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Mueller matrix characterizations of circularly polarized reflections from golden scarab beetles

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Circularly polarized light (CPL) reflections are rare in nature. Only a few animal groups—most notably certain stomatopod crustaceans and certain beetles in the family Scarabaeidae—are known to reflect CPL from incident unpolarized light. Here, we examine five species of metallic scarabs in the genus *Chrysina* that, to the naked human eye, look remarkably similar. Using a spectropolarimetric reflectometer to characterize the complete Mueller matrix elements of the beetles' elytral surfaces, we found that four of the five species were strongly left-handed circularly polarized (LHCP), and only one scarab species, *Chrysina resplendens*, had an overall lower degree of polarization and switched from LHCP to right-handed circularly polarized reflectance depending on wavelength. © 2020 Optical Society of America

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1. INTRODUCTION

Scarab beetles in the genus *Chrysina* (Coleoptera: Rutelinae: Scarabaeidae) have been prized for centuries due to their shiny, jewel-like appearance, and more recently have served as inspiration for biomimetic applications related to structural color [1–8]. These jewel scarabs (not to be confused with jewel beetles in the Buprestidae family) recently underwent a taxonomic revision in which the former genus *Plusiotis* was reclassified as the genus *Chrysina* [9], derived from the Greek word *chrysos* meaning gold. Not only do many of these *Chrysina* beetles have mirror-like golden surfaces, they also demonstrate the rare optical phenomenon of reflecting circularly polarized light (CPL). Ever since the discovery that the golden scarab beetle, *Chrysina resplendens*, reflects CPL [10], there has been interest in understanding how widespread this phenomenon is and what features of the cuticle contribute to CPL reflectance.

Interestingly, most *Chrysina* beetles examined thus far are known to strongly reflect left-handed circularly polarized (LHCP) light [11], but Michelson first noted that *C. resplendens* also reflected right-handed circularly polarized (RHCP) light [10]. The helicoidal layers in the cuticle responsible for the CPL reflectance have since been identified [7,12,13]. *C. resplendens* is unique thus far among other studied species in the *Chrysina* genus because its cuticle has two left-handed helicoidal regions,

separated by a unidirectional layer of birefringent material that acts as a half-wave plate retarder [12,14]. Over a certain wavelength range, this arrangement results in RHCP traveling through the first helicoidal layer, followed by a switch to LHCP upon the first transmission through the unidirectional retarder layer, then reflection by the second helicoidal layer, and then a switch back to RHCP reflectance on the return path through the unidirectional layer [12,14].

Even though transmission electron microscopy has revealed the ultrastructure of the cuticle of several specimens of C. resplendens that results in CPL, we still lack a complete understanding of how these chirped helical layers affect CPL signatures. Different studies have reported wide variations in the handedness and spectral response of CPL reflections in C. resplendens [11,15–21]. Some of these variations are the result of differing methods for measuring the CPL reflectances. For example, Hegedüs et al. used imaging polarimetry to characterize whole specimens of C. resplendens and other scarabs at blue, green, and red wavelengths [15]. Pye examined how widespread the phenomenon of CPL reflectance is in many groups of scarabs including C. resplendens, but this study did not distinguish wavelength-dependent effects [11]. Hodgkinson et al. [16] used ellipsometry methods to determine that the Mueller matrices for C. resplendens correspond with both right-circular and left-circular polarizers in the 400 to 900 nm

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wavelength range. Fernández del Río [17] and Arwin *et al.* [18] also conducted Mueller matrix spectroscopic ellipsometry, but their small (<100 µm) spot size and high angle of incidence (between 25° and 75°) relative to the surface plane measurements may not give us information about what another beetle or predator or prey may see as compared to Mueller matrix measurements that average over the entire elytral surface of the beetle. Additionally, Mendoza Gálvan *et al.* later used data measured from electron microscopy images (which may be influenced by tissue shrinkage that occurs during preparation) to model an expected Mueller matrix for *C. resplendens*, and showed RHCP reflectance peaking at over 800 nm [19] as compared to Goldstein's measured Mueller matrix that showed RHCP peaking at approximately 600 nm [20,21].

