

## Micrographia

# Polarization Microscopy and Infrared Microspectroscopy of Integument Coverings of Diapausing Larvae in Two Distantly Related Nonsocial Bees

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### Abstract

The larvae of the two distantly related nonsocial bees *Ericrocis lata* (Apidae) and *Hesperapis (Carinapis) rhodocerata* (Melittidae), which develop mostly under arid desert areas of North America, and that differ in that they either spin (*E. lata*) or do not spin (*H. rhodocerata*) protective cocoons before entering diapause, produce transparent films that cover the larval integument. To understand the nature of these films, their responses to topochemical tests and their characteristics when examined with fluorescence and high-performance polarization microscopy and microspectroscopy were studied. A positive staining by Sudan black B, birefringence of negative sign, and a Fourier transform-infrared (FT-IR) spectrum typical of lipids were detected for the integument covering of both species. The FT-IR signature, particularly, suggests a wax chemical composition for these lipid coverings, resembling the waxes that are used as construction materials in the honey cells produced by social bees. Considering the arid environmental conditions under which these larvae develop, we hypothesize that their covering films may have evolved as protection against water depletion. This hypothesis seems especially appropriate for *H. rhodocerata* larvae, which are capable of undergoing a long diapause period in the absence of a protective cocoon.

**Key words:** integument covering, wax, nonsocial bees, optical anisotropy, FT-IR spectra

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### Introduction

*Hesperapis (Carinapis) rhodocerata* (Melittidae) and *Ericrocis lata* (Apidae) are two distantly related nonsocial bees. *H. rhodocerata* has been found to nest in normally dry desert areas of North America (Rozen, 1987, 2016), whereas the cleptoparasitic bee *E. lata* has been reported to use the nests of ground-nesting bee hosts like *Centris (Paracentris) caesalpiniae* in SE Arizona (Rozen & Buchmann, 1990), close to the *H. rhodocerata* sites and under similar climatic conditions (Rozen, 2016). Although these species differ in that their larvae either spin (*E. lata*) or do not spin (*H. rhodocerata*) a protective cocoon before entering diapause, they both produce a thin transparent film that covers the integument of the postdefecating larvae (Rozen & Buchmann, 1990; Rozen, 2016). This material takes on the exact contour of the larval body surface and when teased with forceps, it flakes off (Rozen, 2016).

When one of us (J.G.R.) tested the cell wall of *H. rhodocerata* larvae with a water droplet, this wall was found to be readily absorbent, which is an unusual finding because solitary bees that do not spin cocoons usually provide water-retardant brood cell linings. In the particular case of these bee larvae, there was

indication that the cell wall had received no special treatment to protect the diapausing larva from desiccation during its 5-year diapause.

The integument covering material was thus suspected to play some role in protecting the diapausing larvae of these bees especially in the case of absence of a protective cocoon. Considering that the composition and the structural organization of these covering films have not yet been determined, in the present study these materials were analyzed using topochemical methods, fluorescence and high-performance polarization microscopy, and infrared (IR) microspectroscopy. These procedures have recently been seen to be useful for understanding the nature and organization of several biopolymers, including silk elements in bee cocoon walls (Mello et al., 2016).

### Materials and Methods

#### Insects

Mature larvae of *H. rhodocerata* (Cockerell) (Hymenoptera: Apoidea: Melittidae) were collected from a nesting site that was 28 miles south of Animas, Hidalgo Co., New Mexico, on September 8, 2015, by J.G.R. and B. Lawley. *E. lata* (Hymenoptera: Apoidea: Apidae) larvae were excavated from host bee nests on May 17, 1989, from 5 miles east of Sahuarita, Pima Co., Arizona, by J.G.R. (Rozen & Buchmann, 1990). Wax flakes were washed off mature larvae and then dried.

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### Topochemistry and Fluorescence Microscopy

A topochemical test for proteins was performed by treating the bee integument coverings with a 0.025% Ponceau SS (Sigma®, St Louis, MO, USA) solution in 3% acetic acid at pH 2.5 for 10 min (Vidal & Mello, 2005). Ponceau SS not only reveals proteins, but it also can be used to detect the organization of protein electropositive binding sites using polarization microscopy because this dye has the property of exhibiting linear dichroism, when bound to oriented substrates (Vidal & Mello, 2005). The periodic acid–Schiff (PAS) reaction (Lison, 1960; Kiernan, 2008) was also used, considering the possibility that glycoproteins were present in the materials under investigation. In addition, treatment with 0.025% toluidine blue (Merck, Darmstadt, Germany) solutions in 0.1 M citric acid and 0.2 M anhydrous disodium phosphate buffer at pH 4.0 and 2.5 was undertaken. Toluidine blue solutions at these pHs reveal acid glycosaminoglycans (AGAG) or protein–AGAG complexes (proteoglycans) and their stereoarrangements (Lison, 1960; Vidal & Mello, 1974, 1984; Mello & Vidal, 2003). Lipids were investigated using a 15-min staining with Sudan black B (Kiernan, 2008).

