Physical and behavioral adaptations to prevent overheating of the living wings of butterflies

Cheng-Chia Tsai1, Richard A. Childers2, Norman Nan Shi1,5, Crystal Ren1, Julianne N. Pelaez2,6, Gary D. Bernard3, Naomi E. Pierce2,4* & Nanfang Yu1*

The wings of Lepidoptera contain a matrix of living cells whose function requires appropriate temperatures. However, given their small thermal capacity, wings can overheat rapidly in the sun. Here we analyze butterfly wings across a wide range of simulated environmental conditions, and find that regions containing living cells are maintained at cooler temperatures. Diverse scale nanostructures and non-uniform cuticle thicknesses create a heterogeneous distribution of radiative cooling that selectively reduces the temperature of structures such as wing veins and androconial organs. These tissues are supplied by circulatory, neural and tracheal systems throughout the adult lifetime, indicating that the insect wing is a dynamic, living structure. Behavioral assays show that butterflies use wings to sense visible and infrared radiation, responding with specialized behaviors to prevent overheating of their wings. Our work highlights the physiological importance of wing temperature and how it is exquisitely regulated by structural and behavioral adaptations.
The wings of Lepidoptera (butterflies and moths) are important structures both in flight and in signaling, where they have been studied intensively in the context of sexual selection\textsuperscript{1-3}, warning coloration\textsuperscript{4,5}, mimicry\textsuperscript{6-8}, and camouflage\textsuperscript{9,10}. Although fully expanded wings are primarily composed of membranes covered by scales, mature wings also contain complex networks of cellular structures, with, for example, an extensive distribution of sensilla along the wing veins\textsuperscript{11-13}. Campaniform sensilla are used to sense the strain or deformation of the wings\textsuperscript{11-14}, and bristle sensilla found along wing veins and margins sense wing beat frequencies\textsuperscript{15,19}. The afferent feedback from these mechanical sensilla to the central nervous system is critically involved in the generation of normal flight in insects\textsuperscript{11-15}. Although detailed studies focus on only a few species of Lepidoptera, the taxonomic range of the species surveyed suggest that these are general features of lepidopteran wings.

Networks of living tissues typically require a constrained range of temperatures for optimal performance. Temperature regulation is critical in insects, with even minor changes in ambient temperatures having profound effects\textsuperscript{20-23}. Much of the research on thermoregulation in adult Lepidoptera has focused on thoracic temperature and flight\textsuperscript{24-35}. Little research has been done on the thermoregulatory and thermodynamic properties of the wings themselves. This is partly because of the difficulty of applying common temperature measurement techniques to such thin and light objects with small thermal capacity. In a pioneering early work, Wasserman and Schmitz\textsuperscript{36} implanted thermistors into wings and other parts of butterflies and then heated the insects with a calibrated light beam. This technique demonstrated that antennae, wings and thorax heat up at different rates and reach specific excess temperatures. However, these implants have the drawback that they can substantially change the local thermal capacity of the wing and lead to inaccurate measurements of both wing temperatures and the dynamics of temperature change. Additional research on the temperature of butterfly wings has relied upon detecting thermal radiation generated by the wings using infrared cameras\textsuperscript{37}. However, butterfly wings are largely semitransparent in the mid-infrared spectrum (except for the thick veins near the base of the wing), meaning that infrared radiation detected by the camera is only partially contributed by the wings, leading to errors in estimating the temperature over the entire wing.

In this work, we investigate the thermodynamic and thermoregulatory properties of the wing itself, first by observing the distribution and persistence over the entire adult lifespan of living tissues in the wings of butterflies, and then by studying the thermodynamics of the wing and the physical and behavioral adaptations that modulate wing temperatures in ways that protect these tissues. Using a robust ecological model species, Vanessa cardui\textsuperscript{38,39}, we show that circulatory and tracheal systems remain active in the wing veins throughout the entire adult stage. We develop an infrared hyperspectral imaging technique that permits accurate temperature mapping of objects that are both lightweight and translucent in the mid-infrared. We find that wings are extremely heterogeneous in their thermodynamic properties, with wing veins and other living parts of the wing being cooler than inter-vein regions under solar radiation. This is achieved via elevated thermal emissivity due to a thickened chitinous layer and specialized nanostructuring of their scales such that they more efficiently dissipate heat through thermal radiation.

In addition to these structural modifications, butterflies employ behavioral strategies to control the heating of their wings by the external environment. We present the responses of species of butterflies representing six families with different sizes, shapes, and coloration to thermal stimulations of the wings. Even heating by a small localized laser spot is sufficient to elicit displacement behaviors, and there is a tight window of trigger temperatures across families.

Butterflies bask for warmth at cool temperatures, and here again, previous studies have focused on thoracic temperature thresholds necessary for flight\textsuperscript{24-35}. We find that when butterflies settle and convective cooling of the wings is substantially reduced, the wings can overheat quickly (within 10 s) under strong solar radiation, whereas other bulkier body parts (head, thorax, and abdomen), whose temperatures are less volatile, remain in a thermally comfortable window. It is therefore the wing temperatures, rather than the thoracic ones, that determine when basking ceases. These physical and behavioral adaptations underscore the importance of wings as highly responsive living structures and the centrality of wing thermoregulation to lepidopteran biology.

