# Exoskeletal microstructure and tensile behavior of the acorn weevil rostrum

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#### INTRODUCTION

The exoskeleton of Coleoptera (beetles) is a hierarchically-structured fibrous composite characterized or less in revisions, 12 nanofibrils embedded in a heterogeneous protein matrix. Although α-chitin is brittle and strongly anisotropic, argu-ment/nerribeetle cuticle is simultaneously rigid and tough due to its unique laminate microstructure. Beetle cuticle is divided into two structurally and mechanically distinct regions, the (outer) exocuticle and (inner) endocuticle. exocuticle is characterized by a transversely-isotropic helicoidal structure that is comparatively rigid (though brittle) and has been the subject of numerous prior studies. By contrast, the endocuticle of beetles is comprised of large, aligned bundles (macrofibers) of chitin that are strongly anisotropic. Arranged in unidirectional laminae, the macrofibers improve cuticle toughness by inhibiting crack formation and propagation between 30 successive plies [11–13].

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In general, impact-prone areas and exaggerated structures, such as horns and legs exhibit cuticle organization that resists deformation and fracture. Acorn weevils in the genus *Curculio* are typified by an elongate structure of the head, called the rostrum (snout), which instead exhibits unusual distal flexibility. The rostrum <sub>37</sub> is a hollow, strongly curved (over 90° in some species), 38 cylindrical, exoskeletal extension of the otherwise nearly-39 spherical head, which bears at its apex the terminal chew-40 ing mouthparts. Despite being composed of the same material as other rigid body parts, the snout can be repeat-42 edly bent without evident damage. This structure is used 43 by the female to feed and to excavate sites for egg-laying 44 (oviposition); the latter process causes significant, appar-45 ently elastic, deformation of the rostrum. By maintaining constant tension on the snout and rotating around the bore-hole, females are able to flex the rostrum into 49 a near-perfectly straight configuration and thereby pro-

50 duce a linear channel into the fruit; a single female may 51 prepare hundreds of such sites during the adult stage.

While this behavior has been observed in many species of Curculio, it was unclear how the rostrum of female 54 acorn weevils can withstand the repeated, often extreme 55 bending incurred during the process of egg-chamber ex-56 cavation. In this study we characterize the composite 57 profile of the rostral cuticle to account for the observed 58 flexibility of the snout. We show that the relative layer thicknesses and fiber orientation angles of the exocuticle 60 and endocuticle of the rostrum are strongly differenti-61 ated from the head capsule and other body parts, and 62 we estimate the effect of these differences using Classical 63 Laminate Plate Theory (CLPT).

Because recent studies have shown that the yield 65 strength of the weevil rostrum exoskeleton is lower in ten-66 sion than compression, we perform a comparative analy-67 sis of the ultimate tensile strength of the rostrum across 68 species and snout morphotypes; we also report the re-69 sults of displacement-controlled load cycling of the snout 70 in a species with strongly curved morphology. We relate 71 an observed increase in the volume fraction of endocuti-72 cle in the rostrum to higher tensile strength at the ros-73 tral apex in all tested species, and find that a strongly 74 curved rostrum can be flexed repeatedly without harm 75 to the structure.

We additionally describe the fracture mechanics of the 77 snout, as pertains to both cuticle composite structure 78 and tensile behavior, and consider how modification of 79 the cuticle may reduce the risk of rostral fracture dur-80 ing oviposition. Based on our findings, we posit that the composite profile of the rostral apex enables the snout to be flexed until straight while remaining within the elas-83 tic limits of the material, mitigating the risk of structural 84 damage, and without evident alteration of the mechani-85 cal properties of the individual components of the cuti-86 cle across the structure and between species. Thus, the 87 flexibility and tensile strength of the rostrum appear to 88 be derived exclusively from modification of the composite 89 architecture of the exoskeleton. To our knowledge, this is 90 the first time that a modified composite profile has been 91 reported as a means of enhancing structural elasticity in

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# MICROSTRUCTURE OF THE CURCULIO ROSTRUM

In arthropods (including beetles), the exocuticle is comprised of numerous unidirectional laminae of chitin fibers; each layer is the thickness of a single fiber (2-99 4 nm) embedded in a proteinaceous matrix. These layers <sup>100</sup> are stacked at a more or less constant angle to each other, thus forming a quasi-isotropic laminate referred to as the Bouligand structure [7–9]. This layout effectively pro-103 duces a transversely isotropic composite, mitigating the 104 strong anisotropy of  $\alpha$ -chitin to yield a versatile building material for the exoskeleton.

