scale width, scale surface, or scale coverage) for a specific species and wing as Relative investment_{Trait} = Trait_{transparent zone} - Trait_{opaque zone}. Values departing more from zero indicate greater changes: toward a decrease (negative values, economy of investment in the transparent relative to the opaque zone) or an increase (positive values, increase in investment in the transparent relative to the opaque zone). For each structural trait, we took the two relative investment values per species (one per wing) obtained from image analysis as observations and treated species as a random effect. We took the relative investment in a specific trait as the dependent variable. We included as fixed effects category, insertion and color in the transparent zone, scale surface, and coverage in the opaque zone, as well as wing length, clearwing area, and proportion of clearwing area, and relevant interactions between these factors.

Structure-optics relationships.—We explored which structural parameters influenced optical properties in clearwing Lepidoptera. For this purpose, we took the 10 spectral measurements per species as observations and species and wing within species as random effects. We took the mean transmittance over (300–700 nm) or the proportion of UV transmittance as the dependent variable. We included as fixed effects scale surface, density, category, insertion and color, wing length, clearwing area, and proportion of clearwing area and relevant interactions between these factors.

Ecological relevance.—We explored the ecological relevance of transparency for visual detection and more specifically for concealment, for thermoregulation, and for protection against UV. We took the 10 spectral measurements per species as observations and species and wing within species as random effects. First, we examined whether optical properties translated into perceptual transparency. We took the physical property (mean transmittance over (300–700 nm) or proportion of UV transmittance) as the dependent variable. We included as fixed effects the wing, the color contrast and brightness contrast and relevant interactions between these factors.

We expected that greater optical transparency translated into greater concealment (lower visual contrast) to a predator, whatever its visual ability, the ambient light or visual background considered.

Second, we explored the link between transparency and latitude to test a potential role of transparency in thermoregulation or protection against UV. For thermoregulation, Munro et al. (2019) have shown that body and proximal wing coloration vary with latitude in butterflies (lower reflection, hence higher absorption at high latitudes), more strongly in smaller species, and more strongly in the near-infrared range. We thus took the mean transmittance over specific ranges of wavelengths (UV, human-visible, and near-infrared) as the dependent variable and we included latitude (absolute latitudinal distance to the equator), wing length and their interaction as fixed factors. We expected lower transmittance (higher absorption) at higher latitudes if transparency could play a role in thermoregulation.

Protection against UV radiation can be ensured by UV absorption (if reflection was identical, this means lower transmission). Decreasing transmission could be ensured by limiting the absolute transmittance in the UV or, but with a lower expected efficiency, the transmittance in the UV range relative to that in the other wavelength ranges (hence the proportion of UV transmittance). Hence, we took mean transmittance over (300–400 nm) or the proportion of UV transmittance as dependent variables and tested them against latitude. We expected higher absolute or relative UV transmittance (lower absorption) at higher latitudes if transparency could play a role in UV protection.

RESULTS

Diversity of microstructures

Structural investigation showed a high diversity of structural strategies (gathering scale type, insertion, and coloration), as illustrated by a few examples (Fig. 1 and associated spectra in Fig. 2). Transparency could be achieved by the means of a nude wing membrane (N), of piliform scales (P), of lamellar scales (L), or of piliform scales and lamellar scales in combination (PL). When present, scales could be flat (F) or erect (E) on the wing membrane and colored (C) or transparent (T; Fig. 3). Rather counterintuitively, lamellar scales (L) are by far the most common structural type to achieve transparency (69/123 species, 26/31 families), followed by the absence of scales (N, 33/123 species, 12/31 families) and piliform scales alone (P, 27/123 species and 9/31 families, Appendix S1: Table S1 and Fig. S1). Rarer strategies involved either transparent erect lamellar scales (LET, 9/ 123 species, 6/31 families) or the combination of piliform scales and lamellar scales (PL, 12/123 species, 7/31 families). Within this latter category, the combination of erect piliform scales and lamellar scales was the rarest



FIG. 2. (a) Relationship between the proportion of UV transmittance and the mean transmittance over 300–700 nm. Species are represented by their forewing value. Perfect transparency (100% transmittance over 300–700 nm, resulting in a 0.25 proportion of UV transmittance, upper right corner) is represented by an empty red square and species examples from Fig. 1 are indicated by their two-letter code inside the plot. Type is P, piliform scales, or L, lamellar scales (the combination of piliform and lamellar scales was assimilated to lamellar scales); insertion is E, erect, or F, flat; and coloration is C, colored, or T, transparent. The N strategy has no scales, no insertion, and no coloration. The regression line was drawn from the best mixed model for the forewing. (b) Mean transmittance spectra of the five points of the forewing for the species listed in Fig. 1, with standard errors. Names are ordered from top to bottom according to decreasing transmittance values at 700 nm. Small pictures (see Fig. 1 for details) of the species are displayed next to their abbreviated name. Colors and symbols for structural strategies (combination of scale type, insertion, and coloration) are the same in both plots.

(PLE, 5/123 species, 3/31 families). Transparent piliform scales were always associated with lamellar scales (PL) and never alone, be they erect or flat. Overall, colored scales (P, L, or PL) concerned 72 species while transparent scales (L or PL) concerned 30 species.

Evolution of structural features

Phylogenetic signal and phylogenetic clustering.—We examined to what extent structural traits were influenced by common ancestry in their evolution. Wing

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FIG. 3. Phylogeny of the species included in this study and distribution of scale structural traits and structural strategies in these species for the forewing (columns 1, 2, 3, 7, 8) and the hindwing (columns 4, 5, 6, 9, 10). A structural strategy is defined as the combination of scale type, insertion, and coloration. Type is P, piliform scales, L, lamellar scales (piliform scales were assimilated to lamellar scales), or N, no scales. Insertion is E, erect, F, flat, and U, unknown (for N strategy). Coloration is C, colored, T, transparent, and U, unknown (for N strategy). The N strategy has no scales, no insertion, and no coloration.

macrostructure (wing area, wing length, and clearwing area) showed significant phylogenetic signal for both forewing and hindwing (Appendix S1: Table S3). By contrast, optical variables of the transparent zone generally showed no phylogenetic signal, except the mean transmittance and the brightness contrast of the hindwing, which showed significant phylogenetic signal (but only for Pagel's λ for the mean transmittance in the UV range; Appendix S1: Table S3).