Results

Butterfly wings as living structures. We conducted morphological studies of the wing micro-structures after carefully removing scales from both sides of the wings of living Vanessa cardui, Satyrium caryaevorus, and Parnassius m-album butterflies. We selected these species for both pragmatic and biological reasons. Vanessa cardui (Fig. 1a) is a well-known ecological organism because it occurs worldwide, engages in long-distance migration, is highly polyphagous, and can be purchased and raised on artificial diet. The other two species were chosen because they possess androconial scent pads and are locally available and easy to collect and observe in the field.

Campaniform and bristle sensilla similar to those found in other lepidopteran species\textsuperscript{12-15} are present in all the wing veins we have inspected using optical microscopy, including interior and marginal veins, and each vein contains multiple sensilla (Fig. 1b, c and Supplementary Fig. 1d). Staining using methylene blue\textsuperscript{40-42} shows that each sensillum is connected via axons to a network of nerves distributed throughout the wing veins (Fig. 1c-e, Supplementary Figs. 1, 2, and Supplementary Movies 1, 2).

The functional morphology of wings has been described in a number of insect orders\textsuperscript{43-46}, but the relationship between the circulatory and tracheal systems in wings has only been explored in detail for a few groups\textsuperscript{11}. In V. cardui, we observed that wing veins provide pathways for the tidal flow of air and hemolymph (Fig. 1f, g and Supplementary Movie 3). Each of the larger veins contains a central tracheal tube and one or two hemolymph channels (Fig. 1f and Supplementary Fig. 4b). Large, semitransparent cells (i.e., hemocytes) can be observed flowing inside the channels, and enable continuous monitoring of the hemolymph flow direction and speed. The flow of hemolymph changes directions rhythmically. Given that the tracheal tube and the hemolymph channels occupy the rigid wing vein, an increase in volume of one will concomitantly decrease the volume of the other: when the tracheal tube expands, air enters the wing vein, and hemolymph is drawn from the tip to the base of the wing and eventually into the thorax; when the tracheal tube decreases in diameter, air flows out of the wing vein, and hemolymph flows back out to the wing tips (Supplementary Movie 3). The frequency of this tidal flow of hemolymph and air in and out of the wing veins is on the order of once every few minutes when the butterflies are not agitated (Fig. 1g).

In V. cardui, hemolymph flow sustains over the lifespan of the adult. We quantified the hemolymph flow by the dilation and contraction rates of hemolymph channel width as a function of time, taking advantage of the strong correlation between hemolymph flow rate and the change in width of the hemolymph channel (Supplementary Fig. 4c). Dilation and contraction of
hemolymph channels remain active for more than 3 weeks after eclosion, and the hemolymph flow does not weaken appreciably with age (Fig. 1g and Supplementary Fig. 6, \( n = 22 \) individuals tested).

Males of a number of species of Lepidoptera have androconial organs on their wings involved in the generation and dissemination of pheromones\(^ {47-51}\). These alar androconial organs are extremely diverse, and we focused our analysis on only two kinds of androconial organs of the hyperdiverse tribe Eumaeini (Lepidoptera: Lycaenidae), termed “scent pads” and “scent patches”\(^ {52,53}\). Scent pads are unique to the Eumaeini, and occur in small regions of the wing where the wing membranes are not
fused, thereby forming intermembrane pockets; scent patches are found in many other groups of butterflies as well as in the Eumaeini, and occur in regions of the wing where the dorsal and ventral membranes are fused. Both scent pads and patches are covered by modified scales that are distinct from scales on other parts of the wing. When found together, they are almost always adjacent to each other.

We observed that scent pads of *S. caryaevorus* and *P. m-album* butterflies both contain intricate networks of hemolymph channels and living tissues (Fig. 2b, d and Supplementary Fig. 2). The flow of hemolymph through the scent pads is rhythmical and unidirectional: it enters the organ from the costal vein defining the leading edge of the discal cell, spreads throughout the intermembrane pocket, and drains into the anterior vein of the discal cell (Fig. 2b, d, e and Supplementary Movies 4–7). This is
different from the tidal flow pattern observed in the veins of V. cardui. In both S. caryaevorus and P. m-album, we observed a “wing heart” that beats endogenously at a rate of a few dozen times per minute and appears to draw hemolymph unidirectionally through the scent pad (Fig. 2b, d and Supplementary Movies 4, 5).

Whole-spectrum response of butterfly wings. Extensive studies have established that the visible colors of butterfly wings are created by the deposition of pigments in the wing scales and/or optical diffraction by nanostructuring of the scales.54–59. Here we examined butterfly wings not only under light that is visible to the human eye, but also under wavelengths beyond the visible, since the whole-spectrum response of butterfly wings, from the ultraviolet to the visible to the mid-infrared, governs their thermodynamic properties.60–63 (Fig. 3).