Beetle endocuticle, however, is unique among arthropods and is comprised of large (1-5 µm diameter) uni-108 directional bundles of chitin, called macrofibers. Chitin macrofibers are orthotropic (axial:  $E_1 = 8.5 \,\mathrm{GPa}$ , transverse:  $E_2 = E_3 = 0.52 \,\mathrm{GPa} \,[1]$ ) and arranged in INSERIE unidirectional plies, seen in Figs. [12, 13]. Typically, ad-HERE 13 jacent macrofiber plies are paired and pseudo-orthogonal INSER (i.e., angled approx. 90° to each other, see Figs.), with REFS-116 a constant stacking angle between pairs, although other configurations have been observed [11, 12]. This geometric sequence of the macrofiber laminae yields an aplami-nate is 120 proximately transversely isotropic composite, similar to the Bouligand structure. Notably, the resulting lami-122 nate is less rigid than the exocuticle, but exhibits greater 123 toughness because the pseudo-orthogonal plies effectively 124 inhibit crack formation and propagation between succes-125 sive layers [11–13].

Serial thin sectioning and scanning electron microscopy of fractured Curculio specimens has revealed that endo-128 cuticle in the head capsule fits this general profile, with 186 <sup>129</sup> an angle of approximately 30° between successive pairs of pseudo-orthogonal plies. Additionally, in the head cap- 187 INSER Fig.) in two key characteristics:

> endocuticle thickened to offset this reduction to 195 ture. maintain a constant cuticle thickness in the head.

2. The endocuticular macrofibers exhibit no rotation 196 between successive pseudo-orthogonal plies, which 197 are all oriented at approximately  $\pm 45^{\circ}$  to the longitudinal axis of the snout (i.e., an antisymmetric 198  $[\pm 45^{\circ}]$  angle-ply laminate).

150 species, C. longinasus Chittenden, 1927 [1?]. This com- 203 de-ionized water for 24 hours Each specimen was sub-151 posite profile has now been uncovered in the rostral apex 204 jected to force-controlled, uniaxial loading to fracture at

152 of six additional, phylogenetically disparate, species (de-153 tailed in methods), indicating that this is likely a genuswide trait. In all examined species, the portion of the snout between the head capsule and apex of the scrobe exhibits a gradual transition in composite profile along 157 an anterior-posterior gradient.

To investigate the effect of these cuticle modifications, 159 we estimated uni-axial membrane and transverse flex- $_{160}$  ural Young's moduli of the cuticle in both the rostral 161 apex and head capsule using Classical Laminate Plate 162 Theory (CLPT), as detailed in our methods. We previ-163 ously derived the effective elastic constants of the cu-164 ticle regions of C. longinasus, which we used here to 165 construct constitutive equations for the entire cuticle of 166 that species. The cuticle was estimated to have membrane and flexural moduli of  $E_m = 4.77 \,\mathrm{GPa}$  and  $_{168}$   $E_f = 6.04$  GPa, respectively, in the head capsule; how-169 ever, in the rostral apex we found that these values were 170 reduced by approximately 72% and 60%, respectively  $(E_m = 1.36 \,\mathrm{GPa}, E_f = 2.44 \,\mathrm{GPa}).$ 

To assess the individual contributions of layer thickness 173 and stacking angle sequence to cuticle flexibility in the 174 rostral apex, two hybrid cuticle lay-ups were also mod-175 eled, each with only one of the modifications, namely:

- 1. A hybrid cuticle with the angle stacking sequence of regular cuticle (the head capsule) but possessing the layer thicknesses of the rostral apex.  $E_m = 3.73 \,\mathrm{GPa}, E_f = 4.31 \,\mathrm{GPa}, \mathrm{representing}$ 22% and 29% decreases, respectively.
- 2. A hybrid cuticle with the layer thicknesses regular cuticle but possessing the anstacking sequence of the rostral apex angle-ply in the endocuticle).  $= 3.77 \,\mathrm{GPa}, E_f = 5.76 \,\mathrm{GPa}, \mathrm{representing}$ 21% and 4.7% decreases, respectively.

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Each of the cuticle modifications noted in the rostral 131 sule, the thickness of the exocuticle in cross section is 188 apex individually decreased the elastic moduli of the cutithickness nearly equal to that of the endocuticle. However, we 189 cle; however, they appear to have a synergistic combined have also found that the cuticle composite lay-up of the 190 effect on cuticle elasticity, rather than a simple additive here 135 rostral apex differs from that of the head capsule (see 191 effect. This result suggests that both modifications are 192 necessary in order for the snout to function properly in 193 the living animal, where the combined effect allows the The exocuticle is reduced to a thin shell, with the 194 rostrum to bend until completely straight without frac-

# FORCE-CONTROLLED LOADING TO FRACTURE

To better characterize the mechanical behavior of the 199 rostrum at failure, we performed tensile testing on the 200 snouts of six Curculio species, representing a mixture of In previous work we identified these modifications to 201 closely and distantly related taxa. Heads of female specthe composite structure of the cuticle within a single 202 imens of each species were rehydrated by immersion in 205 a constant stress rate (detailed in Methods). We then ex- 263 composite profile of the cuticle in the rostral apex makes and mechanical behavior of the snout, using phylogenetic 265 sistant. linear mixed effects models to account for phylogenetic non-independence in the residual variances of the models.

In general, the specimens exhibited a non-linear vis- 266 III. LOAD CYCLING OF CURCULIO CARYAE 211 coelastic response curve characterized by a sharp increase in stress at higher strains, terminating in apparently brittle fracture after a period of strain hardening. We observed that the maximum force sustained at the site of the break was strongly correlated with the cross-sectional area of the endocuticle, and not the exocuticle, at that site. Consequently, there was a negative correlation between ultimate tensile strength of the specimen and the ratio of exocuticle to endocuticle cross-sectional area at the site of fracture. Inversely, the failure strength of the cuticle is positively associated with the relative proportion of the endocuticle.

Although CLPT predicts that the proportion of exocuticle is positively correlated with the stiffness of generalhavior of the entire rostrum. Because the cross-sectional areas of the cuticle regions vary across the length of the  $_{283}$  amplitude appeared to reach an asymptotic minimum. head along an anterior-posterior gradient, it is not pos-230 sible to correlate measurements from the fracture sur-231 face to the properties of the entire snout. Instead, we found that the uniaxial elastic modulus of the snout unultimate strain and toughness of the snout (as was the to be statistically significant and independent of species 237 membership.

Additionally, we observed a moderate stiffening size- 293 because of the greater volume fraction of exocuticle.

in sclerotization or chitin composition within the cuticle, 312 dissipating the accumulated strain. rather than composite profile, are responsible for the me- 313 <sub>261</sub> cle, likely because the endocuticle is organized in large <sub>317</sub> species ( $F_m ax = 5.02 \,\mathrm{N}$ ). Given this surprising result, 262 bundles of aligned, anisotropic fibers. Thus, the altered 318 it appears that the specimen was undamaged by the test-

amined the correspondence between composite structure 264 the rostrum simultaneously more flexible and fracture re-