At a microscopic level, scale dimensions and density showed significant phylogenetic signal in both transparent and opaque zones, for both forewing and hindwing (except for scale length in the opaque zone of the hindwing, Appendix S1: Table S3). Concerning binary structural variables, both wings showed the same evolutionary patterns (Appendix S1: Table S4): in the transparent zone, the evolution of scale insertion could fit both a Brownian motion model and a random model for the hindwing. In the transparent zone, scale presence, type, and coloration were conserved, i.e., they were significantly more clumped in the phylogeny than the Brownian Motion expectation. In the opaque zone, scale insertion was significantly more overdispersed than the random expectation, but this was likely due to little variation in that trait (1/123 species with different trait; Appendix S1: Table S4).

Several structural features showed significant phylogenetic clustering (i.e., species sharing a structural feature separated by fewer nodes than expected by chance). Specifically, in the transparent zone significant phylogenetic clustering was found on the forewing for the presence of scales, the presence of piliform scales alone or including the mixed category (PL, lamellar scales combined with piliform scales), for the erect insertion, and for the absence of transparency in scales (color C), and on the hindwing for the absence of scales and the presence of transparency in scales (Appendix S1: Table S5). In the opaque zone, there was significant clustering for the presence of lamellar scales alone (S) for the forewing, and the presence of combined piliform scales and lamellar scales (PL) for the hindwing. Considering structural strategies (i.e., the combination of given type, insertion and coloration), only 2/11 appeared phylogenetically clustered in the transparent zone whatever the wing: erect colored piliform scales (PEC), erect colored lamellar scales mixed with piliform scales (PLEC), both strategies clustered when considering wings separately or together. For the hindwing, two structural strategies appeared clustered: the nude membrane (N) and the flat transparent scales (LFT) (Appendix S1: Table S5). For the opaque zone, the structural strategy flat colored lamellar scales (LFC) was clustered for the forewing, while the structural strategy flat colored lamellar scales mixed with piliform scales (PLFC) was clustered for the hindwing.

Overall, structural strategies were less clustered phylogenetically than their composing structural parameters, scale type, insertion, and coloration. This suggested these structural parameters combined into structural strategies in multiple ways.

Correlated evolution between structural traits.—Analyses indicate that the structural strategy found in a zone (transparent or opaque) of one wing was likely to be found in the corresponding zone of the other wing. Scale presence, type (P or L), insertion, and color were correlated between wings in the transparent zone and scale type (P or L), and insertion were correlated between wings in the opaque zone (Appendix S1: Fig. S4 and Table S6). We found similar correlations in both wings: scale type (P or L) of the opaque zone was correlated with the scale type of the transparent zone. It was not the case for scale insertion. Scale type (P or L) and insertion were not correlated, in the transparent zone or the opaque zone, of the forewing or the hindwing. Conversely, in the transparent zone, scale coloration was correlated with scale type for the forewing and for the hindwing, and to scale insertion for the hindwing (Appendix S1: Fig. S4 and Table S6).

Structural constraints and investment in transparency

Analyses reveal some relationships between structural features that suggest the existence of evolutionary constraints.

At a macroscopic level, transparency occupied a larger proportion of wing area when it involved colored than transparent scales, especially in species with small wings (Table 1, Fig. 4a,c). Transparency concerned a larger proportion of wing area for smaller species (lower wing area); in other words, transparency could occupy a small or a large proportion of wing area in species with small wings, while transparency could only occupy a reduced proportion of wing area in species with large wings (Fig. 4a). These effects were stronger for the hindwing than for the forewing (Table 1, Fig. 4a).

At a microscopic level, we assessed the relative investment in transparency as the change in scale structure in the transparent zone compared to the opaque zone. As the wing area concerned by transparency increased, the difference in density between the transparent and the opaque zone decreased (Appendix S1: Table S7). As scale surface or coverage (whichever variable was retained depended on the dependent variable considered) increased, differences in scale density, length, width, surface or coverage between the transparent and opaque zones decreased (Appendix S1: Table S7). Compared to small species, larger species showed smaller changes in scale surface in the transparent zone compared to the opaque zone.

The relative investment in transparency depended on microstructural features. Compared to the opaque zone, the reduction in density in the transparent zone was greater for a nude membrane than for piliform scales (P), and to a lesser extent greater for piliform scales than

	Mixed m	Pavasian		
Factor	Estimate ± SE	t	Estimate [and 95% CI]	
Intercept	41.9 ± 5.4	7.74***	41.5 [27.2, 55.5]	
(FW > HW)	-16.5 ± 5.5	-3.02***	-19.8 [-30.1, -9.9]	
Coloration $(U > C)$	0.15 ± 2.75	0.05	-1.6 [-6.5, 3.2]	
Coloration $(T > C)$	-14.9 ± 3.3	-4.5***	-13.5 [-19.9, -6.9]	
Clearwing area	$\textbf{0.68} \pm \textbf{0.05}$	15.05***	0.71 [0.631, 0.79]	
Wing length	0.12 ± 0.34	0.34	0.39 [-0.26, 1.02]	
Wing area	-0.26 ± 0.03	-7.46 ***	-0.23 [-0.3, -0.16]	
Coloration (U > C) \times clearwing area	0.03 ± 0.02	1.8	0.028 [-0.003, 0.06] I	
Coloration (T > C) \times clearwing area	$\textbf{0.17} \pm \textbf{0.04}$	4.51***	0.134 [0.065, 0.202]	
Wing length \times clearwing area	-0.01 ± 0.001	-10.63***	-0.012 [-0.014, -0.01]	
Wing length \times wing area	$\textbf{0.004} \pm \textbf{0.0004}$	8.39***	0.003 [0.002, 0.004]	
Wing length \times (FW > HW)	1.43 ± 0.41	3.48**	1.41 [0.66, 2.17]	
Clearwing area \times (FW > HW)	0.05 ± 0.02	2.61*	0.059 [0.021, 0.098]	
Wing area \times (FW > HW)	-0.08 ± 0.02	-3.74***	-0.075 [-0.115, -0.033]	
Phylogenetic variance	_	-	146.4 [92.4, 217.2]	
Residual variance	_	-	59.3 [45.6, 76.6]	