Consider a male Bistonina biston (Lycaenidae: Eumaeini) (Fig. 4a). The dorsal forewing surface is covered by at least four types of wing scales (Fig. 4d), and the wing contains androconial organs consisting of a “scent patch” and a “scent pad”, both covered by modified scales. Spectroscopic measurements show that, with dorsal irradiation, the region covered by black scales has stronger solar absorption (i.e., absorptivity integrated over the solar spectrum of 0.71) than the lighter-colored scent patch, scent pad, and structural blue region that have integrated solar absorptivities of 0.60, 0.63, and 0.59, respectively (Fig. 4b upper panel). The lighter-colored ventral wing surface (Fig. 4b lower panel) has reduced solar absorption: the four ventral regions that correspond to the black region, scent patch, scent pad, and blue region on the dorsal side have integrated solar absorptivities of 0.57, 0.59, 0.60, and 0.45, respectively. All wing regions have substantially lower solar absorptivity in the near-infrared ($\lambda = 0.7$–1.7 $\mu$m) than in the visible (Fig. 4b); this helps to reduce the temperature of the wings, which are more susceptible to overheating by the sun than bulkier body parts.

While the visible wing color is related to the absorption and reflection of sunlight, it has little correlation with the wing’s ability to cool down via thermal radiation. Those parts of the heated wing with higher emissivity are better able to dissipate heat through thermal radiation to a cool environment (e.g., the sky and terrestrial environment)60,62,64 (Fig. 3). We conducted hyperspectral imaging of the wings of an adult B. biston in the mid-infrared ($\lambda = 2.5$–17 $\mu$m) and found that different parts of the butterfly wing have dramatically different thermal emissivities. In particular, the scent patch, scent pad, and wing veins have emissivities approaching unity (i.e., the emissivity of a perfect blackbody) (Fig. 4c, e). Our spectroscopic studies of individual wing scales show that the scent patch scales are by far the most emissive among all types of wing scales on B. biston (Fig. 4g). We found that the high thermal emissivity of the scent patch is caused by unique nanostructures of the scales (Fig. 4d, h, i), whereas the high emissivities of the scent pad and the wing veins are due primarily to the physical thickness of their unfused membranes (Fig. 4f). The scent patch scales of other species of Eumaeini are similarly nanostructured as those found on the wings of B. biston, and also possess elevated emissivity (Supplementary Fig. 9). We have characterized the infrared properties of the wings of individual adults representing more than 50 species (Supplementary Tables 6, 7) and found that scent patches, scent pads, and wing veins in all studied species, without exception, have elevated thermal emissivity and thus enhanced radiative-cooling capabilities (Supplementary Figs. 18–31).

Wing temperatures under simulated environmental conditions. We developed a noninvasive technique based on infrared
Hyperspectral imaging to accurately determine wing temperatures under experimental conditions that simulate the butterflies’ natural environment (Supplementary Fig. 33). By measuring mid-infrared emissivity, transmissivity, and reflectivity spectra of the butterfly wing, we can distinguish thermal radiation generated directly by the wing from environmental thermal radiation that reflects from and transmits through the wing (Supplementary Fig. 33c), and thus derive temperature distributions across the entire wing (Supplementary Fig. 36).

Temperature maps for the forewings of six Eumaeini butterflies are shown in Fig. 5. We observed that, despite the wings’ wide variation in visible coloration and pattern, the average temperature of the scent patches, pads, and wing veins that contain living cells (including hemolymph cells, androconial cells, and mechanical and thermal sensilla) is always lower than that of the remaining “non-living” parts of the wings (i.e., regions with fused membranes). We also observed that a cooler radiative background (e.g., \( T_c = -40 \, ^\circ C \)) leads to reduced overall wing
temperatures (Fig. 5 and Supplementary Fig. 38) because of a larger net radiative heat transfer from the heated wings to the background. Under the condition of ventral irradiation of light equivalent to the full sun, the scent patches, pads, veins, and margins rarely exceed 45 °C. This offers a plausible explanation for why many Eumaenini butterflies tend to close their wings when resting or basking, and infrequently expose their dorsal wing surfaces.

Bright visible and near-infrared coloration does help reduce solar absorption and thus reduce temperatures (e.g., the light gray ventral side of Strephonota tephraeus, Fig. 5c), and the white bands on the ventral side of Panthiades aeolus, Fig. 5f), but the largest temperature reduction is achieved by a combination of low solar absorption and efficient radiative heat dissipation: the white scent pad of Rekoa meton is ~15 °C cooler than the hottest part of the wing (Fig. 5a), and the light-colored scent patch and pad of B. bistom are ~10 °C cooler than the hottest part of the wing (Fig. 5b). The cryptic scent patch of Theritas hemon becomes visible in the temperature map due to its higher thermal emissivity and cooler temperature (Fig. 5d).