To confirm that repeated, proscribed straightening of 268 the rostrum does not result in damage to the cuticle, we 269 performed displacement-controlled fatigue testing on a 270 typical female specimen of *Curculio caryae*, a species that exhibits extreme  $(80 - 90^{\circ})$  rostral curvature. The head 272 was removed and rehydrated for 24 hours in de-ionized  $_{273}$  water, then coated in grease to prevent loss of moisture 274 and stiffening of the specimen during the duration of the 275 test. The specimen was then aligned so that uniaxial 276 tension would induce elongation of the apical portion of 277 the rostrum. The strain per cycle was fixed at an am-278 plitude sufficient to completely elongate the snout and generate a tensile load equal to  $0.2\times$  the average tensile ized cuticle, we found no associations between the cross- 280 strength of conspecifics in the straightened configuration, sectional properties of the fracture site and the gross be- 281 at a frequency of 0.33 Hz. The test was terminated after 282 a period of two weeks (ca. 400K cycles) when the strain

We observed viscoelastic behavior in the rostrum, as 285 indicated by histeresis in the stress-strain relationship 286 during each cycle. Strain amplitude decreased logarith-287 mically with cycle number, and the specimen initially der low strain (E-low) was inversely correlated with the 288 appeared to have deformed plastically during the test. 289 We believed that this indicated damage to the specimen; secant modulus, E-sec) These associations were all found 290 however, after cleaning the specimen in a 24 hour wash 291 with ethanol and water, we observed that rostrum re-292 turned to its original shape.

We cannot fully account for the stress relaxation of effect with respect to rostral length. This observation 294 the rostrum after testing, but we speculate that it arose was initially quite puzzling, as we had expected that a 295 from the same mechanism as the viscoelastic behavior longer (and typically more strongly curved [insert eco- 296 observed in tensile testing. The endocuticle is made of morph paper ref here]) snout would need to be more 297 aligned α-chitin nanofibrils whose crystalline structure is flexible to avoid fracture during oviposition. Based on 298 enforced by hydrogen bonds between individual chitin preliminary confocal microscopy data, we speculate that 299 chains along their length. The viscoelasticity of the cuthis may be the result of a longer transition gradient from 300 ticle is thought, in part, to come from slippage between the basal profile to the apical profile in longer rostra.  $_{301}$  these chains as the hydrogen bonds break and reform in We hypothesize that a decrease in exocuticle thickness 302 response to shearing between the chitin molecules. We along a longer portion of the base might reinforce the 303 believe that repeated strain may have caused such slipsnout against buckling; however Young's modulus of the  $_{304}$  page in the endocuticle of the rostral apex during the rostrum would be comparatively higher in these species 305 fatigue test; however, without sufficient time for the ma-306 terial to completely relax after deformation, the specimen From these results, we infer that the gross elastic 307 would slowly accumulate strain and consequently deform behavior of the cuticle is consistent across the genus, 300 viscoelastically (REF). After 24 hours soaking in ethanol in agreement with our current understanding of cuticle 310 and water, the hydrogen bonds would relax sufficiently to mechanobiology. It is therefore unlikely that differences 311 allow the specimen to return to its original configuration,

The specimen did not show any evidence of fractures, chanical behavior of the rostrum in different species. Fur- 314 micro-tears, or shear cusps anywhere in the surface of thermore, the endocuticle apparently contributes more 315 the exocuticle, and, furthermore, the tensile strength of to the tensile strength of the rostrum than the exocuti- 316 the specimen was consistent with other members of its

history

319 ing. We therefore expect that under normal conditions 374 areas of endocuticle, this was often obscured by delami-320 in life, repeated bending of the snout does not exceed 375 nated endocuticular macrofibers.  $_{\rm 321}$  the yield strength of the cuticle, and the bending strain  $_{\rm 376}$ 322 is purely elastic.

#### FRACTOGRAPHY OF TEST SPECIMENS

Due to the complex failure patterns present across the fracture surface, it was not always possible to identify void nucleation and crack initiation sites (both primary and secondary). Nonetheless, we observed several patterns characteristic of both the micro-scale behavior of the cuticle and the meso-scale behavior of the rostrum during uniaxial tensile failure, which we describe here.