Because studies have suggested that certain metallic or golden beetles besides C. resplendens may also have wavelengthdependent CPL reflectance that may switch handedness, our study aims to further characterize these polarization properties, such as degree of polarization, handedness, and ellipticity, in additional species of scarab beetles. Our study is the first direct follow up to Goldstein's study [21] using the same spectropolarimetric reflectometer to measure the complete Mueller matrices of five species of golden scarabs (all from similar habitats in mountainous forests in Central America). While C. resplendens is arguably the beetle species that has received the most attention in the past and is included in this current study, we also include an additional four golden species that have never been measured before with this spectropolarimetric reflectometer method. By taking measurements that average over the entire elytral surface at normal incidence, we can compare our results with those obtained from ellipsometry methods and with those studies that characterize handedness based on viewing the beetles through polarizing filters in order to gain new insights into any ecological relevance of these polarization signatures.

2. EXPERIMENTAL DETAILS

A. Specimens

Spectral Mueller matrices were measured of five species of golden scarab beetle in the genus Chrysina: Chrysina argenteola [22], Chrysina batesi [23], Chrysina chrysargyrea [24], Chrysina resplendens [23], and Chrysina strasseni [25]. These beetle specimens were obtained on loan from the Florida State Collection of Arthropods in Gainesville, Florida, and had originally been collected in cloud forests in Costa Rica, Colombia, or Guatemala [26]. While we only report the complete Mueller matrices for one individual from each of the five species, we additionally report the Mueller matrices from eight specimens of a species previously measured by Goldstein, Chrysina gloriosa [9,27], to estimate the amount of variability that can exist within a single species. All beetle specimens were photographed using a Canon EOS 40D digital camera, with no polarizing filter initially, and then subsequently using a RHCP filter prior to the specimen being mounted and aligned in the spectropolarimetric reflectometer (Fig. 1).

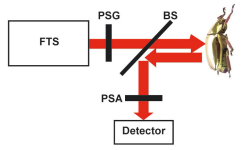


Fig. 1. Spectropolarimetric reflectance measurement configuration. FTS is the Fourier transform spectrometer, PSG is the polarization state generator, BS is the beam splitter, and PSA is the polarization state analyzer. The red arrows represent the light beam (that is larger than the beetle) hitting the dorsal curved surface of the beetle sample at normal incidence in this lateral view.

B. Dual Rotating Retarder Mueller Matrix Spectropolarimetric Reflectometer

The spectropolarimetric reflectometer used herein for Mueller matrix measurements is an instrument that measures spectral polarization properties of materials in transmission or reflection modes (Fig. 1). Its core is based on a commercial Fourier transform spectrometer (FTS), and it was previously described [28], patented [29], and used for precision measurements [20,21] for a variety of applications. For purposes of the measurements described here, the spectrometer is used with one source, an Oriel xenon lamp, and one silicon detector, such that the light collected is light that has been retroreflected at normal, or close to normal, incidence from the back of the beetles. Note that the light source has slightly lower intensity at shorter wavelengths, and the detector is also less responsive at shorter wavelengths. Together, these contribute to a lower signal and higher noise at shorter wavelengths compared to other wavelengths, as demonstrated in previous measurements [20,21,28]. The spectrometer serves as a radiation source for the polarimetric portion of the instrument and is operated in the conventional absorption spectroscopy mode. The radiation generated by the spectrometer is brought out through the spectrometer's external port.

Figure 1 shows the basic optical schematic of the instrument for monostatic reflectance measurements. The beetle sample was mounted vertically a short distance past the beam splitter (BS). The spot size was larger than any beetle specimen, which was pinned against a piece of black foam smaller than the beetle and so the beetle appeared suspended in air, such that any stray light that did not reflect off the dorsal surface of the beetle would go off several meters in the dark room to a beam dump and not be picked up by the detector. The optical system that collects light for the detector consists of an off-axis parabolic mirror. This mirror is oriented to look toward the beam splitter and to focus light on the detector that is mounted perpendicularly to the light coming from the beam splitter. The parabolic mirror, detector, and mounting devices form the detector assembly. The spectrometer was also used without any modification of the polarization of the source radiation (i.e., no added polarization elements). In this mode, it functions as a spectral reflectometer.

