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## INTRODUCTION

The exoskeleton of Coleoptera (beetles) is a hierarchically-structured fibrous composite characterized or less in by variously arranged  $\alpha$ -chitin (N-acetylglucosamine) sions, in 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 struc-32 tures, such as horns and legs exhibit cuticle organization that resists deformation and fracture. Acorn weevils in the genus Curculio Linnaeus, 1758<sup>1</sup> are typified by an elongate structure of the head, called the rostrum (snout), which instead exhibits unusual distal flexibility. The rostrum is a hollow, strongly curved (over 90° in 38 some species), cylindrical, exoskeletal extension of the 39 otherwise nearly-spherical head, which bears at its apex 40 the terminal chewing mouthparts. Despite being com-41 posed of the same material as other rigid body parts, the 42 snout can be repeatedly bent without evident damage.

While this behavior has been observed in many species 54 of Curculio, it was unclear how the rostrum of female 55 acorn weevils can withstand the repeated, often extreme 56 bending incurred during the process of egg-chamber ex-57 cavation. In this study we characterize the composite 58 profile of the rostral cuticle to account for the observed 59 flexibility of the snout. We show that the relative layer 60 thicknesses and fiber orientation angles of the exocuticle 61 and endocuticle of the rostrum are strongly differentiated 62 from the head capsule and other body parts, and we esti-63 mate the effect of these differences on the elasticity of the 64 cuticle using Classical Laminate Plate Theory (CLPT).

Because recent studies have shown that the yield 66 strength of the beetle exoskeleton is lower in tension than 67 compression, we perform a comparative analysis of the 68 ultimate tensile strength of the rostrum across species 69 and snout morphotypes; we also report the results of 70 displacement-controlled load cycling of the snout in a 71 species with strongly curved morphology. We relate an  $_{72}$  observed increase in the volume fraction of endocuticle in 73 the rostrum to higher tensile strength at the rostral apex in all tested species, and find that a strongly curved ros-75 trum can be flexed repeatedly without harm to the struc-76 ture.

We additionally describe the fracture mechanics of the 78 snout, as pertains to both cuticle composite structure 79 and tensile behavior, and consider how modification of 80 the cuticle may reduce the risk of rostral fracture dur-81 ing oviposition. Based on our findings, we posit that the 82 composite profile of the rostral apex enables the snout to 83 be flexed until straight while remaining within the elas- $_{\it 84}$  tic limits of the material, mitigating the risk of structural 85 damage, and without evident alteration of the mechani-

trated

<sup>43</sup> This structure is used by the female to feed and to exca-44 vate sites for egg-laying (oviposition); the latter process 45 causes significant, apparently elastic, deformation of the rostrum. By maintaining constant tension on the snout 48 and rotating around the bore-hole, females are able to 49 flex the rostrum into a near-perfectly straight configura-50 tion and thereby produce a linear channel into the fruit; 51 a single female may prepare hundreds of such sites during 52 the adult stage.

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Pursuant to the International Code of Zoological Nomenclature, the first mention of any specific epithet will include the full genus and species names as a binomen (two part name) followed by the author and date of publication of the name. This is not an in-line reference; it is a part of the name itself and refers to a particular species-concept as indicated in the description of the species by that author.

the insect exoskeleton.

# MICROSTRUCTURE OF THE CURCULIO ROSTRUM

In arthropods (including beetles), the exocuticle is 97 comprised of numerous unidirectional laminae of chitin 98 nanofibrils; each layer is the thickness of a single fiber (2-4 nm) embedded in a proteinaceous matrix. These layers 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 produces a transversely isotropic composite, mitigating the strong anisotropy of  $\alpha$ -chitin to yield a versatile building material for the exoskeleton.