The examination of unstained fragments of the integumental coverings by autofluorescence was performed using a Carl Zeiss Axiophot 2 photomicroscope (Carl Zeiss, Oberkochen, Germany), equipped with an HBO-mercury short-arc 103 W/2 (Osram GmbH, Berlin, Germany) as a UV source and Carl Zeiss filter sets 02 (G exciter filter transmitting a  $\lambda = 365$  nm exciting light, FT 395 beam splitter, and LP 420 barrier filter) and 38 HE (BP exciter filter transmitting a  $\lambda = 470/40$  nm exciting light, FT 495 beam splitter, and BP 525/50 barrier filter).

### High-Performance Polarization Microscopy

Birefringence was investigated on unstained flakes of the integument coverings using a Carl Zeiss Axiophot 2 microscope equipped with an Axiocam HRc camera and Kontron KS400-3 software (Carl Zeiss), and an Olympus BX51 high-performance polarizing microscope (Olympus America, Center Valley, PA, USA) equipped with a Q-color 5 camera (Olympus America). Some of the birefringence images were obtained using differential interference contrast (DIC) optics (Vidal et al., 2015). Images of the birefringence intensity brilliance were captured and analyzed using Image-Pro Plus 6.3 software (Media Cybernetics Inc., Silver Spring, MD, USA), as previously described (Vidal & Mello, 2010). The birefringence sign was detected using a first-order red compensator (Axiophot 2 microscope) and DIC red band interference color (BX51 polarizing microscope) (Vidal & Mello, 2010).

### Fourier Transform-Infrared (FT-IR) Microspectroscopy

FT-IR spectra were obtained from dry fragments of the larval integument coverings using an Illuminat IR II™ microspectroscope (Smiths Detection, Danbury, CT, USA) that was equipped with a liquid nitrogen-cooled mercury–cadmium–telluride detector and GRAMS/AI™ 8.0 spectroscopy software (Thermo-Electron Co., Waltham, MA, USA). An Olympus BM51 microscope (Olympus America) with an attenuated total reflection diamond objective (magnification 36×) was part of the microspectroscope equipment. A low signal-to-noise ratio of 7,929:1 indicated the performance validation of the equipment (Vidal & Mello, 2011).

The absorbances were evaluated within a 3,500–650  $\text{cm}^{-1}$  wavenumber spectral range using a square of 25  $\mu\text{m}$  per side as the measurement site area. The spectral resolution was equal to

4  $\text{cm}^{-1}$  as quoted by the equipment supplier. The absorbances of the preparations and background were estimated using 64 scans for each individual spectral profile (Vidal & Mello, 2011). Twelve and eight spectral signatures were obtained for the cover material of *H. rhodocera* and *E. lata*, respectively. A plus-zero baseline correction was obtained for each spectrum, followed by the production of the average spectral profiles and normalization to the highest absorption peak of the spectra (2,921 and 2,916  $\text{cm}^{-1}$  for *H. rhodocera* and *E. lata*, respectively). All these procedures were performed in accordance with the instructions provided by the GRAMS/AI software.

## Results

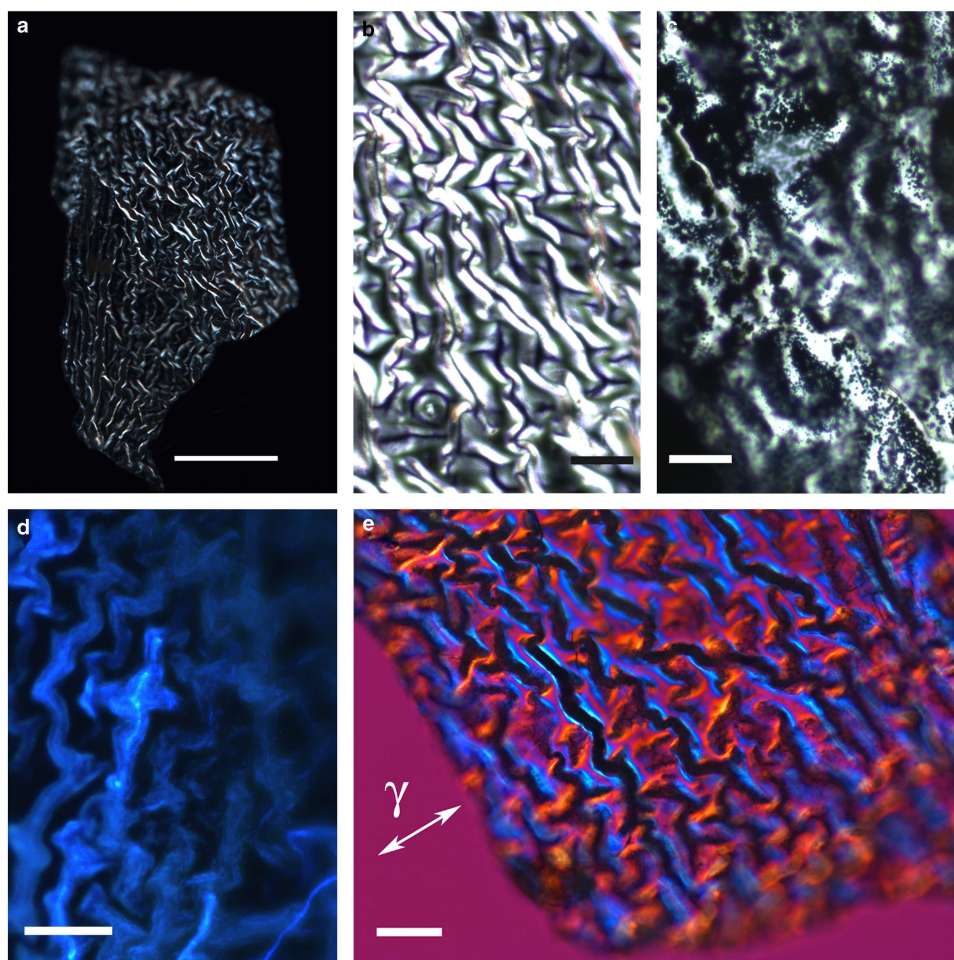
### Topochemistry, Fluorescence, and High-Performance Polarization Microscopy