To obtain spectropolarimetric measurements, a dual rotating retarder Mueller matrix polarimeter, described by Azzam [30],

is included in the system. This polarimeter consists of a polarization state generator (PSG) before the sample and a polarization state analyzer (PSA) after the sample. The PSG consists of a linear polarizer followed by a quarter-wave retarder. The PSA consists of a quarter wave retarder followed by a linear polarizer in front of the detector assembly. Although achromatic retarders are used that were nominally quarter wave in the spectral region being measured, the exact retardance is not critical since errors in retardance from quarter wave are known from the calibration of the instrument and are compensated for during processing of the sample measurements (for complete calibration details, see [28,31]). When the retarders are rotated in a 5:1 ratio, all 16 elements of the sample Mueller matrix are encoded onto 12 harmonics of the detected signal; these are then Fourier analyzed to recover the Mueller matrix elements. Other previous implementations of this Mueller matrix polarimeter have been described elsewhere [32].

C. Theory and Calculations

The Mueller matrix formalism determines the reflected Stokes vector S_r in response to any incident Stokes vector S_i according to the equation

$$S_r = \mathbb{M}S_i, \tag{1}$$

where the Mueller matrix is

$$\mathbb{M} = \begin{pmatrix} m_{11} & m_{12} & m_{13} & m_{14} \\ m_{21} & m_{22} & m_{23} & m_{24} \\ m_{31} & m_{32} & m_{33} & m_{34} \\ m_{41} & m_{42} & m_{43} & m_{44} \end{pmatrix},$$
(2)

where m_{21} is the linear polarizance in the horizontal/vertical, m_{31} is the linear polarizance at $\pm 45^{\circ}$, and m_{41} is the circular polarizance [33].

If unpolarized light illuminates the elytral surface of the beetles, our incident Stokes vector is

$$S_i = \begin{pmatrix} 1 \\ 0 \\ 0 \\ 0 \end{pmatrix}, \tag{3}$$

and therefore the reflected beetle Stokes vector is

$$S_r = \begin{pmatrix} m_{11} \\ m_{21} \\ m_{31} \\ m_{41} \end{pmatrix}. \tag{4}$$

By making spectral measurements using the spectropolarimetric reflectometer, all Mueller matrix elements are wavelength dependent and span visible wavelengths from 0.4– $0.7~\mu m$. This permitted us to calculate spectral polarization-related metrics. We determined the polarization state and degree as a function of wavelength. These results showed the amount of circular polarization reflected from the entire elytral surface of the beetle was wavelength dependent. More specifically, and similar to Arwin $\it et al.$'s study [18], we calculated the spectral degree of polarization (circular and linear) and the spectral ellipticity. Again, assuming unpolarized light illuminates the surface, these parameters of reflected light are

Degree of Circular Polarization, or $DoCP = m_{41}$, (5)

Degree of Polarization, or DoP =
$$\sqrt{m_{21}^2 + m_{31}^2 + m_{41}^2}$$
, **(6)**

and

Degree of Linear Polarization, or DoLP =
$$\sqrt{m_{21}^2 + m_{31}^2}$$
. (7)

Equations (5)–(7) combine to show that the square of the degree of total polarization is equal to the sum of the squares of the degree of linear polarization and degree of circular polarization, or

$$DoP^2 = DoLP^2 + DoCP^2.$$
 (8)

This relationship is useful for interpreting some of the results presented in the next section.

The ellipticity of the polarization ellipse is given by

$$e = \tan\left(\frac{1}{2}\sin^{-1}\left(\frac{m_{41}}{\sqrt{m_{21}^2 + m_{31}^2 + m_{41}^2}}\right)\right).$$
 (9)

Pure LHCP light reflectance would have a DoCP (m_{41}) value and an ellipticity value of -1, while the values for pure RHCP light reflectance would be +1. All polarization metrics were calculated from the raw unfiltered Mueller matrices (normalized to $m_{11}=1$). The matrix elements were then smoothed along the wavelength axis using a median moving average filter (order 20). Additionally, the spectral reflectances of all beetles were measured using the same setup without polarization elements. The relative reflectance is normalized to the maximum spectral reflectance and smoothed using a median moving average filter (order 16). Thus, the reported reflectance measurements of the five beetle species are by necessity relative comparisons of the spectral peaks between species, since the spot size is larger than the beetles and the curved backs of the beetles, which are of different sizes, cannot be estimated as flat uniform panels.