Notes: Retained classic mixed model and Bayesian phylogenetically controlled mixed model for the proportion of surface occupied by the clearwing area. We removed the two outliers (corresponding to the saturnid *Attacus atlas*) from the data analyzed. FW, forewing; HW, hindwing, scale coloration (T, transparent; C, colored; U, unknown [for absent scales]); –, not applicable for the analysis considered. Values in boldface type are statistically important factors associated with 95% CI excluding zero in Bayesian models, less important factors are associated to 90% CI excluding zero (symbol I) and *P* values (*P < 0.05; **P < 0.01; ****P < 0.001) for the mixed model adjusted for type III sums of squares.

for lamellar scales (alone or in combination with piliform scales L or PL; Fig. 5). Scale length was reduced to zero for the nude membrane, and more reduced when scales were erect than flat, flat scales showing an increase in scale length (Fig. 5, Appendix S1: Fig. S5 and Table S7). The reduction in scale width or surface was maximal for the nude membrane, less important in piliform scales, and even less important in lamellar scales. Scale width was even increased in transparent lamellar scales compared to colored lamellar scales. Finally, strategies that involved transparent lamellar scales (alone or in combination with transparent piliform scales) showed an increase in scale surface, and an increase in scale coverage in the transparent zone compared to the opaque zone.

Overall, two trends appear: except for transparent lamellar scales, transparency was associated with a reduction in density and in scale dimensions and coverage, which was maximal for the nude membrane (Fig. 5, Appendix S1: Fig. S5). Transparent lamellar scales followed the reverse trend, with an increase in scale width, surface and coverage on the membrane compared to the opaque zone of the same wing and species (Fig. 5, Appendix S1: Fig. S5).

Optical transparency and influence of structure

Mean transmittance over (300–700 nm) was higher for the forewing than the hindwing; it was positively correlated with the proportion of UV transmittance (Fig. 2a) and this relationship did not differ between forewing and hindwing (Appendix S1: Table S8). However, a Pearson's correlation between mean transmittance over (300–700 nm) and proportion of UV transmittance yielded a moderate coefficient (r = 0.47, t = 8.18, P < 0.001), suggesting that species can play on these aspects independently, to some extent.

At a macroscopic level, mean transmittance over (300–700 nm) and the proportion of UV transmittance were higher when transparency occupied a higher proportion of wing area and for smaller species (Fig. 4d, Table 2). In addition, mean transmittance was also higher for the forewing compared to the hindwing (Table 2), which was not caused by a higher proportion of clearwing area in the forewing as the proportion of clearwing area was higher for the hindwing than for the forewing (Table 1).

At a microscopic level, mean transmittance and the proportion of UV transmittance were higher for lower scale density, for a nude membrane than for erect scales and for erect scales than for flat scales, and for transparent scales than for colored scales (Figs. 6, 7, Table 2). Mean transmittance was higher for a nude membrane or a membrane covered with piliform scales than for a membrane covered with lamellar scales or a combination of lamellar and piliform scales, while the variation in the proportion of UV transmittance was not statistically important in relation to scale type, but only in relation to scale presence with nude membrane. Most effects depended on the wing considered and some effects (e.g. scale coloration, scale insertion (U > E)) were only statistically important when controlling for phylogeny, a



FIG. 4. Proportion of clearwing area (a) in relation to wing area and structural strategies, (b) in relation to clearwing area and wing area, and (c) in relation to scale coloration. (d) Mean transmittance over 300–700 nm in relation to the proportion of clearwing area and structural strategies. Lines were drawn from the coefficients of the best mixed model for the forewing (plain line) and the hindwing (dashed line). A structural strategy is defined as the combination of scale type, insertion, and coloration. Type is P, piliform scales, or L, lamellar scales (the combination of piliform and lamellar scales were assimilated to lamellar scales); insertion is E, erect, or F, flat; and coloration is C, colored, T, transparent, and U, unknown (for N strategy). The N strategy has no scales, no insertion, and no coloration. The color codes of structural strategies, described in plot a, are identical in plot d. Species are represented by two points, one for forewing and one for hindwing. For clarity reasons, two points were removed from the plots shown in panels a and b and from analyses.

fact that may be explained by a high optical variation between closely-related species with similar microstructural features. The large variation in transmittance within each structural category shown in Fig. 6 suggests that many key features contribute to building the optical signal. Overall, optical transparency, as estimated by the mean transmittance over (300–700 nm) and the proportion of UV transmittance, depended on both wing macrostructure and wing microstructure. It increased as it occupied a larger proportion of wing area and as membrane scale coverage decreased.