The thin, transparent, and friable flakes of the integument covering from *H. rhodocera* larvae revealed a wrinkled structure that did not stain with Ponceau SS or with toluidine blue at pH 4.0 or 2.5 or with the PAS reaction, but that stained with Sudan black B and displayed blue autofluorescence (Figs. 1a–1d). In some parts of the fragments, the wrinkles obeyed a curved spatial distribution (Figs. 1a–1c, 1e), whereas in other parts the wrinkles appeared to be distributed linearly, acquiring an extended folder shape that, when examined in detail, revealed fine granular components (Figs. 2a, 2b). Circular protuberances were observed amidst the main material in some of the integument covering samples examined using fluorescence and polarization microscopy (Figs. 2c–2e). These protuberances showed an ordered spatial distribution, which suggests that they are probably imprints of pits from which setae arise. All these elements exhibited intense negative birefringence brilliance demonstrated by interference colors when they were examined with a first-order red compensator. When the long axis of the linear folders was positioned perpendicular and parallel to the  $\gamma$  direction of the compensator, blue and yellow colors were observed, respectively (Figs. 1e, 2b, 2e), thus revealing the negative sign of the birefringence exhibited by the flakes.

The integument covering fragments of *E. lata* larvae revealed a variable spatial distribution of membranous elements, resembling a lacy structure (Figs. 3a–3e) that responded to the above-mentioned topochemical tests in the same way as *H. rhodocera* samples. Mold spores, which were previously reported for the membranous structure of the larval covering of this bee species (Rozen & Buchmann, 1990), were detected in Sudan black B-stained preparations (Fig. 3c). A deep autofluorescence was revealed when using both 02 and 38 HE filter sets (Figs. 3d, 3e). Birefringence was detected in the covering flakes, highlighting the evenly distributed papillae images that correspond to places from where setae will arise (Figs. 3a, 4b, 4c). When adequately oriented with respect to the  $\gamma$  direction of the first-order red compensator, components of the flake fragments also exhibited negative birefringence (a blue interference color) (Fig. 4a). Examining the cover flakes using DIC optics revealed various interference colors that resulted from the varied distribution and thickness of the macromolecularly oriented material of the cover film (Figs. 4b, 4c).

### FT-IR Microspectroscopy

The highest absorptions in the FT-IR spectra of the larval integument covering films that were obtained from both bee species



**Figure 1.** Images of wrinkled integument cover fragments from *Hesperapis rhodocercata* larvae. Unstained samples (a,b,d,e) and a Sudan black B-stained preparation (c) are observed by polarization (a,b,e), ordinary (c), and fluorescence (d) microscopy. A polarized light image using the first-order red compensator is shown (e) to demonstrate the negative birefringence sign of the oriented covering elements, which is typical of lipid-containing substances. The blue interference color of birefringence is detected when the long axis of the filamentous distribution of the film was oriented perpendicular to the gamma direction of the compensator ( $\gamma$ ). Scale bars equal 50  $\mu\text{m}$  (a) and 20  $\mu\text{m}$  (b–e).