3. RESULTS

A. Appearance Through CPL Filter

The five species of scarab beetles from the genus *Chrysina* examined in this study were chosen for their golden appearance and for comparison of their Mueller matrices to previously published results [15–21]. Photographs of these species without any polarizing filter (top row, Fig. 2) and through a RHCP filter (bottom row, Fig. 2) revealed that four of the five species showed a loss of color (becoming dark brown) through the RHCP filter. *C. resplendens* was the only beetle of these five that did not dramatically change appearance through the RHCP filter, as has also been shown in previous studies [15,21].

B. Reflectance

The spectral reflectances of the five species of beetles show similar reflectance peaks in the yellow to red part of the spectrum

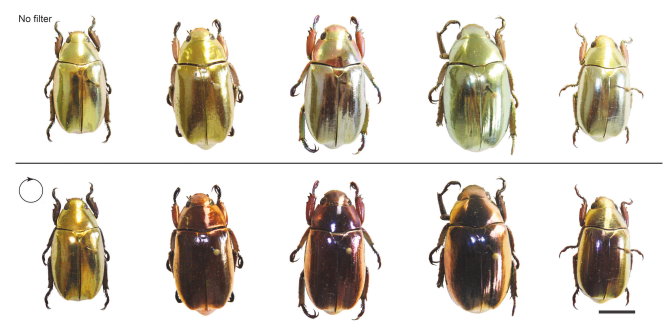


Fig. 2. Photographs of the five scarab species. The top row shows the beetles with no polarization filter. The bottom row shows the beetles through a RHCP filter. From left to right: *Chrysina resplendens, Chrysina argenteola, Chrysina chrysargyrea, Chrysina batesi*, and *Chrysina strasseni*. Scale bar is 10 mm.

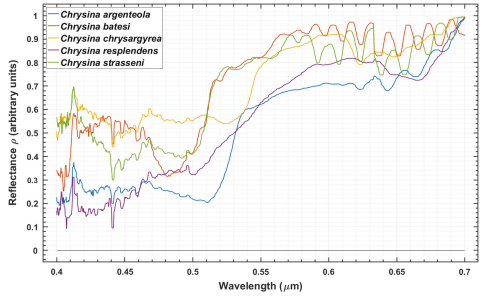


Fig. 3. Normalized reflectance spectra for the five golden beetle species at normal incidence.

that match with their "gold" appearance (Fig. 3). The oscillations indicate interference that is often seen in "metallic" colored animals that is a result of their cuticle structure [34,35]. As described in Section 2.C, these reported values are normalized values with arbitrary units, allowing for comparisons of spectral peaks, rather than absolute reflectance measurements (Fig. 3).

C. Mueller Matrix

The complete spectral Mueller matrix for *C. argenteola* is shown in Fig. 4 (see Fig. S1 for the other four species measured). *C. argenteola*, *C. chrysargyrea*, *C. batesi*, and *C. strasseni* were

all similar to Goldstein's measurement of a different metallic (silver) species, *C. clypealis* [21]. *C. resplendens* was the only species out of the five examined that shows additional features in the Mueller matrix (Fig. S2 C). These are described in more detail along with other derived metrics below. Every gold species other than *C. resplendens* resembled a textbook Mueller matrix example of a wavelength-independent homogenous left-circular polarizer over most of the visible spectrum, i.e.,

$$\begin{pmatrix} 1 & 0 & 0 & -1 \\ 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 \\ -1 & 0 & 0 & 1 \end{pmatrix}.$$

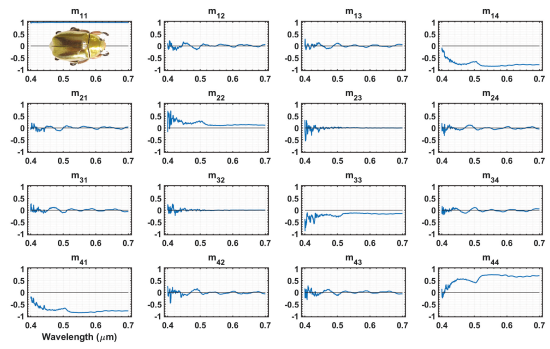


Fig. 4. Mueller matrix spectra for *C. argenteola* at normal incidence.

D. Left- and Right-Handedness and Other Polarization Parameters

First, to determine the amount of variability that can be found within one species of beetle, we measured the Mueller matrix and calculated derived metrics for eight specimens of *C. gloriosa*—a species previously reported in [21] (Fig. S2). Because overall variation between specimens of this species was

low (Fig. S2), the following results are reported only from a single specimen from each of the five golden species.