Scanning electron and light microscopy of testing spec-332 imens revealed that the fracture mechanics of the exocuticle and endocuticle differed according to their microstructure. In transverse view, the exocuticle consistently presented a nearly continuous fracture surface, indicative of comparatively brittle failure, likely due to the helicoidal arrangement of  $\alpha$ -chitin laminae and resultant transverse isotropy of the Bouligand structure. The exocuticle typically appeared to fracture at lower strains 397 macrofibers are less brittle than the exocuticle due to 404 tire thickness of the laminate. Although other, more cuticular shear-cusp formation in uniaxial tension sug- 408 ates tension along the ventral surface of the rostrum. 351 gests extension-shear coupling within individual endocuticle laminae, which caused ply-splitting via mode II fracture between macrofibers at high strain. This intralaminar extension-shear coupling also yielded off-axis in- 409 plane resultant forces as a function of lamina orientation angle; mode III shearing then occurred between laminae with opposing in-plane resultant forces, causing inter-ply delamination. Tensile failure of the macrofiber laminae would ultimately occur via mixed-mode I/II (transverse tension/intra-laminar shear) fracture due to an increase in applied stress caused by ply-splitting in adjacent laminae. 362

At the meso-scale, most specimens fractured along a 417 364 single plane across and between the occipital sulci, which 418 endocuticle deforms viscoelastically along the longitudi-

The first layer of endocuticle usually fractured along 377 the same plane as the exocuticle, and ventrally the en-378 docuticle laminae typically converged toward a scarp-like 379 region characterized by severe delamination and numer-380 ous de-bonded macrofibers. Additionally, macrofibers 381 oriented in the direction of crack propagation exhibited extensive ply-splitting with intermittent transverse shearing, while macrofibers oriented against the direction of crack propagation primarily displayed fracture by 385 transverse shear along the plane of ply-splitting in adja-386 cent laminae. Thus, contralateral fibers in the same lam-387 ina displayed opposing fracture modes. In addition, the 388 ventrolateral surfaces often exhibited extensive inter-ply 389 delamination and fiber de-bonding in scarp-like promi-390 nences, likely due to a combination of tensile failure and 391 shearing along the dorsally-radiating crack front. Dor-392 sally, the coalescent crack fronts often caused significant de-bonding and ply-splitting followed by broom-like ten-394 sile failure; in some specimens, the contralateral crack 395 fronts were out of plane and coalesced via transverse 396 shearing through a large dorsal section of cuticle.

From these patterns we hypothesize that the than the endocuticle, with shear-cusp formation evident 398 exocuticle-rich occipital sulci are the most likely site for across the fracture surface and over unbroken endocuti- 399 the initiation of void nucleation and catastrophic failure cle on the rostral surface adjacent to the plane of frac- 400 of the integrated rostral cuticle in cross section, as illusture. Conversely, the endocuticle exhibited severe de- 401 trated in Fig REF. According to this model, structural lamination, off-axis ply-splitting, and fiber-pulling away 402 failure would take place as cracks propagate through the from the fracture surface, indicating that endocuticular  $_{403}$  endocuticle from these sutures, which penetrate the enthe unidirectional alignment of their  $\alpha$ -chitin fibers. Be-  $_{405}$  complex failure modes have been observed, we posit that cause the exocuticle of weevils is anchored to the en- 406 in live specimens this is the most likely mechanism of docuticle by cross-linking fibers, the presence of exo- 407 tensile failure because typical bending behavior gener-

# CONCLUSIONS

In this study, we... raising the intriguing possibility 411 that this system evolved via negative selection of breakage, rather than positive selection of flexibility.

evolutionary constraint (like occipital sulci)

The macrofibers are pulled longitudinal axis, explains 416 fmax, toughness, and upcurved stress-strain curve.