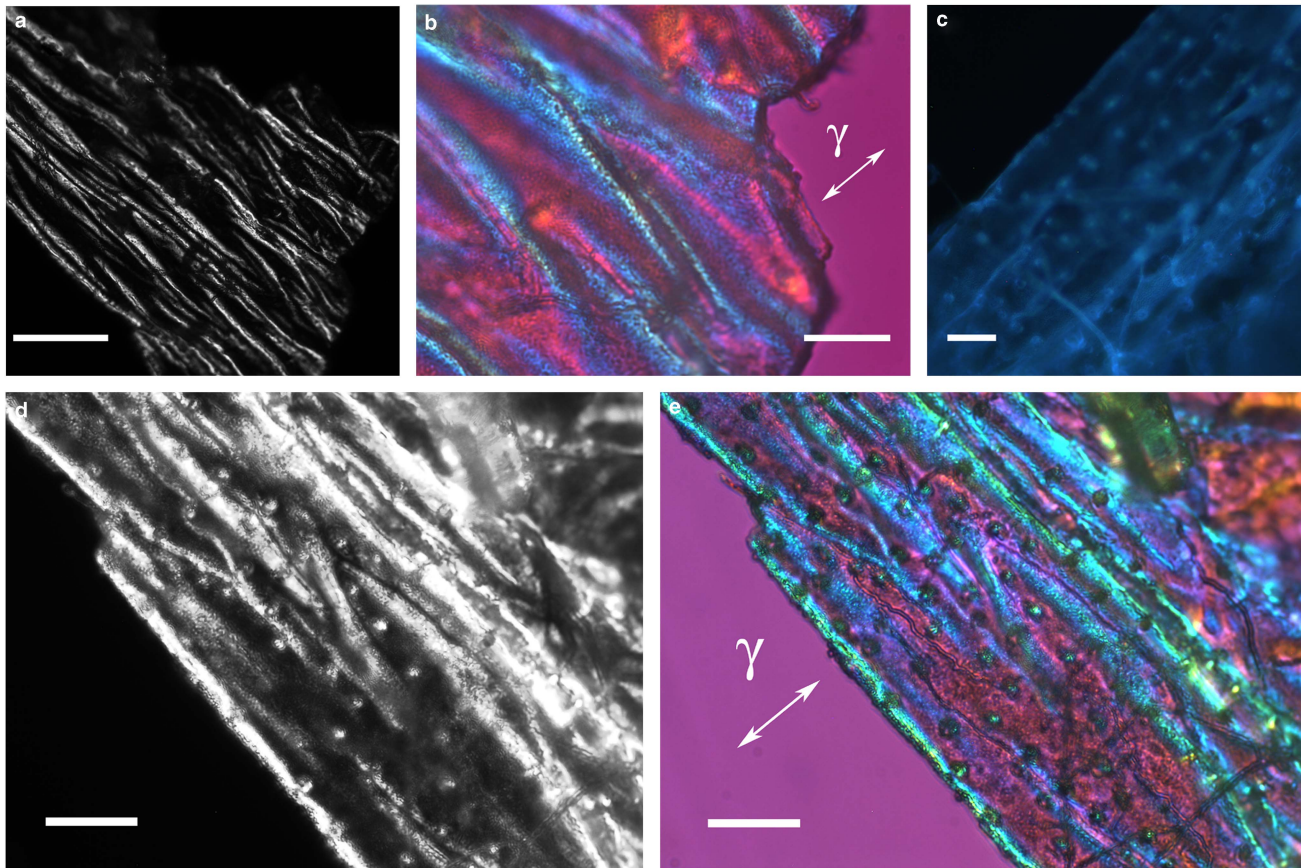
were detected as sharp band peaks in the 2,921–2,916  $\text{cm}^{-1}$  and 2,851–2,849  $\text{cm}^{-1}$  spectral regions (Fig. 5a). These bands correspond to hydrocarbon  $-\text{CH}_2$   $\nu_{\text{as}}$  and  $\nu_{\text{s}}$  vibrations, respectively, similar to the FT-IR spectral characteristics of the wax material used by *Apis mellifera* and some other social bees for building honeycomb cells (Pinzon et al., 2013; Svecnjak et al., 2015).

Regarding the FT-IR fingerprint spectral region, a low broad band at the 1,740–1,690  $\text{cm}^{-1}$  spectral window was detected, with more elevated absorbances at  $\sim 1,694 \text{ cm}^{-1}$  for *H. rhodocercata* and  $\sim 1,720 \text{ cm}^{-1}$  for *E. lata* (Fig. 5b). The irregular shape of this band contrasts with the well-defined band peak exhibited by the *A. mellifera* wax at this spectral region (Pinzon et al., 2013; Svecnjak et al., 2015). However, it resembles the spectral band assigned to ester C=O stretching bands (Pavia et al., 2009) and reported for the waxes of the stingless bees *Plebeia sp.* and *Scaptotrigona sp.* and the bumblebee *Bombus pullatus* (Pinzon et al., 2013).

When examined in detail, a sharp band peak which appeared at the 1,480–1,460  $\text{cm}^{-1}$  wavenumber region in Figure 5a, revealed two sharp components, with one at 1,473  $\text{cm}^{-1}$  and another at 1,462  $\text{cm}^{-1}$  (Fig. 5b). This characteristic was not reported for the wax spectra of *A. mellifera* and other social bees (Pinzon et al., 2013; Svecnjak et al., 2015), but was evident in the FT-IR spectrum of a Polish bee wax 10 min after the samples were heated to

melting point temperature and quenched to room temperature (Zimnicka & Hacura, 2006). In *H. rhodocercata*, the peak at 1,473  $\text{cm}^{-1}$  was more prominent than that at 1,462  $\text{cm}^{-1}$ , and the inverse occurred for *E. lata* peaks (Fig. 5b). These peaks are usually associated with the hydrocarbon  $-\text{CH}_2$  during in-plane bending or scissoring (Pavia et al., 2009; Svecnjak et al., 2015).