Change in the transparent zone							
relative to the							LDID
opaque zone	Ν	PEC	PFC	LFC PLFC	LEC PLEC	LET PLET	LFT PLFT
Change in density	Maximal decrease	Slight decrease	Slight decrease	No change	No change	Increase	Increase
Change in length	Maximal decrease	No change	Increase	Increase	No change	No change	Increase
Change in width	Maximal decrease	Decrease	Decrease	Slight decrease	S l ight decrease	Increase	Increase
Investment in pigments	Absent	Present	Present	Present	Present	Absent	Absent
Investment in chitin	Absent	+	+	+	+	++	+++

FIG. 5. Diagram showing the main results on relative investment in transparency detailed in Appendix S1: Table S7. Relative investment in transparency measures to what extent scale structural features in the transparent zone are modified relatively to the opaque zone, considering scale density, length, and width, as well as total investment in pigments and in chitin. Structural strategies are ranked from left to right from minimal investment to maximal investment. Notice that strategies based on transparent scales (be they lamellar scales alone or in combination with piliform scales) involve a relative investment higher than in the opaque zone. A structural strategy is defined as the combination of scale type, insertion and coloration. Type is P, piliform scales, or L, lamellar scales (the combination of piliform and lamellar scales was assimilated to lamellar scales); insertion is E, erect, or F, flat; and coloration is C, colored, or T, transparent. The N strategy has no scales, no insertion, and no coloration. The color codes of structural strategies are identical as in other figures.

Ecological relevance of transparency

Relevance for visual detection and concealment.—As expected, a greater optical transparency, estimated by mean transmittance or by proportion of UV transmittance, yielded a reduced color and brightness contrast between the transparent zone and the visual background, whatever the background (sky, leaf, trunk), the illuminant (woodland shade, forest shade, large gap) and the considered bird visual system (UVS or VS) (Appendix S1: Tables S9, S10, S11, S12). The positive interaction term between color and brightness contrasts was statistically important in all models, indicating that the reduction in visual contrast decreased as optical transparency increased (Fig. 8). Moreover, we found that mean transmittance was higher for the forewing compared to the hindwing, as in previous analyses (Table 2). For a given increase in mean transmittance, the reduction in brightness contrast was stronger for the forewing than for the hindwing (Appendix S1: Tables S9, S10). For a given increase in mean transmittance, the reduction in color contrast was stronger for the hindwing, whatever the background for the woodland shade, but only for the sky background for the forest shade and large gap ambient lights (Appendix S1: Tables S9, S10). Similar results were found for both bird visual systems.

Overall, a greater optical transparency yields greater concealment in various terrestrial backgrounds and lights, as perceived by bird predators.

Relevance for thermoregulation.—We found that mean transmittance in the near infrared range (700–1,100 nm) and in the human-visible range (400–700 nm) decreased with increasing latitude (Fig. 9c, d, Appendix S1: Table S13), a pattern that supported the hypothesis that transparent wing elements have an impact on thermoregulation. The mean transmittance in the UV range

did not vary with latitude (Fig. 9a, Appendix S1: Table S13). Mean transmittance in the near infrared range (700–1,100 nm) correlated more tightly to transmittance in the human-visible range (400–700 nm) than to transmittance in the UV range (300–400 nm; 1,421 AIC counts below; Appendix S1: comparing Fig. S6B and S6A). This was compatible with a greater thermal impact of infrared wavelengths. Such variations were independent of species size (factor not retained in the analyses).

Overall, mean transmittance decreases with increasing latitude, in the near infrared range and in the humanvisible range, supporting the hypothesis of a potential role of transparency in thermoregulation.

Relevance for UV protection.—All transmittance spectra showed a decrease in transmittance in the UV range, resulting in a proportion of UV transmittance lower than 0.25 (Fig. 2). This can be interpreted as potential UV protection. Yet, this hypothesis is not supported by the analysis of the patterns of UV variation with latitude. The mean transmittance in the UV range and the proportion of UV transmittance showed no significant variation with latitude (Fig. 9a, b, Appendix S1: Table S13). These results do not support the hypothesis that transparency offered UV protection, as an increase with latitude was expected for both variables.

DISCUSSION

Diverse structural strategies produce transparency

Transparent or translucent wings are present in at least 31 families, representing a quarter of the extant Lepidoptera families, which collectively harbor a striking majority of opaque species. Transparency has thus likely evolved multiple times independently. With these

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TABLE 2. Relationships between mean transmittance over (300-700 nm) or the proportional UV transmittance and wing microstructure.