We therefore infer that, under uniaxial tension, the are cuticle invaginations that traverse the entire length of 419 nal axes of the macrofibers, causing ply-splitting and dethe rostrum. These sulci increase the volume fraction of 420 lamination at high strain, while the overlying exocuticle exocuticle in the ventral part of the snout and create large 421 exhibits brittle fracture, at a comparatively low strain, interfaces ideal for void nucleation. The exocuticle of the 422 due to shearing between the stretching endocuticle fibers occipital sulci usually displayed shear cusps oriented out- 423 to which it is anchored. This outcome is consistent ward from the center of the invagination, continuing dor- 424 with behavior shown in previous studies and is congrusolaterally and ventromedially. Ventrally the cusps con- 425 ent with theoretical consideration of cuticle microstrucverged toward a prominent scarp where the crack fronts 426 ture in CLPT, which predicts extension shear-coupling  $_{373}$  joined, although in specimens with large cross-sectional  $_{427}$   $(A_{16}, A_{26} \neq 0)$  for individual off-axis macrofiber laminae.

(maybe this in conclu-sions)

#### VI. METHODS

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Methods, including statements of data availability and any associated accession codes and references, are available in the online version of this paper.

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### **AUTHOR CONTRIBUTIONS**

477 **Andrew Jansen:** Conducted sectioning and staining, 478 microscopy and imaging, tensile and fatigue testing, statistical analysis, and participated in manuscript preparation.

Jason Williams: Conducted tensile and fatigue testing, participated in manuscript preparation.

Nikhilesh Chawla: Facilitated microscopy, tensile and fatigue testing, and participated in manuscript preparation.

Nico Franz: Facilitated specimen acquisition and imaging, participated in manuscript preparation.

#### ADDITIONAL INFORMATION

Supplementary information is available in the online version of the paper. Reprints and permissions information is available online at www.nature.com/reprints.
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# COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

#### **METHODS**

# Histological sectioning

# Tensile and fatigue testing

# Specimen imaging and microscopy

#### Cold-fracture of rostra

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# Constitutive modeling of the cuticle

General Approach

Cuticle profile of model The cuticle of C. longinasus 503 is 50 microns thick; we use C. longinasus because we have constitutive models for the cuticle regions of this species. We assumed equal layer thicknesses in the endocuticle of the basal cuticle, and equal thicknesses of exocuticle and endocuticle. In the apical cuticle we assumed the exocuticle and upper 8 layers of the endocuticle were each 5 microns thick, with 4 thinner layers of endocuticle, each 1.25 µm thick. The angles of the endocuticle layers for both types of cuticle are as described above.

Permutations We additionally calculated models for two hypothetical hybrid cuticles: one model has the layer thicknesses of the apex, but fiber orientations of the base, while the second has the fiber orientations of the apex, but the layer thicknesses of the base.

Classical Laminate Plate Theory We begin by calcu-519 lating the 2D reduced stiffness matrix for each part of the 520 cuticle. For orthotropic materials with the principal axes 521 parallel to the ply edges, the reduced stiffness matrix is 522 defined as follows:

$$[Q] = \begin{bmatrix} Q_{11} & Q_{12} & 0 \\ Q_{21} & Q_{22} & 0 \\ 0 & 0 & Q_{66} \end{bmatrix} , \tag{1}$$

and where: 523

$$\begin{split} Q_{11} &= \frac{E_1}{1 - \nu_{12}\nu_{21}} \,, \\ Q_{12} &= \frac{E_1\nu_{21}}{1 - \nu_{12}\nu_{21}} = Q_{21} \,, \\ Q_{21} &= \frac{E_2\nu_{12}}{1 - \nu_{12}\nu_{21}} = Q_{12} \,, \\ Q_{22} &= \frac{E_2}{1 - \nu_{12}\nu_{21}} \,, \\ Q_{66} &= G_{12} \,. \end{split} \tag{2}$$

527 formed stiffness matrix according to:

$$[\bar{Q}] = [T]^{-1}[Q][T]^{-T},$$
 (3)

where the transformation matrix [T] is defined as:

$$[T] = \begin{bmatrix} \cos^2 \theta & \sin^2 \theta & 2\cos\theta\sin\theta\\ \sin^2 \theta & \cos^2 \theta & -2\cos\theta\sin\theta\\ -\cos\theta\sin\theta & \cos\theta\sin\theta & \cos^2 \theta - \sin^2 \theta \end{bmatrix}. \quad (4)$$