A very low peak at 1,377  $\text{cm}^{-1}$  was present in the spectral profile of both bee materials (Fig. 5b). It could be attributed to alcohol C–H symmetric flexion and/or deformation (Pinzon et al., 2013). Low peaks at 1,110  $\text{cm}^{-1}$  (*H. rhodocercata*, exclusively) and 887  $\text{cm}^{-1}$  (*E. lata*, exclusively) were also detected (Fig. 5c); these peaks have been assigned to C–O stretching vibrations of ethers and alcohols (Pinzon et al., 2013). One apparently sharp band peak which occurred in the 740–715  $\text{cm}^{-1}$  spectral region in Figure 5a, was found to comprise twin sharp peaks (734 and 718  $\text{cm}^{-1}$ ) (Fig. 5c). The peak at 718  $\text{cm}^{-1}$  was higher than that at 734  $\text{cm}^{-1}$  in the spectra of both bee materials (Fig. 5c). Vibrations at  $\sim 720 \text{ cm}^{-1}$  are frequently assigned to the bending (rocking) motion associated with four or more  $-\text{CH}_2$  groups in an open chain (“long-chain band”) such as that of nujol (Pavia et al., 2009). These vibrations have also been detected in the FT-IR spectra of *A. mellifera* wax (Knuutinen & Norrman, 2000; Pinzon et al., 2013; Svecnjak et al., 2015).



**Figure 2.** Images of folded integument cover fragments from *Hesperapis rhodocerata* larvae. Unstained preparations are observed by polarization (a,b,d,e) and fluorescence (c) microscopy. Polarized light images were obtained using the first-order red compensator to reveal the negative birefringence sign ( $\gamma$ ) of the covering components and the details of a fine granular distribution of birefringent elements (b,e). Autofluorescence (c) and birefringence brilliance (d,e) are also verified on circular projections that appear with an ordered spatial distribution and which are attributed to the imprints of pits from which setae will arise. Scale bars equal  $50\mu\text{m}$  (a) and  $20\mu\text{m}$  (b–e).

No characteristic bands in the  $3,600\text{--}3,200\text{ cm}^{-1}$  spectral range or at  $1,040\text{ cm}^{-1}$  that could be attributed to O–H stretching vibrations (Pavia et al., 2009) were verified in the spectral profiles of the samples from both bee species (Fig. 5a). This result resembles an observation by Pinzon et al. (2013) for the Africanized honeybee wax, and it differs from that reported for pure beeswax (Knuutinen & Norrman, 2000; Zimmnicka & Hacura, 2006) and the waxes of some other social bees (Pinzon et al., 2013).

## Discussion

A lipid-based chemical composition is assumed for the integument covering films that were obtained from the diapausing larvae of both *H. rhodocerata* and *E. lata*. This conclusion is based on the positive Sudan black B staining of such films, demonstration of negative sign birefringence, and specific FT-IR spectral profiles, that are all typical of lipids. No evidence of overall proteins, glycoproteins, and AGAG or proteoglycans was found when using the Ponceau SS dye, the PAS reaction, and the toluidine blue tests, respectively. Under the same Ponceau SS staining conditions used here, proteins stain in a multitude of materials, from vertebrate collagens and noncollagenic proteins of the extracellular matrix to insect cocoon proteins (Vidal & Mello, 2005; Mello et al., 2016). Clear images of Ponceau SS-stained proteins have been recently reported in cocoons of the bee *Lithurgus chrysurus* (Mello et al., 2016). Ponceau SS is also widely used in electrophoresis studies as a dye of choice for locating