We explored derived metrics from the spectral Mueller matrix that are believed to be of biological significance: the degree of circular polarization (DoCP), ellipticity (e), degree of polarization (DoP), and degree of linear polarization (DoLP) (Fig. 5). The DoCP is the element m_{41} . Values below zero indicate LHCP, and values above zero indicate RHCP. The ellipticity

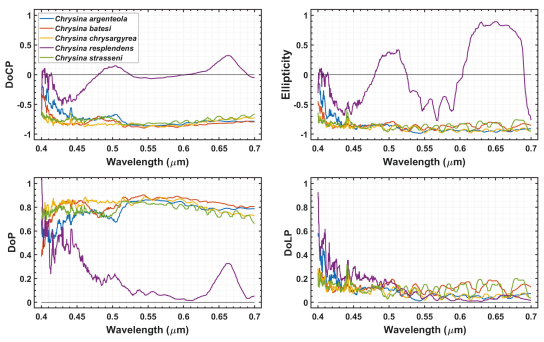


Fig. 5. Polarization metrics calculated from the Mueller matrices for the five species of beetles. Clockwise from top left: the degree of circular polarization (DoCP), ellipticity, degree of linear polarization (DoLP), and degree of polarization (DoP).

also tells us how circular the polarization is, with pure LHCP values at -1 and pure RHCP values at +1. While it may seem that DoCP and the ellipticity are closely related, there are cases where DoCP can be small but ellipticity is still close to one (pure circular). We must also consider how much of the total reflected light is polarized (DoP) which includes both DoCP and DoLP [see Eqs. (5)-(8)].

Four of the five beetle species showed similar signatures, with DoCP values between -0.7 and -0.9 and ellipticity values between -0.8 to -1.0 (Fig. 5). This indicates that these beetles were strongly LHCP spanning most of the visible spectrum. *C. resplendens* was the only species of the five that showed any wavelength-dependent shift to RHCP (Fig. 5).

C. resplendens showed LHCP from 400 to 480 nm, then switched to RHCP from 480 to 525 nm. Then there was another switch back to LHCP from 525 to 600 nm, and another switch to RHCP from 600 to 690 nm (Fig. 5). These wavelength-dependent switches in handedness are similar to what Goldstein [20,21] measured in a different C. resplendens specimen (see Fig. S3). During the first switch to RHCP (480–525 nm), the DoCP and ellipticity were approximately 0.15 and 0.4. During the second switch to RHCP (600–690 nm), the DoCP and ellipticity were approximately 0.33 and 0.9.

Interestingly, *C. resplendens* has a low DoP and low DoLP whereas the other four species have a very high DoP and low DoLP, consistent with their high "amount" of LHCP (where DOCP approaches -1). Referring to Eq. (8), *C. resplendens* shows a low DoLP and a low DoP, which must indicate low DoCP (i.e., closer to 0). The other four golden species show a low DoLP and a high DoP, which must indicate a high DoCP (i.e., closer to -1 in this case). Therefore, while *C. resplendens* does demonstrate RHCP, the overall DoP is very low relative to the other four species. This result is also consistent with the RHCP-filtered photographs (Fig. 2), where *C. resplendens* does not change appearance as dramatically as the other four species.

4. DISCUSSION

A. Reflectance and Polarization Properties of the Beetles

While the five species of golden scarabs examined in this study look similar to the naked human eye (Fig. 2), *C. resplendens* is clearly distinct from the other four species in several ways. Beginning with reflectance (Fig. 3), *C. resplendens* shows a more gradual reflectance increase spanning 500 to 600 nm. The other four species exhibit a "cuton" phenomenon at ~500, 520, or 540 nm and show a rapid increase in reflectance over the 500 to 540 nm spectral range. *C. batesi* and *C. strasseni* appear to have the same "cuton" and show similar interference features at higher wavelengths (580 to 700 nm), suggesting a similar cuticle ultrastructure.