Using the lay-ups specified for the cuticle permutations, we calculate the extensional stiffness matrix [A], bending stiffness matrix [D], and bending-extension cou- $_{532}$  pling matrix B for each laminate consisting of n layers at a distance z from the laminate mid-plane. The elements of these matrices can be found according to:

$$A_{ij} = \sum_{k=1}^{n} (\bar{Q}_{ij})_k (z_k - z_{k-1}),$$

$$B_{ij} = \frac{1}{2} \sum_{k=1}^{n} (\bar{Q}_{ij})_k (z_k^2 - z_{k-1}^2),$$

$$D_{ij} = \frac{1}{3} \sum_{k=1}^{n} (\bar{Q}_{ij})_k (z_k^3 - z_{k-1}^3).$$
(5)

These stiffness matrices relate vectors of resultant forces  $\{N\}$  and bending moments  $\{M\}$  to mid-surface strains and curvatures  $\{\epsilon^{\circ}\}$  and  $\{\kappa\}$ , respectively, in the 538 laminate according to the following relationship:

$${\{N\} \atop \{M\}\}} = \begin{bmatrix} [A] & [B] \\ [B] & [D] \end{bmatrix} {\{\epsilon^{\circ}\} \atop \{\kappa\}}.$$
 (6)

For symmetric laminates, [B] = 0, and therefore:

$$\{N\} = [A]\{\epsilon^{\circ}\},$$
  
$$\{M\} = [D]\{\kappa\},$$
  
(7)

or, in expanded form:

$$\begin{cases}
N_{xx} \\
N_{yy} \\
N_{xy}
\end{cases} = \begin{bmatrix}
A_{11} & A_{12} & A_{16} \\
A_{21} & A_{22} & A_{26} \\
A_{61} & A_{62} & A_{66}
\end{bmatrix} \begin{Bmatrix}
\epsilon_{xx}^{\circ} \\
\epsilon_{yy}^{\circ} \\
\gamma_{xy}^{\circ}
\end{Bmatrix},$$

$$\begin{cases}
M_{xx} \\
M_{yy} \\
M_{xy}
\end{Bmatrix} = \begin{bmatrix}
D_{11} & D_{12} & D_{16} \\
D_{21} & D_{22} & D_{26} \\
D_{61} & D_{62} & D_{66}
\end{bmatrix} \begin{Bmatrix}
\kappa_{xx} \\
\kappa_{yy} \\
\kappa_{xy}
\end{Bmatrix}.$$
(8)

If we make the simplifying assumptions (see ref.) that 543 (1) the laminate experiences pure axial loading and transverse bending (i.e.,  $N_{yy}=N_{xy}=0$  and  $M_{yy}=$ For each layer k, the reduced stiffness matrix is trans-  $M_{xy} = 0$ , respectively) and (2) the laminate is a beam  $_{525}$  formed to account for the layer orientation angle  $\theta$  within  $_{546}$  of sufficiently high aspect ratio to minimize the Poisson 526 the laminate coordinate system, yielding a reduced trans- 547 effect and anisotropic shear coupling (i.e., below we ef-548 fectively let  $A_{12}^* = A_{16}^* = 0$  and  $D_{12}^* = D_{16}^* = 0$ , then

549 we can calculate the in-plane effective flexural and axial 571 Young's moduli of the laminate along the x-axis.