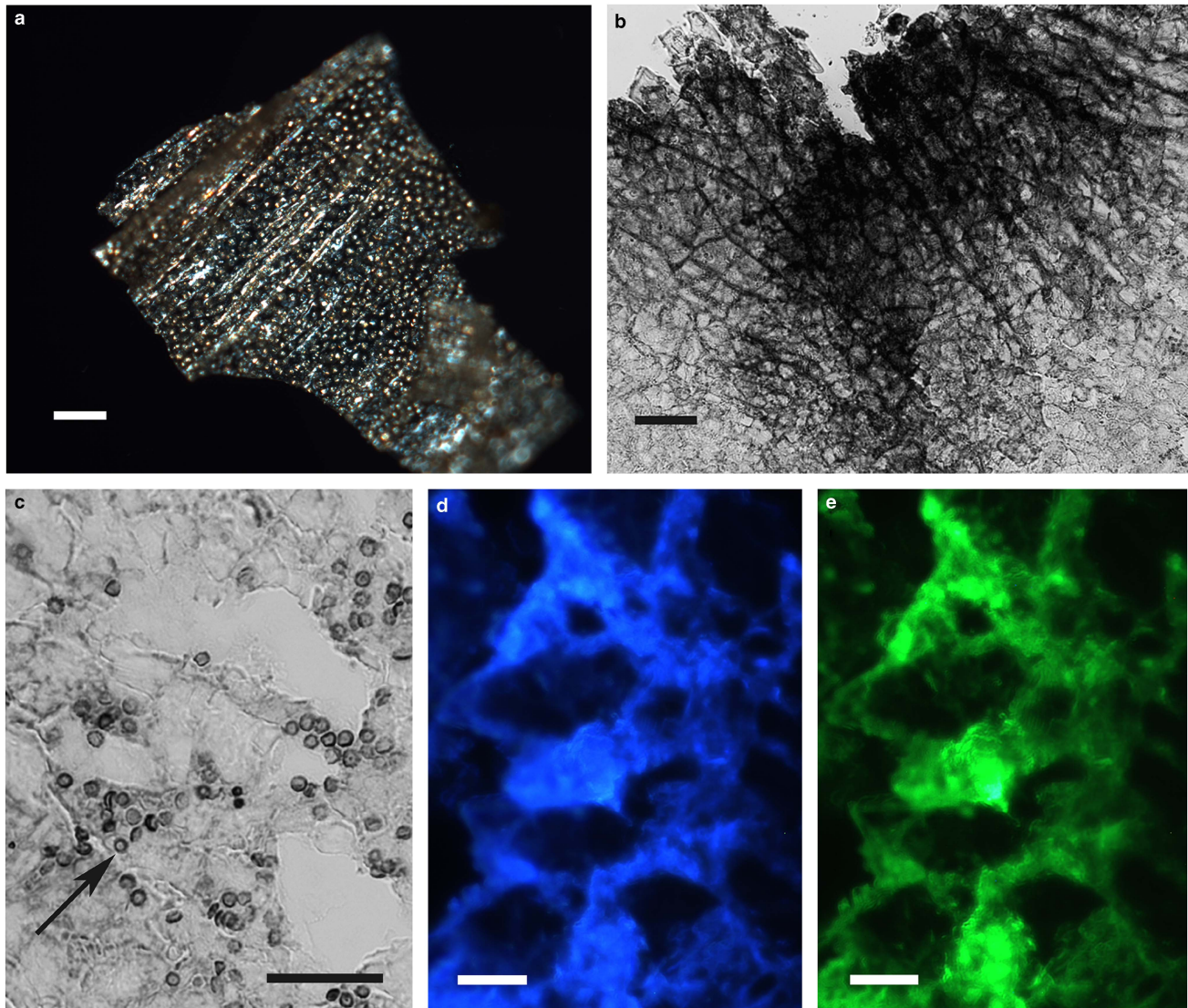
polypeptides on Western blots for blot-sequencing. Regarding PAS reaction, it is a classic assay for glycoproteins (Lison, 1960; Kiernan, 2008). Toluidine blue staining usually reveals proteoglycans in extracellular matrices, mucins, bee salivary and Malpighian tubule secretions, and cocoon walls of bumblebees and solitary bees (Mello & Vidal, 1971, 1979, 2003; Vidal & Mello, 1974, 2016; Mello, 1982; Mello & Garófalo, 1986; Mello et al., 2016).

The Sudan black B test, that provided a positive response in the integument covering films studied here, is a classic topographical assay for lipids (Kiernan, 2008).

Lipid structures with an ordered organization are characterized by negative birefringence, when studied using polarization microscopy (Schmidt, 1936; Bennett, 1967; Vidal et al., 1980). This phenomenon was detected in the integument coverings of both *H. rhodocerata* and *E. lata*.

Finally, the proposal of a lipid composition for the integument coverings of *H. rhodocerata* and *E. lata* is confirmed by analyzing the FT-IR spectral signatures of these materials. FT-IR microspectroscopy is known to offer a reliable method for qualitative analysis of lipids (Pidgeon et al., 1989; Gibbs, 1998; Zimmnicka & Hacura, 2006). FT-IR spectral profiles for proteins as assessed in several databases are different from those revealed for lipids.

The characteristic lipid bands in the IR spectra are those that are related to the aliphatic group's symmetric and antisymmetric stretching vibrations ( $3,000\text{--}2,800\text{ cm}^{-1}$ ), the ester C=O stretching vibrations ( $\sim 1,740\text{--}1,690\text{ cm}^{-1}$ ), the hydrocarbon  $-\text{CH}_2$  in-plane bending or scissoring ( $1,480\text{--}1,460\text{ cm}^{-1}$ ), the alcohol



**Figure 3.** Images of fragments of the lacy integument cover film from *Ericrocis lata* larvae. Unstained preparations are observed using polarization (a) and fluorescence microscopy (d,e), and the Sudan black B-stained samples are observed by ordinary microscopy (b,c). Mold spores recovered by intense Sudan black B-positive staining are viewed amidst the cover elements (c, arrow). Scale bars equal 20  $\mu\text{m}$ .