	Mean transmittance over 300–700 nm			Proportional UV transmittance [†]			
	Mixed model			Mixed model			
Factor	Estimate \pm SE	t	[and 95% CI]	Estimate \pm SE	t	[and 95% CI]	
Intercept (FW > HW) Scale surface	$71.7 \pm 5.2 \\ 10.8 \pm 3.1 \\ -$	13.88*** 3.48*** -	67.3 [41.5, 92.3] 9.3 [6, 12.7] -	$\begin{array}{c} 18.2\pm1.2\\ 1.4\pm0.5\\ -0.00018\pm0.00008\end{array}$	15.24*** 2.49* -2.15*	19.4 [11.8, 26.7] 1.6 [1, 2.2] -0.00016 [-0.00025, -0.00007]	
Scale density	-0.025 ± 0.006	-4.43***	-0.01 [-0.018, -0.003]	-0.00208 ± 0.00123	-1.69	-0.002 [-0.0034, -0.00058]	
Scale type (L,PL) > (P,N)	-10.4 ± 3.3	-3.12**	-12 [-17.6, -6.4]	0.4 ± 0.9	0.39	-0.2 [-1.4, 1]	
Insertion ($F > E$)	$-10.1~\pm~2.5$	-4.09***	-11.5 [-14.7, -8.1]	-2.7 ± 0.6	-4.69***	-3.3 [-4, -2.6]	
Insertion (U > E)	-4.3 ± 3.6	-1.19	-7.2 [-12.2, -2.3]	-2.3 ± 0.8	-2.82**	-2.7 [-3.7, -1.8]	
Proportion of clearwing area	$\textbf{0.12} \pm \textbf{0.048}$	2.49*	0.119 [0.055, 0.182]	0.03736 ± 0.00746	5.01***	0.03451 [0.0262, 0.04269]	
Clearwing area	0.007 ± 0.015	0.45	0.018 [-0.004, 0.039]	-	-	_	
Color $(T > C)$	2.4 ± 3.2	0.74	8.8 [4, 13.6]	0.9 ± 0.8	1.06	2.2 [1.2, 3.2]	
Wing length	-0.2 ± 0.1	-2.01*	-0.2 [-0.4, -0.1]	-0.02623 ± 0.02232	-1.18	-0.04793 [-0.07705, -0.01899]	
(FW > HW) × scale surface	-	-	-	-0.0001 ± 0.00007	-1.41	-0.00011 [-0.00019, -0.00004]	
$(FW > HW) \times$ scale density	-0.015 ± 0.005	-2.74**	-0.013 [-0.019, -0.007]	-0.00135 ± 0.00111	-1.22	-0.00156 [-0.00271, -0.00038]	
$(FW > HW) \times$ scale type (L,PL) > (P,N)	2.9 ± 2.4	1.21	4 [1.5, 6.4]	0.7 ± 0.5	1.37	0.7 [0.2, 1.1]	
(FW > HW) × insertion (F > E)	1.4 ± 2.1	0.67	0.5 [-1.7, 2.8]	-0.9 ± 0.4	-2.29*	-0.9 [-1.3, -0.5]	
(FW > HW) × insertion (U > E)	-8.7 ± 3.3	-2.61*	-7.9 [-11.5, -4.6]	-1.8 ± 0.6	-3.04**	-1.9 [-2.5, -1.3]	
(FW > HW) × clearwing area	-0.031 ± 0.011	-2.89**	-0.03 [-0.042, -0.018]	-	-	-	
$(FW > HW) \times color$ (T > C)	-6.2 ± 2.5	-2.44*	-6.6 [-9.2, -3.9]	-0.4 ± 0.5	-0.69	-0.5 [-1,0.1]	
Phylogenetic variance	_	-	786.8 [593.9, 1035.9]			72.1 [55.8, 93.2]	
Residual variance	_	-	55.3 [50.8, 60.4]			1.9 [1.7, 2.1]	

Notes: Best classic mixed model and Bayesian phylogenetically controlled mixed model for mean transmittance over 300–700 nm and proportion of UV transmittance (multiplied by 100 for the analysis). For all analyses, we used all 10 measurements per species. We treated species and wing within species as random factors in classic mixed models and species as a random factor in Bayesian models. A dash (-) indicates factors not retained in the best model or of no concern (for phylogenetic and residual variances). FW, forewing; HW, hindwing. Scale type, P, piliform scales; L, lamellar scales; PL, piliform scales and lamellar scales in combination; N, none. Scale insertion on the wing membrane, U, unknown (for absent scales); F, flat; E, erect. Scale coloration T, transparent; C, colored. All factors relating to scales concerned scales from the transparent zone. Values in boldface type are statistically important factors associated with 95% CI excluding zero in Bayesian models, less important factors are associated to 90% CI excluding zero (symbol **1**) and *P* values (*P < 0.05; **P < 0.01; ****P < 0.001) for the mixed model adjusted for type III sums of squares.

The proportion of transmittance was multiplied by 100 before analysis.

multiple evolutions comes a massive diversity of structural strategies (combination of scale type, insertion, and coloration), expanding the range of strategies reported in the literature in the scarce studies conducted so far in clearwing Lepidoptera. Counterintuitively, the most common structural strategy for transparency is not a nude membrane but flat lamellar scales, either colored and in lower densities or transparent and packed in high densities. While flat colored lamellar scales have been recorded in the literature



FIG. 6. Variations of transparency (mean transmittance) with structural strategies. A structural strategy is defined as the combination of scale type, insertion, and coloration. Type is P, piliform scales, or L, lamellar scales (the combination of piliform and lamellar scales was assimilated to lamellar scales); insertion is E, erect, or F, flat; and coloration is C, colored, or T, transparent. The N strategy has no scales, no insertion, and no coloration. Mean transmittance was computed over 300–700 nm; for the graph, we took one average per wing and species.

(in the nymphalid Parantica sita and the papilionid Parnassius glacialis Goodwyn et al. 2009), flat transparent lamellar scales (LFT) have not been recorded in any clearwing Lepidoptera species so far, although they have been recorded in the opaque colored papilionid Graphium sarpedon (Berthier 2007, Stavenga et al. 2012). The nude membrane, already recorded in the sphingid Cephonodes hylas (Yoshida et al. 1997), can be fully nude (one-third of cases) or have scale sockets (twothirds of cases), which are the remnants of fully developed scales shedding at first flight, as in the sphingid genus Hemaris (Schmidt 2018, Koshkin and Yevdoshenko 2019), in Cephonodes hylas (Yoshida et al. 1996, 1997) and in Heliconisa pagenstecheri (Zikán 1927, Farina 2001). Piliform scales alone have been previously reported in the saturniid Rothschildia leheau (Hernandez-Chavarria et al. 2004) or the nymphalids Greta morgane oto (Binetti et al. 2009, Siddique et al. 2015) and Cithaerias menander (Berthier 2007). The term "piliform scale" is an umbrella term that comprises monofid (bristle-like) scales and/or bifid (forked-like) scales, two types that may correspond to modifications of cover and ground scales, respectively. Indeed, both types can be found in the same transparent zone, for instance in Ithomiini species and in the drepanid Deroca hyalina. Moreover, when combined with lamellar scales, piliform scales are always monofid, which suggests they could be modified cover scales, a hypothesis that should be tested by distinctive marking of cover and ground scales during development. Piliform scales are never transparent when alone (no strategy PFT or PET), which likely results from structural or functional constraints. Finally, combined piliform scales and lamellar scales (PL) represents the rarest strategy in our data set (12 species, seven families), with erect colored piliform scales and lamellar scales (PLEC) showing a significant phylogenetic clustering. Another rare structural strategy involves transparent erect lamellar scales (LET, nine species, six families), which show no phylogenetic clustering, indicating that it evolved independently in distinct lineages. This is the first record of this LET structural strategy in the literature.