For axial Young's modulus of the laminate, we first define the average membrane stresses in the laminate as:

$$\{\bar{\sigma}^m\} = \frac{\{N\}}{z_1 - z_n}.$$
 (9)

By substitution in Eq. 7, we find: 553

$$\begin{cases} \bar{\sigma}_{xx}^{m} \\ \bar{\sigma}_{yy}^{m} \\ \bar{\tau}_{xy}^{m} \end{cases} = \frac{1}{(z_{1} - z_{n})} \begin{bmatrix} A_{11} & A_{12} & A_{16} \\ A_{21} & A_{22} & A_{26} \\ A_{61} & A_{62} & A_{66} \end{bmatrix} \begin{cases} \epsilon_{xx}^{m} \\ \epsilon_{yy}^{m} \\ \gamma_{xy}^{m} \end{cases}, \quad (10) \ _{579}^{578}$$

and, by inverting this equation (let  $A^* = A^{-1}$ ) and 555 substituting based on the assumptions above, we infer:

$$\epsilon_{xx}^m = (z_1 - z_n) A_{11}^* \bar{\sigma}_{xx}^m \,.$$
 (11) 586

We therefore define Young's modulus for effective axial 557 elasticity as:

$$E_{xx}^{m} = \frac{\bar{\sigma}_{xx}^{m}}{\epsilon_{xx}^{m}} = \frac{1}{(z_{1} - z_{k})A_{11}^{*}}$$
(12)

To find the transverse flexural Young's modulus of the 559 laminate, we first specify the moment-curvature relation  $_{560}$  of an Euler-Bernoulli beam:

$$M = EI\kappa. (13)$$

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Along the x-axis, the second moment of area for a rect-562 angular cross-section is:

$$I_{yy} = \frac{b(z_1 - z_n)^3}{12} \,, \tag{14}$$

564 ment along the x-axis is related to the moment of the 607 pared to models with only cross-sectional area of either 565 beam by:

$$M = M_{xx}b. (15)$$

Thus, given the assumption that  $D_{12}=D_{16}=0$ , 613 567 Young's modulus for the effective transverse flexural elas- 614 568 ticity of the laminate can be found by making Eq. 13 spe-569 cific to transverse flexure of the x-axis and rearranging 570 the terms:

$$E_{xx}^f = \frac{12M_{xx}}{(z_1 - z_n)^3 \kappa_{xx}} \tag{16}$$

From inversion of Eq. 7 (let  $D^* = D^{-1}$ ) this reduces

$$E_{xx}^{f} = \frac{12}{(z_1 - z_n)^3 D_{11}^*}$$
Statistical analysis (17)

General approach To explore the relationships be-575 tween the composite structure and mechanical proper-576 ties of the cuticle, we fit phylogenetic linear mixed-effects 577 models to the data using maximum likelihood estimation.  $_{\rm 578}$  In order to control for phylogenetic non-independence in the data, we included the species of each specimen as a random effect in all models. We also allowed for corre-581 lation in the error term of the models, as specified by 582 a variance-covariance matrix generated from a Brownian motion model of trait evolution along the phylogeny. Response variables and covariates were natural-log transformed, as needed, to ensure model residuals were normally distributed and homoscedastic. In all models, we 587 tested whether the inclusion of phylogenetic correlation in the model error produced significantly better model 589 fit, using a likelihood-ratio test and  $R_{\sigma}^2$ -difference test 590 between the fully-specified model and a model lacking 591 the phylogenetic effect.

Hypothesis testing The following three hypotheses 593 were tested using PGLMMs fitted using ML estimation:

- 1. The maximum sustained tensile force is proportional to the cross-sectional area of the endocuticle, and not that of the exocuticle.
- 2. The ultimate tensile strength of the samples is inversely proportional to the ratio of exocuticle to endocuticle at the location of fracture.
- 3. Young's modulus of the samples is proportional to the length of the snout.

We fitted a fully-specified model with the cross-603 sectional area of endocuticle and exocuticle at the site 604 of fracture as fixed effects, including an interaction term, 605 and with maximum tensile force sustained prior to frac-Given the assumption that  $M_{yy} = M_{xy} = 0$ , the mo- of ture as a response variable. This model was then com-608 endocuticle or exocuticle as the sole fixed effect in the 609 model. We then tested the first hypothesis by using 610 likelihood-ratio tests and  $R_{\beta*}^2$ -difference tests between (15) 611 each of the three models.

> The hypothesis that Model selection and fitting Estimating phylogenetic signal

> > Code availability

Data availability

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# REFERENCES

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