The importance of high thermal emissivity in reducing wing temperatures is further exemplified in the case of dorsal irradiation of T. mavors and P. aeolus: the temperature of their black scent patches and pads, which have high solar absorptivity but also near-unity thermal emissivity, is 2–5 °C cooler than the hottest part of the wing (Fig. 5e, f).

Wings as temperature sensors that modulate behavior. The importance of wing temperatures can also be demonstrated by behavioral experiments. We conducted a study of species representing 6 of the 7 recognized families of butterflies (all but Hedylidae) that were readily available to us. During each experiment, a collimated laser beam (either in the visible or near-infrared spectrum) was directed onto a butterfly wing with normal incidence to increase the wing temperature locally. Laser intensity was selected such that the temperature increased from 25 to 45 °C in 7–8 s. Butterflies were observed to move in characteristic ways to displace thermal stimuli applied to their wings (Fig. 6a–g). These reactions could be reliably elicited by positioning the laser spot anywhere along the wing veins, or on the androconial organs, indicating that the “living” components in the wings contain a distributed network of thermal sensors (Supplementary Fig. 44 and Supplementary Movies 8, 9).

We used thermal camera videos to record the peak wing temperature, $T_{\text{trigger}}$, at which butterflies started to show displacement responses. We observed that butterflies across diverse families converge on a $T_{\text{trigger}}$ of about 42 °C ($n = 50$ species, $T_{\text{trigger}} = 41.89 \pm 1.54 ^\circ \text{C}$) (Fig. 6h). Butterflies in the family Papilionidae show the highest family average $T_{\text{trigger}}$ of 43.55 ± 1.53 °C; butterflies in the Hesperididae and Lycaenidae show lower average $T_{\text{trigger}}$ of 41.15 ± 1.04 °C and 40.78 ± 1.00 °C, respectively. Further research on wing solar absorptivity and thermal emissivity in the context of the ecology of each species would be necessary to explain the subtle differences in $T_{\text{trigger}}$. In 10 out of 11 species with sufficient sample sizes for comparison, females have a higher median $T_{\text{trigger}}$ than males (Fig. 6h). The exception to this pattern among the butterflies we analyzed is the Zabulon skipper, P. zabulon, where females have a lower $T_{\text{trigger}}$ than males. These butterflies are sexually dimorphic, and the dark brown females may overheat more readily than light-colored males.

We also studied the basking behavior of butterflies under experimental conditions simulating natural environmental conditions (Supplementary Movies 10, 11). Adults of three species of hairstreak, P. m-album, S. favonius, and S. caryaevorus exhibit lateral basking24,25 behavior by closing their wings dorsally and tilting sideways to present their ventral wing surfaces at right angles to the light source (Fig. 7a–c). In laboratory experiments, we used a Xenon lamp shining uniformly onto a substrate from above to simulate sunlight. We observed that the percentage of time the butterflies spent basking decreased as the ambient temperature, $T_a$, increased (Fig. 7e). Under the same experimental conditions, adults of S. caryaevorus basked far less than did those of P. m-album and S. favonius. Additional work would be needed to explain differences in basking behaviors, but differences in heating caused by coloration can help explain the observed variation in basking time between these species.

We monitored the thoracic and wing temperatures of individuals of S. caryaevorus under experimental conditions (ambient temperature of 21.5 °C and light intensity of 66.0 mW/cm², corresponding to 2/3 of full sun in the field) and found that the effect of basking on heating up the wings was considerably stronger and faster than its effect on heating up the thorax because of the relatively small thermal capacity of the wings (Fig. 7f). For example, basking increased the temperature of the wing veins by 4.48 ± 2.25 °C, reaching 39.5 ± 1.52 °C, and increased the temperature of the scent pads by 6.60 ± 2.19 °C, reaching 40.0 ± 1.54 °C, while during the same time frame, the temperature of the thorax increased by only 1.01 ± 0.54 °C. The data suggest that cessation of basking occurs when wing temperatures are too high (i.e., ~40 °C), whereas thoracic temperatures remains relatively constant during basking. This thoracic homeostasis may allow the butterflies to remain flight-ready.

Similar experiments to monitor the thoracic and wing temperatures of the two hairstreaks, S. favonius (n = 4) and S. caryaevorus (n = 6) when basking under a lamp corresponding to 2/3 of full sun and at a variety of ambient temperatures showed that the peak wing temperature at the moment when the butterflies terminated basking was 4.08 °C higher than that of the thorax, for S. favonius (Fig. 7g), and 8.04 °C higher than that of the thorax, for S. caryaevorus (Fig. 7h), again supporting the
hypothesis that butterflies stop basking when wing temperatures have become too high.