Overall, structural strategies seem less clustered phylogenetically than scale type, insertion, and coloration, suggesting that structural parameters combine into structural strategies in multiple ways. Yet, strategies are similar between forewing and hindwing, suggesting that they arise from genetic and physiological pathways that are implemented in both wings during development.

Investment in transparency differs between structural strategies

Analyses on relative investment show that structural strategies imply joint changes in different structural features. On the one hand, some strategies follow the general intuitive picture of transparency consisting of a "reduction" in scale density, dimensions, membrane coverage.



FIG. 7. Relationship between optical transparency, estimated either by (a, b) mean transmittance over 300–700 nm or (c, d) the proportion of UV transmittance, and (a, c) scale density or (b, d) scale surface. Species are represented by two values, one for forewing and one for hindwing. Lines were drawn from the coefficients of the best mixed model for the forewing (plain line) and the hindwing (dashed line). In panels b and d, for clarity reasons, two points were removed from the graph but not from the analyses. A structural strategy is defined as the combination of scale type, insertion, and coloration. Type is P, piliform scales, or L, lamellar scales (the combination of piliform and lamellar scales was assimilated to lamellar scales); insertion is E, erect, or F, flat; and coloration is C, colored, or T, transparent. The N strategy has no scale, no insertion, and no coloration.

Reduction is apparently maximal for the nude membrane, although the initial investment can be high when scales fully develop and shed subsequently. Piliform scales (always colored when alone) are longer (especially when flat), thinner and in lower density than scales in the opaque zone, resulting in a low membrane coverage. Reduction seems minimal for colored lamellar scales, which are slightly thinner than their opaque counterparts, and associated with a slight decrease in scale surface and wing coverage. On the other hand, strategies based on transparent lamellar scales show an increased investment in chitin compared to the opaque zone. Wider and shorter (when erect) or wider and longer (when flat), transparent lamellar scales increase in surface and are more densely packed. We show clear correlations between scale coloration (colored vs. transparent) and scale morphology; while such correlations have been observed in opaque butterflies (e.g., Janssen et al. 2001, Matsuoka and Monteiro 2018) and



FIG. 8. Relationships between mean transmittance as an optical property and (a, c) color or (b, d) brightness contrasts, for the forewing (dark turquoise) and the hindwing (light turquoise). Color and brightness contrast values were obtained from perceptual vision modeling (see *Methods* for details) on all transmittance measurements and are expressed in JNDs, just discriminable differences. We show two examples of viewing conditions: a clearwing patch seen against the sky and under a large gap ambient light (a, b) or seen against a green vegetation background and under a forest shade ambient light (c, d), by a UVS vision bird predator. Detailed conditions and results are presented in Appendix S1: Figure S3 and Table S9 to S11, respectively.

investigated in genetics and development, this remains to be done in the case of transparency. If we use scale coverage to estimate the investment in chitin, transparency seems maximally economical for a nude membrane without sockets, whereas it seems costlier than the opaque zone when transparency is achieved through transparent scales. But in the latter case, the lower investment in pigment for these transparent scales may compensate the higher investment in chitin. While transparency can be intuitively thought as economical because based on an absence of scale, it is far from the case. Some strategies (like LE or LF) require more material overall than the opaque zone of the same wing and even the nude membrane can result from scales fully developing before shedding. Transparency may thus be costlier than classically viewed and future studies should quantify the energetic costs of transparency during development.

Patterns common to different species emerge, which may help to reveal structural constraints. For instance, transparent lamellar scales are packed in higher densities

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FIG. 9. Tests of the potential role of wing transparency as a protection against UV radiation (left column) and in thermoregulation (right column). Left column: UV-B radiation decreases as distance increases from the equator (Beckmann et al. 2014), and so does the need for UV absorption. We thus predict an increase in transmittance in the ultraviolet 300–400 nm range, or at least in the proportion of UV transmittance with increasing distance from the equator. Corresponding results are shown in left column (a, b). Right column: Ambient temperature decreases as distance increases from the equator. Absorbing radiation helps thermoregulation at cold temperatures. We thus predict a decrease in transmittance in the near-infrared 700–1,100 nm range and at shorter wavelengths, such as in the 400–700 range or even in the UV range with increasing distance from the equator. Corresponding results are shown in right column (c, d). Lines were drawn from the coefficients of the best mixed models. For clarity reasons, we plotted only two points per species, forewing and hindwing mean transmittance values but models were run on all points.

and broadened, especially when erect. Why so? We can invoke two non-mutually exclusive reasons. First, melanization may enhance mechanical resistance, as suggested in birds (Butler and Johnson 2004) and in Lepidoptera (only mentioned as personal communication in Brakefield 1987). In insects, cuticle melanization and cuticle hardening (sclerotization) occur with little time lag during development and share very similar pathways with common precursors and some sclerotization components provide different coloration in insect cuticles, from colorless to yellow and brown (Sugumaran 2009). Knockout mutations in genes that function in the melanin pathway affect both scale coloration and scale fine structure (Matsuoka and Monteiro 2018). Shortening melanin-deprived transparent scales may be a way to reinforce their mechanical resistance; it would be interesting to explore whether these shape changes come with fine structural changes that reinforce scale mechanical resistance. This may also explain why transparent piliform scales are absent from our data set, as long and thin melanin-deprived piliform scales may be too fragile. Second, shortening erect lamellar scales and enlarging them (see the hesperid Oxynetra semihyalina in Fig. 1) may be beneficial for water repellency and mechanical resistance. Erect scales (PE piliform scales alone, LE lamellar scales alone or PLE, the combination of piliform scales and lamellar scales) provide hierarchically structured geometry with asperities and air pockets between them. Such a multiscale roughness is crucial to ensure superhydrophobicity, as shown both in theoretical studies (Nosonovsky and Bhushan 2007) and in natural systems, such as the cuticle of water striders (Goodwyn et al. 2008).