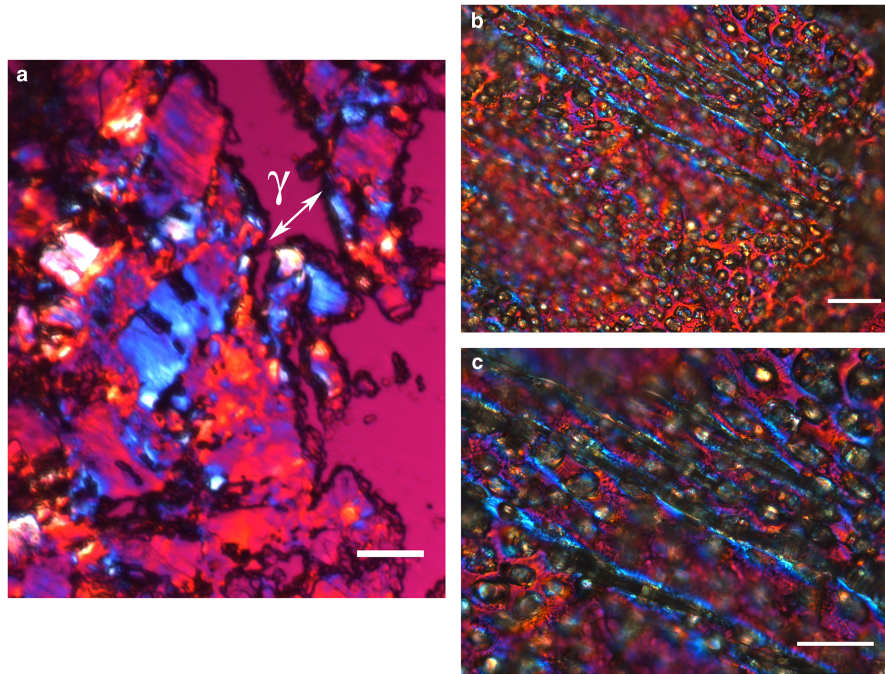
C–H symmetric flexion ( $1,377\text{ cm}^{-1}$ ), the C–O stretching vibrations of ethers and alcohols ( $1,110$  and  $887\text{ cm}^{-1}$ ), and the rocking motion associated with four or more  $-\text{CH}_2$  groups in a long-chain band ( $720\text{ cm}^{-1}$ ). All these bands are present in the FT-IR spectra of the integument cover films in *H. rhodocerata* and *E. lata* larvae.

On the basis of the FT-IR spectral results, this was the first report on the nature of the integument films that cover diapausing larvae of nonsocial bees and that are suggestive of the waxes used as construction materials in the honey cells of *Apis mellifera* and other social bees (Zimmnicka & Hacura, 2006; Pinzon et al., 2013; Svecnjak et al., 2015), although they are not fully identical. A few differences revealed between the FT-IR spectra of *H. rhodocerata* and *E. lata*, involved the vibronic frequency corresponding to ester C=O stretching bands, the prominence of the higher band peak assigned to  $-\text{CH}_2$  in-plane bending, and the intensity of band peaks assigned to ether and alcohol C–O stretching vibrations. These findings are not unexpected because of the complexity of the lipid-containing elements in bee waxes. A complex mixture of fatty acid esters and the esters of long-chain aliphatic alcohols have long been known to be present in

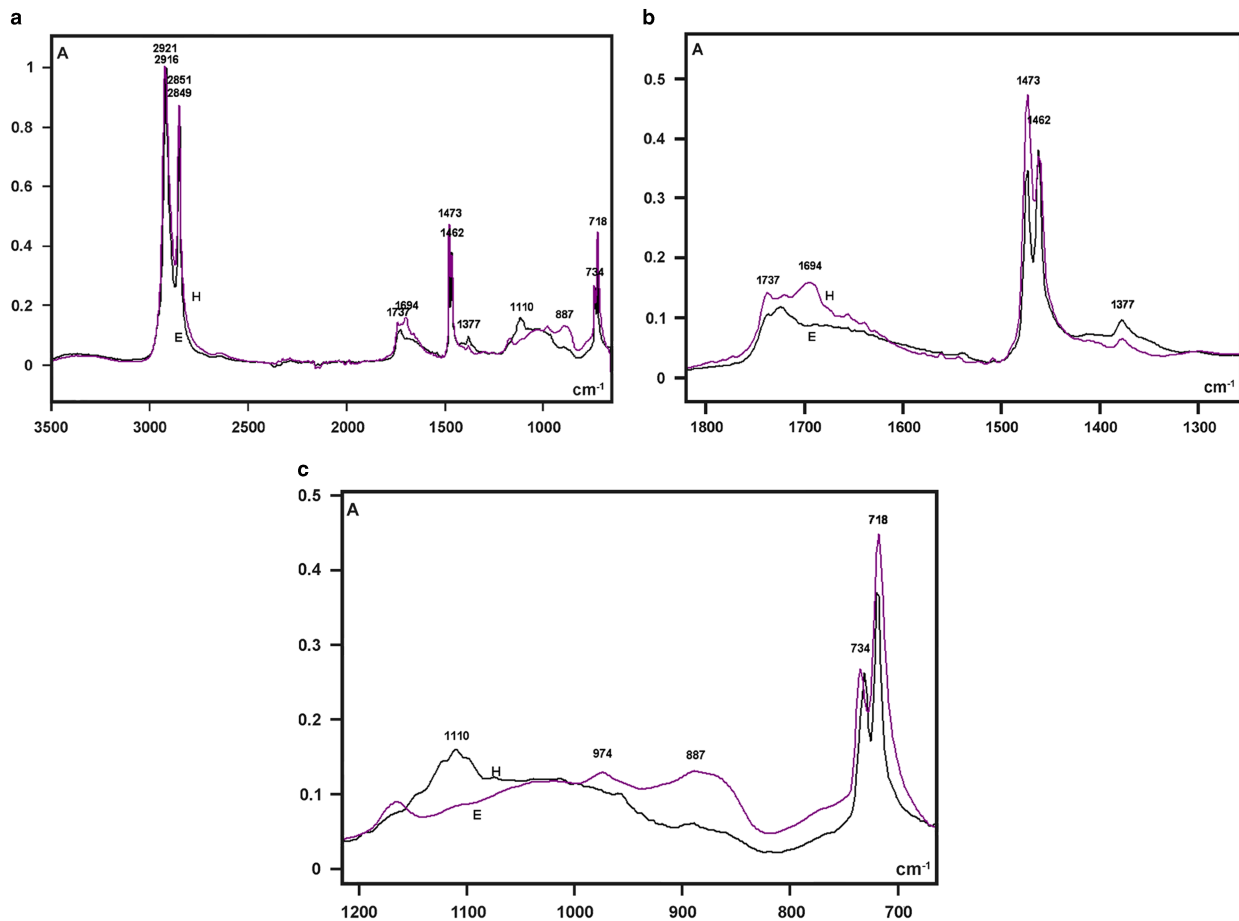
honeybee wax (Blomquist & Jackson, 1979; Tulloch, 1980). Approximately 300 different compounds have been estimated to be components of beeswax (Pinzon et al., 2013).