In our field studies using a thermal camera, we found that *S. caryaevorus* butterflies almost always bask in the sun within a few seconds after ceasing flight (Supplementary Fig. 45 and Supplementary Movie 12). At that moment, the thoracic temperature is reasonably high (>30 °C) as a result of the metabolic heat generated by the flight muscles, but the wings, due
Fig. 5 Temperature distributions on the forewing of six species of Eumaeini butterflies. The temperature distributions were obtained by processing infrared flux images taken by a thermal camera with measured two-dimensional emissivity, transmissivity and reflectivity maps of the butterfly wings (Supplementary Fig. 36). The thermodynamic steady state was reached within 10 s of the exposure of the wings to the illumination due to their small thermal capacity (Supplementary Fig. 39). Within each sub-figure, the top shows photos of the dorsal and ventral forewing, and the bottom shows wing temperature distributions under four experimental conditions (i.e., light simulating full sun incident on the dorsal or ventral wing surface, and cryostat temperature $T_c$, of $+20 ^{\circ}C$ or $-40 ^{\circ}C$). $T_c = +20 ^{\circ}C$ simulates an environmental condition with a relatively high ambient temperature and humidity; $T_c = -40 ^{\circ}C$ simulates an environmental condition with a relatively low ambient temperature and humidity. The exact combinations of ambient temperature and humidity simulated by $T_c = +20 ^{\circ}C$ or $-40 ^{\circ}C$ are provided in Supplementary Fig. 35. One individual per species was tested for this study.

to strong convection during flight, have temperatures close to the ambient temperature. Our lab experiments confirmed that at the moment when the S. cayaevarus butterflies stopped flight, their thoracic temperatures were in the range of 30.9 ± 0.73 °C, while their wing temperatures were only 0.73 ± 0.47 °C above the ambient temperature, which varied from 25 to 28 °C (Fig. 7i). Basking behavior may thus be elicited by cold wings: presumably the temperature of the wings must be sufficiently high for optimal function of sensilla, which are crucial for such agile flying insects.

Discussion

While previous work on lepidopteran wings has primarily focused on visible patterns involved in inter-specific and intra-specific signaling, this work emphasizes the interactions of the wings as physical structures with the abiotic electromagnetic environment, including solar and thermal radiation. Our results show that butterfly wings play a largely unappreciated role in sensing solar radiation, and that their spectral properties across wavelengths beyond the visible, including the infrared solar and thermal spectra, are tailored to help them better adapt to the electromagnetic radiation environment. Selection has favored adaptations such as heterogeneous cuticle thickness, specialized scale nanostructures, contrasting dorsal and ventral wing coloration, and finely-tuned behavioral reactions to protect the vulnerable living tissues within wings.

When butterflies close their wings, exposing only their ventral sides, the thermal emissivity of the closed wings is larger than that of a single wing due to their increased physical thickness. In the Eumaeini (Lycaeidae), such an increase of thermal emissivity is large for the non-living regions with fused membranes (e.g., from 0.30 to 0.51), moderate for the wing veins (e.g., from 0.70 to 0.91), and small for the scent patches and pads (e.g., from 0.95 to 0.998). Thus, closing the wings would reduce primarily the temperature of the regions with fused membranes, and because wing veins, scent pads and patches are bounded by regions with fused membranes, their temperatures would also decrease via thermal conduction.

An additional effect of wing closure is that the time constant of temperature changes increases substantially because of the increased thermal capacity. Thus dorsal basking with open wings and lateral basking with closed wings could have different effects on the thermodynamics of butterfly wings. In tropical and temperate climates, lateral basking with enhanced thermal emissivity and increased thermal capacity of the stacked wing pairs enables butterflies to bask longer to warm up their thoraces while reducing the risk of overheating the wings; in contrast, butterflies at high latitudes or altitudes could warm up their wings efficiently by dorsal basking in the sunlight with their wings fully or partially open.

In The Naturalist on the River Amazon, Henry Walter Bates writes about butterfly wings, "On these expanded membranes Nature writes, as on a tablet, the story of the modifications of species, so truly do all changes of the organization register themselves thereon..." This study offers a perspective on butterfly wings as living structures through an analysis of their thermodynamic and thermoregulatory properties. We have considered the synergistic effects that the electromagnetic environment has had in shaping the morphology, physiology and behavior of butterflies, and identified physical properties and behaviors that are likely to be applicable to the flying wings of most insects when exposed to sunlight.

Methods

Observing micro-structures and hemolymph flow in wings. Vanessa cardui, Satyrium caryaevarus, and Parrhasius m-album butterflies were anesthetized with CO$_2$ and scales were carefully removed from both sides of the wings with a paintbrush. Each specimen was then secured using a low melting temperature wax to two microscope slides, one on either side of the body (Supplementary Fig. 4a). Photos and videos were taken with a microscope equipped with a CCD camera. The direction and speed of hemolymph flow (Figs. 1g, 2b, d, e and Supplementary Fig. 5c) were inferred from the movement of semitransparent cells (hemocytes) inside the hemolymph channels and scent pads (Supplementary Movies 3–7). Stacking photography was sometimes used to better resolve fine structures within the wings (Figs. 1b–e, 2b, d and Supplementary Figs. 1–3, 5, 10–17).