Structural strategies differ in their optical efficiency

Optical properties of transparency evolved more rapidly than structures, as shown by the absence of phylogenetic signal in most color descriptors. Structural strategies are correlated with their efficiency at transmitting light. The nude membrane category is the most efficient while lamellar scales alone or in combination with piliform scales are the least efficient. Erect scales are more efficient than flat scales, transparent are more efficient than colored scales. Yet, the large variation in transmittance within structural strategies indicates that other parameters are crucial: scale density, dimension and surface are key features. The lower the density, scale surface and coverage on the wing membrane, the higher the transmittance. Additional parameters not measured here, e.g., the pigmentation of scales and of the wing membrane, the dimension and density of nanostructures, likely play a role, as shown in recent studies (Pinna et al. 2020, Pomerantz et al. 2021). For instance, wing nanostructures like nanopillars found in the glasswing nymphalid *Greta morgane oto* create by their very shape a progressive air-chitin gradient from the air interface toward the chitinous membrane that facilitates light transmission and reduces light reflection, acting as effective antireflective devices (Binetti et al. 2009, Siddique et al. 2015). Both the density and shape of the nanorelief are crucial to determine the amount of reduction of light reflection.

Ecological relevance of transparency for visual detection, differences between small and large species

Increasing transmittance reduces the visual contrast both brightness and color contrast - offered by Lepidoptera with various visual backgrounds (sky, leaf, trunk), in various light conditions encountered in closed or open habitats, and as seen by two examples of tetrachromatic visual systems. This underlines the potential protective role offered by transparency on land and illustrates the principle of transparency, namely giving access to the visual background, whatever the illuminant, the background and the visual system. The reduction in visual contrast gets lower as mean transmittance increases, which may explain why mean transmittance never exceeds 90.25% in our data set. While the translation of mean transmittance into brightness contrast seems intuitive, the translation of mean transmittance into color contrast is more surprising and underlines that transmittance spectra are not achromatic. Indeed, even if transmittance can reach up to 95% in the (400-700 nm) range, it is lower in the UV range, which can be due either to diffuse transmittance or to absorption by molecules, such as chitin or pigments, in the scales or the wing membrane. Chitin shows higher absorption at short than long wavelengths (Azofeifa et al. 2012, Stavenga et al. 2014). Melanin has a higher specific absorption at short than at long wavelengths (Wolbarsht et al. 1981) and even a weak pigmentation produces a loss of transmittance in the UV range first.

Are there selective processes that could explain the lower transmittance in the UV range? It can result from non-mutually exclusive reasons: first, weaker selection by visually hunting predators in this range, where they are less sensitive (low number of UVS or VS sensitive cones in the retina of birds, leading to a higher relative noise, as shown in Vorobyev and Osorio [1998]). Second, wavelengths in the UV range are cut out by vegetation (Endler 1993), thereby attenuating the importance of this range for species living in closed habitats. Third, the need for communication or thermoregulation (see the last two sections of the Discussion) may offset the visual gain brought by an increase in transmittance in the UV range. Species can exploit the UV range for communication with conspecifics, as suggested in opaque species (Bybee et al. 2012).

Interestingly, our analyses reveal size-dependent strategies that can be interpreted in the context of a role of transparency in visual detection and concealment. (1) In small species (with low values of wing area), transparency spans a wide range of proportions of wing area, from spots to entire wing, but in large species (with high values of wing area), transparency is always restricted to small proportion of wing surface. (2) The efficiency of transparency at transmitting light positively correlates with the proportion of clearwing area. Hence, small and poorly transmitting spots can be present in wings of all species, such as the large saturniid Rothschildia lebeau (Hernandez-Chavarria et al. 2004) and the small thyridid Dysodia speculifera, but almost entirely transparent wings highly efficient at transmitting light can only be found in small species, such as the psychiid Chalioides ferevitrea, the wings of which are extremely difficult to detect. Why so? For small species, having large proportions of clearwing area can be profitable given their overall small size and low detectability. Conversely, large species may not benefit from high proportions of clearwing area, for several non-mutually exclusive reasons.

First, optical benefits of having large transparent areas may be limited in large species. Kang et al. (2017) have found in erebid moths that small species have cryptic coloration in both wings, while large species have cryptic coloration on the forewing, and hidden contrasting color signals on the hindwing, probably involved in startle displays. They experimentally show that crypsis fails as body size increases and secondary defenses like startle displays are more effective in large prey. If such an anti-predator switch applies to transparency, crypsis by the means of transparency could potentially fail in large species for two reasons: a conspicuous large opaque body, and a wing transparency too limited in efficiency by wing membrane thickness. More studies are needed to test whether such a size-dependent switch of transparency in wing pattern exists and to unravel its structural determinants.

Second, optical benefits may be offset by costs entailed by large transparent areas for other functions. Among the Lepidoptera species investigated so far (Wagner et al. 1996, Zheng et al. 2007, Goodwyn et al. 2009, Wanasekara and Chalivendra 2011, Fang et al. 2015), large species like the nymphalid *Parantica sita* and the papilionid *Parnassius glacialis* show moderate to high hydrophobicity (Goodwyn et al. 2009, Fang et al. 2015). Conversely, the species with the highest proportion of clearwing area, the nymphalid *Greta oto*, has one of the lowest hydrophobicity values found (Wanasekara and Chalivendra 2011). The structural determinants of hydrophobicity in clear wings remain to be identified, and potential trade-offs between optics and hydrophobicity should be investigated.