Regarding the evidence for autofluorescence emissions by the wax films of *H. rhodocerata* and *E. lata*, no similar reference to other bee waxes has been found. The autofluorescence characteristics reported here might indicate the presence of unsaturated fatty acids with four or more  $-\text{C}=\text{C}-$  double bonds (Stockert, JC, 2016, personal communication). However, if that were the case, a band at  $1,680\text{--}1,600\text{ cm}^{-1}$  would be revealed in the FT-IR spectra (Pavia et al., 2009), which was not observed. Thus, the meaning of the autofluorescence findings could not be resolved with the present methods.

Although the integument covering films are conformed to the contour of the larvae of *H. rhodocerata* and *E. lata*, they are slightly separated from their bodies. The wrinkled and folded structure of the covering film from *H. rhodocerata* and the lacy material of the integument covering of *E. lata* that were especially demonstrated using polarization microscopy, probably result from the distribution of the secreted material over the entire body surface,



**Figure 4.** Polarized light images obtained for the cover film from *Ericrocis lata* larvae. A first-order red compensator demonstrates the negative birefringence sign typical of lipid-aligned elements (a,  $\gamma$ ), whereas polarized light/differential interference contrast optics images against a black background not only demonstrate this interference color but also show different colors that indicate the various directional distributions and thicknesses of the birefringent materials (b,c). The ordered distribution of papillae images (b,c) probably results from the imprints of growing insect setae. Scale bars equal  $20\ \mu\text{m}$ .



**Figure 5.** Fourier transform-infrared spectra of larval cover films are compared in the  $3,500\text{--}700\ \text{cm}^{-1}$  range (a) and detailed in the  $1,800\text{--}1,300\ \text{cm}^{-1}$  (b), and  $1,200\text{--}700\ \text{cm}^{-1}$  (c) spectral windows. A, absorbance; H, *Hesperapis rhodocera* (purple line); E, *Ericrocis lata* (black line).

which even imprints the sites where the larval setae arise on its surface. Only the spiracle openings remain uncovered. Particularly in the case of *H. rhodocera*, the larval wax integument covering has possibly evolved as an important protection against water depletion during a long diapause (Hadley, 1980); the larvae of this bee species must withstand arid environmental conditions like those of the territories of the Arizona, New Mexico, and NE Mexico in the absence of a cocoon (Rozen, 2016). The diapausing larvae of *H. rhodocera* are located ~30 cm below the surface in normally dry desert ground for at least 10 months, from the end of August to the following early July. The larval period may extend for as long as 5 years depending on the rains at the nesting site (J.G.R., unpublished data). It is suggested that surface waxes help protect plants and animals from a deadly rate of desiccation (Hadley, 1980). Indeed, it would not be surprising if wax coating on diapausing larvae would eventually be found characteristic of the entire subfamily Dasypodainae of the Melittidae.

Regarding *E. lata*, this bee may produce a cocoon that is not effective in protecting its larvae against desiccation (Rozen & Buchmann, 1990), for these cocoons apparently lack the sheet-like silk observed in bees like certain Megachilidae (Rozen et al., 2011). Presumably, a wax integument covering endows the diapausing larvae with a protective shield, establishing a waterproofing barrier that helps to preserve the water balance (Hadley, 1980).

Although the role of these coverings in water conservation could not be demonstrated through an assay in which larval specimens were manipulated and deprived of their integument coverings, due to technical experimental difficulties, we still believe that this is a role of these coverings.

The acquisition of a wax covering in *H. rhodocera* and *E. lata* diapausing larvae may represent an example of parallel evolution between these two distantly related bee species. An outstanding case of parallel evolution with respect to the appearance of waxes with similar chemical compositions has already been proposed for two widely divergent groups such as plants and arthropods (Hadley, 1980). Further studies involving integument coverings and cocoons in the diapausing larvae of nonsocial bee species other than those analyzed here would certainly provide additional support for this hypothesis.

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