Our measurements of polarization properties further reveal that *C. resplendens* is unique among the metallic beetles measured. *C. argenteola*, *C. batesi*, *C. chrysargyrea*, and *C. strasseni* have a high, nearly constant, left-handed DoCP spanning most of the visible spectrum. Together, the DoCP and the ellipticity indicate the four species have a very high amount or DoCP of \sim 0.8 and ellipticity of \sim 0.9. Since \sim 1 for either of these metrics indicates pure LHCP, the four species are technically

elliptically polarized; however, they are approaching a full degree of circular polarization, and the ellipse is geometrically approaching a circle. In contrast, C. resplendens has a unique DoCP signature as compared to those other four species. The DoCP is relatively very low (\sim -0.4 to 0.3) as compared to a near constant DoCP ~ -0.8 for the other four species. The DoCP spectrally shifts twice from left-hand elliptically polarized to right-handed elliptically polarized. This spans an ellipticity from -0.75 (left-hand elliptical) to +0.4 and +0.9 (right-hand elliptical) as a function of wavelength. The C. resplendens is the only beetle observed in our study that exhibited these properties. There are slight differences in the measurement results between the C. resplendens specimen measured in this study and a different individual measured in 2006 by Goldstein [21]. These differences can likely be explained by nutritional or environmental conditions that affect the exact composition of the cuticle (Fig. S3). The overall trend of the switch between LHCP and RHCP is demonstrated in both specimens (Fig. S3).

These spectral transits of *C. resplendens* into the RHCP regime add to the total DoP. This is prominently noted by a large spectral DoP feature approaching 0.33 at 663 nm and one of lesser prominence of 0.24 at ~500 nm. Additionally, *C. resplendens* has a slightly higher DoLP of 0.15–0.5 (400–500 nm) and a slightly lower DoLP of 0–0.1 (550–700 nm) as compared to the other four species. The other four species have a very low DoLP of 0.05–0.20 spanning the entire visible spectral region. For low DoLPs, the total DoP will be dominated by the behavior of the DoCP.

B. Comparison of Methods for Determining Mueller Matrices in Beetles

While we have not yet performed transmission electron microscopy to examine the ultrastructure of the cuticle of these five particular specimens, other studies have examined *C. resplendens* [12,14] and C. chrysargyrea [7] as well as other LHCP species of scarabs [13]. Based on our Mueller matrix measurements, we expect that the four specimens of C. argenteola, C. batesi, C. chrysargyrea, and C. strasseni have similar cuticle architecture and that *C. resplendens* is the only golden species with a different structure that consists of a birefringent layer that is sandwiched between two helicoidal layers. Interestingly, while we can use measured Mueller matrices to infer characteristics of the cuticle ultrastructure, we suggest that going in the opposite direction or using ultrastructure measurements alone to infer the Mueller matrix may result in inaccurate spectral peaks for both LHCP and RHCP reflectances (e.g., as reported by [19]). This may be due to the fact that exact thickness of layers in the cuticle of beetle specimens (dead or alive) can be difficult to determine because of the shrinkage of tissues that occurs when preparing a specimen for examination using electron microscopy.

Additionally, visual observations in the past generally have declared the characteristic of LHCP or RHCP by use of a like polarizer without reference to the degree of left or right elliptical character [11,15]. Our work provides a quantitative assessment of the amount that these five species of golden scarab beetles are elliptically/circularly polarized. Importantly, our measurements of Mueller matrix spectra account for the optical signature of the entire dorsal surface of the beetle. Because beetles have curved

backs, "normal incidence" refers to the fact that the beam of light (larger than the beetle) is positioned normal to the entire specimen. Since Goldstein presented the first complete Mueller matrix data for three species of scarab beetles [21], several studies have relied on ellipsometry methods to obtain Mueller matrix spectra for different scarabs at oblique incidences, using a small spot size (<100 µm) on the beetle's elytral surface [16–18,36–39]. It is interesting that our measurements of C. argenteola at normal incidence differed significantly from Arwin et al.'s [18] ellipsometer measurements, which were collected at non-normal incidence. These ellipsometer measurements at specific angles are most valuable in the determination of the internal structure along with validated models based on physical measurements of the layered structures; however, they are not necessarily representative of what one sees on observation of the beetles through circular polarizers by eye or with photographic equipment, or what one measures with the spectropolarimetric reflectometer with the specimens reflecting light off the entire surface. Our measured DoCP for C. argenteola was close to -1 across most of the visible spectrum, quite different from Arwin et al.'s measured DoCP that, at angles above 45°, showed a switch from LHCP to RHCP values. At 65°, C. argenteola showed high RHCP DoCP values between +0.6 and +0.8 at wavelengths between 600 and 800 nm [18]. That the handedness can change depending on angle of incidence requires further investigation. In C. resplendens, for example, we expect that the optical pathlength through the wave retarder in the beetle elytra will not be the same at different angles.