Methylene blue, a dye that has been widely used to stain nervous tissue, including the neurons of Lepidoptera, was used to stain the neurons in wing veins. Adult V. cardui and S. caryaevarus butterflies were anesthetized. A saturated aqueous solution of methylene blue (Aldron Contraction) was injected into the thoraces: 50–100 µl for individual V. cardui and 10 µl for individual S. caryaevarus. The stained wings were removed 6 h after injection, when the butterflies had recovered from anesthesia and the injected stain had been circulated by the hemolymph into the wing veins. The scales were removed from both sides of the wings and stacking photography was used to take pictures of stained tissues.

Infrared hyperspectral imaging of butterfly wings. The infrared optical properties of butterfly wings were measured using a hyperspectral imaging technique. To measure the transmissivity distribution, the wing was secured by two pairs of thin threads and then raster scanned under an infrared microscope (Bruker Hyperion 2000), with one transmission spectrum taken at each position. The raster scan comprised two-dimensional ($x$, $y$) data cube, where $x$ includes 1763 wavelengths ranging from 2.5 to 16.7 µm and the step of scan is $\Delta x = \Delta y = 90$ µm (comparable to the sizes of individual wing scales). Similarly, an optical reflection hyperspectral data cube was generated by collecting spectra of reflected infrared light at all $(x, y)$ positions over the butterfly wing. An emissivity hyperspectral data cube was then created by calculating the absorption spectrum at each position using 1 – transmissivity – reflectivity, as Kirchoff’s law of thermal radiation states that radiative emission and absorption are reciprocal processes, and thus emissivity equals absorptivity. The hyperspectral data cubes (simplified version of the data cube for B. bistort shown in Fig. 4e, Supplementary Fig. 3b) of other species in Supplementary Figs. 18–31) enable us to visualize butterfly wings at different infrared wavelengths. Features that are not discernible to the naked eye stand out, and these features are not always correlated with visible patterns of butterfly wings. The hyperspectral data cubes were used to derive the temperature distribution on the butterfly wings.

Infrared properties of chitin used in full-wave simulations. The membranes and scales of butterfly wings are mainly composed of the long-chain polymer chitin (C$_x$H$_{2n+2}$O$_{n+1}$), which is the primary component of the exoskeletons of arthropods, including some crustaceans (such as crabs and shrimps) and insects. The optical properties of chitin have not been studied thoroughly due to the difficulty in the production of optical-grade materials and its various derivatives depending on the sources. An average refractive index of 1.56 has been used in many publications and previous studies reported the dispersive complex refractive indices of shrimp chitin samples only in the UV and visible spectral range ($\lambda =$ 250–750 nm). Here, the dispersive complex refractive indices of chitin composing the butterfly
wings in the mid-infrared range (λ = 2.5–20 μm) were derived from measured spectra and applied in full-wave simulations to investigate the interactions between mid-infrared light and nanostructured wing scales. The chemical composition and molecular structure of chitin have been identified using various methods including infrared spectroscopy and X-ray diffraction. Thirteen resonance frequencies in the mid-infrared spectral range corresponding to the major vibrational and rotational bands of chitin molecules have been characterized (Supplementary Table 2) and applied in the Lorentz oscillator model:

$$\epsilon(\omega) = \frac{\epsilon_0}{1 + \sum_m \epsilon_m (\omega_{\text{lo}}^2 - \omega^2)^{\gamma_m} + i \epsilon_m (\omega_{\text{lo}}^2 - \omega^2)^{\gamma_m} \omega}$$

where $\epsilon_0$ is the dielectric constant in the absence of the electric field, $\omega$ is the angular frequency of the incident light, $\omega_{\text{lo}}$ is the low-energy resonance frequency of the Lorentz oscillator, and $\gamma_m$ is the Lorentzian line width of the $m$th resonance.

**Fig. 6 Displacement response induced by local heating of butterfly wings.** a-g Photos and thermal camera images showing the butterflies and their typical displacement responses when the wings are locally heated by a laser beam. For example, adults of *Eumaes alata* tend to tilt their wings to reduce light intensity and thus temperature on the wing; those of *Panhasus m-album* either walk away from the laser beam path or change their body orientation to reduce wing temperature. Numbers in the thermal camera images indicate time in seconds. Supplementary Movies 8, 9 show a number of species of butterfly wings. h Wings with laser beam. A line within the box marks the median, and a square within the box marks the mean. Whiskers below and above the box indicate the 25th and 75th percentiles, a line within the box marks the median, and a square within the box marks the mean. Supplemetary Tables 2 and 3 provide a Source Data file. Source data are provided as a Source Data file.
Fig. 7 Basking behavior of three species of hairstreak butterflies. a–c Photos showing lateral basking of *P. m-album*, *S. favonius*, and *S. caryaevorus*. d Thermal camera image of an adult of *S. caryaevorus* (uncorrected for the emissivity distribution on the butterfly) at the end of a representative basking event, showing that the scent pads and veins are much hotter than the thorax. e Summary of the basking experiments showing that the percentage of time spent basking decreases as the ambient temperature, $T_a$, increases for the three species of hairstreak butterflies, and that under the same conditions, the *S. caryaevorus* butterflies ($n = 6$) require far less basking than do those of *P. m-album* ($n = 10$) and *S. favonius* ($n = 4$). Supplementary Movies 10, 11 show, respectively, basking of *S. caryaevorus* and *P. m-album* at low and high ambient temperatures. Error bars represent one standard deviation of uncertainty. f Recorded temperature increases as a function of time during 20 basking events for an individual of *S. caryaevorus*. The experimental conditions are $T_a = 21.5 \, ^\circ C$ and lamp power of 66.0 mW/cm² (corresponding to 2/3 of full sun). g, h Temperatures of the wing and thorax at the cessation of basking for *S. favonius* ($n = 4$) and *S. caryaevorus* ($n = 6$), respectively, with lamp intensity corresponding to 2/3 of full sun and for a variety of ambient temperatures. i Data showing the temperatures of the thorax, scent pad, and wing veins at the point when a *S. caryaevorus* butterfly stops flying. Source data are provided as a Source Data file.