Furthermore, what are the potential benefits for large species to have small transparent spots, which sometimes account for only a few percent of wing total area? We can propose several hypotheses: (1) clear spots can create false margins, resulting in internal edges more salient than the animal true outline that function as surface disruption patterns (Stevens et al. 2009b). As suggested by Costello et al. (2020), these false internal edges can create false holes, especially when edges are enhanced; this creates false depth planes and incorrect visual categorization as depth cues. (2) Clear-spotted, cryptically colored Lepidoptera species may be protected by masquerade, predators taking them for non-edible objects of their environment, like damaged or rotten leaves, an hypothesis formulated by Janzen (1984) but never tested. (3) When transmission is poor, small spots could resemble white-pigmented spots, which could be involved in communication. More generally, small clear spots may function as eyespots, which are efficient at limiting predator attack and divert them from vital parts (Stevens et al. 2008, 2009a) and used in mate choice (Robertson and Monteiro 2005) in opaque species. Experimental studies are needed to clarify this point.

Ecological relevance of transparency for thermoregulation

The general patterns of variations in transmittance with latitudinal distance to the equator support the idea that transparency may be involved in thermoregulation. While transmittance shows its largest variation in the tropics, it decreases as at higher latitudes, with an average loss of 4-5% transmittance per 10° increase in latitude. Such variations concern the near-infrared (700-1,100 nm) range and the human-visible (400-700 nm) range, but not the UV range (300-400 nm). Given that transmission + reflection + absorption = 1, it is reasonable to think that lower direct transmittance values likely translate, at least in part, into higher absorption levels, if reflection values and diffuse transmittance values are roughly maintained at the same levels. In this context, we can interpret the decrease in transmittance with increasing latitude as an increase in absorption with increasing latitude. Our results are in agreement with previous support of the thermal melanism hypothesis (Zeuss et al. 2014, Heidrich et al. 2018, Xing et al. 2018, Stelbrink et al. 2019). Contrary to Munro et al.'s (2019) study in Australian opaque butterflies, where reflectance decreases (absorption increases) with increasing latitude more strongly in the infrared range than at shorter wavelengths, we find similar variations in the near infrared and human-visible ranges. Such similarity in the two ranges could be due to several reasons. First, chitin and melanin refraction indexes and absorption coefficients show lower wavelength-dependent variations at long than at short wavelengths (Azofeifa et al. 2012, Stavenga et al. 2014), resulting in more similar properties at longer wavelengths. Second, light intensity levels determine the maximal sighting distance at which a transparent object can be detected (Johnsen and Widder 1998, Ruxton et al. 2004). A poorly transmitting target seen in dimmer light can be as difficult to detect as a highly transmitting target seen in brighter light. Light intensities are higher in the tropics than in the temperate zone (Gueymard and Ruiz-Arias 2016) and visual systems may therefore select for higher transmittance in the tropics. As a consequence, thermoregulation (in the infrared range) and vision (at shorter wavelengths) may act concurrently and contribute to creating an autocorrelation. More studies are needed to further explore the variations in absorption of clearwing Lepidoptera, and to decipher the relative contribution of vision and thermoregulation to variation in transparency at large geographical scale.

Munro et al. (2019) have found that climate correlates to body and proximal wing coloration. Given that we have measured five points at various locations within the transparent zone, and that the transparent zone is not always located close to the insect body, finding a general correlation between latitude and transparency suggests it does not depend on the location of the transparent zone on the wing, and its proximity to the insect body. Whether optical variations exist within transparent zones in relation to the distance to the insect body requires further investigation. Contrary to Munro et al.'s (2019) study, we did not find that latitudinal variations were more pronounced in smaller species.

Ultimately, whether transparent patches are involved in thermoregulation depend not only on intrinsic thermal wing properties, but also on species behavior (basking or seeking shade, flight activity, shivering). Selection can act on both components. Although the potential thermal benefit offered by transparency may seem small compared to that of opaque patches, benefits accumulated over a life-time can be substantial (Stuart-Fox et al. 2017). Studies are needed to explore the relative contribution of physics and behavior to the thermoregulation in clearwing Lepidoptera.

Ecological relevance of transparency for UV protection?

Transmittance spectra show a constant pattern, whereby UV transmittance is always lower than transmittance at longer wavelengths. This suggests an absorption in the UV range, as shown by chitin and melanin specific absorption curves (Wolbarsht et al. 1981, Azofeifa et al. 2012, Stavenga et al. 2014). This can be the by-product of selection for a transparent material (which inevitably absorbs more at shorter wavelengths) or this could be interpreted as a protection against UV irrespective of latitudinal variations. If transparency plays a role in UV protection (and in protection against pathogens), we should expect lower transmission (a higher absorption with similar reflection levels) at lower latitudes, where UV radiation levels are higher and pathogens more abundant. Instead, our results show that UV transmission in the UV range (in absolute or in proportion) does not vary with latitude, suggesting that thermoregulation rather than protection against UV or pathogens drives transparency features in Lepidoptera. In the literature on opaque coloration, agreement with thermoregulation but not with UV protection has been found in the coloration of bird eggs, for which thermoregulation is crucial to ensure embryo development (Wisocki et al. 2020).

CONCLUSION

Transparency has evolved multiple times independently in an insect order characterized by wing opacity. These multiple gains have led to a large diversity of structural strategies that enable transparency. Optical transparency is determined by both macrostructure (length, area, proportion of clearwing area) and microstructure (scale dimensions, type, insertion, coloration, and density). Microstructural traits are tightly linked in their evolution, leading to differential investment in chitin and pigments between structural strategies. Physical transparency translates into visually effective concealment with interesting size-dependent differences. The links between transparency and latitude are consistent with thermal benefits, and much less with UV protection. In many ways, our findings echo the results that have been found in opaque Lepidoptera, showing that transparency is more complex than just enhancing concealment and is likely a multifunctional compromise. Future studies should expand measurement configurations, explore the developmental pathways of transparency, its intraspecific variations, and experimentally test its putative adaptive functions and potential trade-offs with other vital functions such as water repellency or flight ability.

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SUPPORTING INFORMATION

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Data (Gomez 2021) are available on the Open Science Framework at: https://doi.org/10.17605/OSF.IO/Q8TWE.