C. Biological Significance

The above comparisons bring into question what ecologically relevant polarization signals can be determined from the various methods of measuring Mueller matrices in beetles. While ellipsometry measurements can certainly reveal information about the nanostructure of the beetle's cuticle and other useful optical information, they may not be relevant for understanding what a beetle or its predators or prey would actually see when viewing another beetle against a leafy background in a montane forest canopy. The spectropolarimetric reflectometer is designed to approximate the polarization and reflection signatures that may be observed in a natural environment, such as in the case of a beetle flying above another beetle and looking down at its dorsal surface.

Currently, we have little to no information about whether any of these golden scarab beetles may be able to distinguish any CPL signature. Overall, CPL reflections are rare in nature. However, recently marine crustaceans called stomatopods have been shown to use CPL signals in communication [40]. In fact, these crustaceans are able to distinguish LHCP from RHCP [41]. Whether or not CPL is detectable or serves a visual function in beetles remains unclear. Brady and Cummings [42] claimed that the beetle *C. gloriosa*, a strongly LHCP beetle, is able to differentiate unpolarized light from CPL, but the beetle *C. woodi*, a more elliptically polarized beetle, was not able to differentiate CPL. This suggests that the degree of polarization and ellipticity may play an important biological role. However, this study did not eliminate brightness cues as a confounding factor. Miao *et al.* [43] showed that *Anomala copulenta* use green

cuticle-reflected light for mate choice, but this approach did not allow the effects of color and polarization to be separated. Blahó *et al.* [44] tested four different species of scarab beetles and found no evidence that they were attracted to CPL when feeding or looking for mates or conspecifics.

Our study shows that C. resplendens has a very low DoP and DoLP, whereas the other four species have very high DoP and low DoLP consistent with a "high toward the left" DoCP (approaches -1). Whether or not the DoP exhibited by any species of golden scarab is enough to be detectable by the beetles requires further investigation into their visual capabilities, including their spectral sensitivities. For example, most insects are not able to see red; yet, all the beetles we measured show spectral reflectance peaks in the yellow to red part of the spectrum, and the second switch to RHCP in C. resplendens takes place around 650 nm. It was recently shown that an increased level of specular reflection of leaf backgrounds contributes to the survival of a certain species of shiny iridescent beetle, Sternocera aequisignata [45]. The camouflage of this beetle's iridescence is enhanced, at least according to human vision, by the beetle choosing glossier backgrounds [45]. Additionally, Feller et al. [46] suggested that the golden reflectors in C. resplendens illustrate that intrinsic polarization properties can function to improve the overall reflectivity of the structure. These findings beg the question about whether our studied scarab species are under selection to increase their golden reflectivity to appear camouflaged to predators but perhaps use circular polarization (which is highly unlikely to be seen by any bird or mammal predator) as a covert signal.

Exploring whether any scarab beetles are actually sensitive to CPL and whether their eyes possess any potential morphological mechanisms for detecting and analyzing CPL is an important next step before we can then address questions regarding whether circular polarization functions as a covert signal. Generally, *Chrysina* species are found to occur in sympatric assemblages [2]; thus, the need to distinguish between closely related species may be an important evolutionary driver to CPL reflectance patterns.

5. CONCLUSIONS

This study compared five species of golden scarabs that—to humans at least—look similar and that have overlapping habitat distributions in mountainous forests in Central America. Previously, individual studies have been undertaken to characterize one species, or compare a few species of different colors, or examine the Mueller matrices at specific high angles of incidence. We wanted to follow up on Goldstein's 2006 study to further explore the polarization properties of these "gold bugs" to determine if a common feature of being a golden scarab is having circular polarization that switches handedness dependent on wavelength. We conclude that C. resplendens is the only golden scarab species known that demonstrates a switch from LHCP to RHCP dependent on wavelength. Our measurements account for the entire elytral surfaces of the beetles, which is representative of what another beetle or predator would see. We conclude that C. resplendens has different polarization signatures than the other golden scarabs we measured, and future work will

examine their eyes to determine if they can discriminate LHCP or RHCP light.

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Disclosures. The authors declare no conflicts of interest.

See Supplement 1 for supporting content.

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