to calculate the complex refractive indices $n(\omega) = \Re(\omega) + i\Delta(\omega)$, which were then used in the transfer matrix method to calculate the absorption spectrum of a thin film of chitin. In Eq. (1), $m$ labels the various vibrational and rotational bands of chitin, $\epsilon_0 = 8.854 \times 10^{-12} \text{ C}^2 \text{ N}^{-1} \text{ m}^{-2}$ is the permittivity of free space, and $\omega_m$ is the $m$th resonance frequency. The values of plasma frequency, $\omega_p$, and collision frequency, $\gamma_m$, were simultaneously tuned to obtain the best fit between the calculated absorption spectrum and a measured spectrum from a piece of butterfly wing membrane with scales carefully removed, which can be considered as a relatively flat slab with a uniform thickness. The absorption spectrum of the scale-less wing membrane was measured using a Fourier transform infrared spectrometer and its thickness was determined using a scanning electron microscope. The obtained complex refractive indices of chitin were used in full-wave simulations to investigate the interactions between infrared light and microstructures of wing scales (Fig. 4h, i).

Experimental setup simulating natural radiative environment. A butterfly wing absorbs sunlight and exchanges energy via thermal radiation with the ambient environment, including the terrestrial environment beneath the wing and the sky above (Fig. 3). The radiative energy loss from the heated wing to the cold sky represents a major channel of heat dissipation in addition to heat losses via convection.

In the thermodynamic experimental setup (Supplementary Fig. 33), the light produced by a Xenon lamp shining from above at a distance of about half a meter was used to simulate sunlight. The lamp occupied only a small solid angle in the top hemisphere. The lab space over the top hemisphere was maintained at $T_a = 25 \, ^\circ C$ to simulate the terrestrial environment. A temperature controllable cryostat equipped with an infrared-transparent window underneath the wing specimen was used to simulate the sky in different weather conditions. The cryostat consists of a vacuum cryostat chamber, a cooled plate located at the center of the chamber, and a potassium bromide (KBr) window. The cooled plate is in contact with a liquid-nitrogen-cooled cold finger and its temperature can be controlled within 0.01 °C of a set temperature by using a Lake Shore cryogenic temperature controller. The top surface of the cooled plate that faces the window is coated with a thick layer of paint and has a near-unity mid-infrared emissivity. The KBr window, 1.5 inches in diameter, has a transmissivity >90% over a broad mid-infrared spectral range ($\lambda = 2.5–20 \, \mu m$). The negligible reflectivity of the window minimizes reflections within the window and between the window and the wing.
and with downward atmospheric thermal radiance calculated using theoretical planar substrate with a vacuum speed of light; the cryostat setup ranges from 289 to 415 W/cm², when the temperature of the cryostat behind the wing: 

\[ \Phi(T_w) = \frac{2\pi \kappa(T_w, T_{cam})}{4} \int_0^\lambda c_\kappa(\lambda, \theta) \sin \theta \, d\theta \]

\[ \Phi(T) = \frac{2\pi \kappa(T, T_{cam})}{4} \int_0^\lambda c_\kappa(\lambda, \theta) \sin \theta \, d\theta \]

\[ \Phi(T_y) = \frac{2\pi \kappa(T_y, T_{cam})}{4} \int_0^\lambda c_\kappa(\lambda, \theta) \sin \theta \, d\theta \]

The sum of these three terms equals the radiator received by the camera: 

\[ \Phi_{total}(T_{cam}) = 2\pi \int_0^\lambda c_\kappa(\lambda, \theta) \sin \theta \, d\theta \]

\[ \Phi_{total}(T) = 2\pi \int_0^\lambda c_\kappa(\lambda, \theta) \sin \theta \, d\theta \]

\[ \Phi_{total}(T_y) = 2\pi \int_0^\lambda c_\kappa(\lambda, \theta) \sin \theta \, d\theta \]

Deriving wing temperatures from thermal camera images. A noninvasive technique based on the results of infrared hyperspectral imaging was developed to map temperature distributions on butterfly wings. The infrared flux detected by the thermal camera (FLIR SC660) (examples for B. bistorta shown in Supplementary Fig. 36a) has three contributions (Supplementary Fig. 33c):

\[ \Phi_{cam} = \Phi_0 + \Phi_1 + \Phi_2 \]

where \( \Phi_0 \) is the thermal radiation produced by the wing, \( \Phi_1 \) is the thermal radiation produced by the cryostat and transmitted through the wing, and \( \Phi_2 \) is the thermal radiation produced by the surrounding environment and reflected by the wing:

\[ \Phi_0(T_w) = 2\pi \int_0^\lambda c_\kappa(\lambda, \theta) \sin \theta \, d\theta \]

\[ \Phi_1(T_w) = 2\pi \int_0^\lambda c_\kappa(\lambda, \theta) \sin \theta \, d\theta \]

\[ \Phi_2(T_w) = 2\pi \int_0^\lambda c_\kappa(\lambda, \theta) \sin \theta \, d\theta \]

The rate at which the temperature increased was chosen to prevent overheating. Displacement in response to local heating of butterfly wing. In the experiment (Fig. 6), an intact adult butterfly was placed in an arena where it could walk freely. The lid of the container was composed of a thin polyethylene membrane that is transparent to mid-infrared radiation (to facilitate observation using a thermal camera). The temperature at which this experiment was recorded as the threshold temperature, \( T_{threshold} \). The laser power was controlled such that the temperature increased from 25 to 45 °C in 7–8 s. The rate at which the temperature increased was chosen to prevent over-stimulation that could either startle butterflies into flight or cause irreversible damage to the sensory neurons in their wings.

Basking experiment. In the experiment (Fig. 7), a dark-adapted adult butterfly was placed in an arena with a thermally transparent window. The temperature of the arena was controlled by a Pelletier thermoelectric cooler. A Xenon lamp, which simulates sunlight, illuminated the substrate of the arena uniformly from above with an angle of –15° to the vertical. All the species used in this experiment perform "lateral basking" when the lamp was turned on, the butterflies closed their wings and tilted sideways to expose their thoraces and ventral wing surfaces to the illumination. The beginning of a lateral basking event was scored when the wings were tilted to be approximately perpendicular to the direction of illumination. The cessation of basking was scored when the butterflies restored the vertical orientation of the wings. As the temperature of the region illuminated by the laser beam increased to a threshold value, the butterfly would move to avoid the beam, and this response was scored as a displacement event (Supplementary Movies 8, 9). The temperature at which this experiment was recorded as the threshold temperature, \( T_{basking} \). The laser power was controlled such that the temperature increased from 25 to 45 °C in 7–8 s. The rate at which the temperature increased was chosen to prevent over-stimulation that could either startle butterflies into flight or cause irreversible damage to the sensory neurons in their wings. Each thermal camera was used to record videos of the response, from which \( T_{limiter} \) was determined. Wing veins have thermal emissivity close to unity, so direct readings from the thermal camera were used as the accurate temperature of the wing vein. Similarly, we used the thermal camera to record temperatures during basking both in the field and lab, again taking advantage of the fact that both the thorax and the wing veins have thermal emissivity approaching unity.

Different butterfly species move their wings in characteristic ways to avoid thermal stimuli applied to the wings. For example, butterflies in the family Papilionidae with wings opened initially at rest tend to flap their wings, and butterflies in the family Lycaenidae with wings closed initially at rest tend to walk forward or turn around to avoid the laser spot (Supplementary Movie 8). Once the movement of the butterfly was observed, we blocked it using a small laser beam to prevent further increase of the wing temperature and possible damage caused by overheating.

Care was taken to ensure that the response was solely due to the temperature of the illuminated wing region exceeding a threshold value. Specifically, the laser spot was never positioned on wing regions close to the base of the wing, which is also close to the insect's eyes. Movement and lighting changes in the surrounding environment were minimized to prevent accidentally stimulating the butterfly during the experiment. Control experiments were conducted that indicated butterflies may not be able to see or perceive the laser spot on their wings during the laser-spot heating experiments (Supplementary Fig. 43).

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

The data source underlyings Figs. 1, 2, 4, 6, and 7 and Supplementary Figs. 4-9, 32, 33, 35, 37-39, 42, and 43 are provided as a Source Data file.
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Author contributions

C.C.T., G.D.B., N.E.P. and N.Y. conceived and designed the experiments. C.C.T., N.N.S., J.N.P., N.E.P. and N.Y. carried out the experiments. C.C.T., R.A.C., C.R., N.E.P. and N.Y. developed theoretical models, conducted numerical simulations, and analyzed the data. C.C.T., R.A.C., N.E.P. and N.Y. wrote the manuscript, with input from all co-authors. All authors discussed the results and commented on the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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Correspondence and requests for materials should be addressed to N.E.P. or N.Y.

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