Beetle endocuticle, however, is unique among arthro-107 pods and is comprised of large (1-5 µm diameter) uni-108 directional bundles of chitin, called macrofibers. Chitin 109 macrofibers are orthotropic (axial:  $E_1 = 8.5\,\mathrm{GPa}$ , transverse:  $E_2 = E_3 = 0.52 \,\mathrm{GPa} \,[1]$ ) and arranged in unidirectional plies, seen in Figs. [12, 13]. Typically, ad-HERE 13 jacent macrofiber plies are paired and pseudo-orthogonal INSERT (i.e., angled approx. 90° to each other, see Figs.), with REFS-115 a constant stacking angle between pairs, although other configurations have been observed [11, 12]. This geowhat kind of 19 metric sequence of the macrofiber laminae yields an approximately 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 successive layers [11–13].

Serial thin sectioning and scanning electron microscopy of fractured Curculio specimens have revealed that endo-128 cuticle in the head capsule fits this general profile, with <sup>129</sup> an angle of approximately 30° between successive pairs of 130 pseudo-orthogonal plies. Additionally, in the head cap-131 sule, the thickness of the exocuticle in cross section is thickness nearly equal to that of the endocuticle. However, we have also found that the cuticle composite lay-up of the 135 rostral apex differs from that of the head capsule (see INSER Fig.). Distally the exocuticle is reduced to a thin shell, 195 with the endocuticle thickened to offset this reduction and maintain a constant cuticle thickness in the head.  $_{
m here}^{
m measure}$  Moreover, the endocuticular macrofibers exhibit no ro-  $_{197}$ 146 laminate).

86 cal properties of the individual components of the cuti- 148 the composite structure of the cuticle within a single cle across the structure and between species. Thus, the 149 species, C. longinasus Chittenden, 1927 [1?]. This comflexibility and tensile strength of the rostrum appear to 150 posite profile has now been uncovered in the rostral apex be derived exclusively from modification of the composite 151 of six additional, phylogenetically disparate, species (dearchitecture of the exoskeleton. To our knowledge, this is 152 tailed in methods), indicating that this is likely a genusthe first time that a modified composite profile has been 153 wide trait. In all examined species, the portion of the 92 reported as a means of enhancing structural elasticity in 154 snout between the head capsule and apex of the scrobe 155 exhibits a gradual transition in composite profile along 156 an anterior-posterior gradient.

> To investigate the effect of these cuticle modifications, we estimated uni-axial membrane and transverse flex-159 ural Young's moduli of the cuticle in both the rostral 160 apex and head capsule using Classical Laminate Plate 161 Theory (CLPT), as detailed in our methods. We previ-162 ously derived the effective elastic constants of the cu-163 ticle regions of C. longinasus, which we used here to 164 construct constitutive equations for the entire cuticle of 165 that species. The cuticle was estimated to have mem- $_{166}$  brane and flexural moduli of  $E_m \ = \ 4.77\,\mathrm{GPa}$  and  $_{167}$   $E_f = 6.04$  GPa, respectively, in the head capsule; how-168 ever, in the rostral apex we found that these values were 169 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 172 and stacking angle sequence to cuticle flexibility in the 173 rostral apex, two hybrid cuticle lay-ups were also mod-174 eled, each with only one of the modifications. A hy-175 brid configuration with the angle stacking sequence of 176 typical cuticle (i.e., in the head capsule) but possess-177 ing the layer thicknesses of the rostral apex had effec-178 tive moduli  $E_m = 3.73 \,\mathrm{GPa}, E_f = 4.31 \,\mathrm{GPa}, \mathrm{repressure}$ 179 senting 22% and 29% decreases from unmodified cuticle, 180 respectively. Similarly, a hypothetical cuticle with the 181 layer thicknesses of ordinary cuticle but possessing the angle stacking sequence of the rostral apex (i.e.,  $\pm 45^{\circ}$ angle-ply in the endocuticle) exhibited elastic moduli  $E_m = 3.77 \,\mathrm{GPa}, E_f = 5.76 \,\mathrm{GPa}, \mathrm{representing} \,21\% \,\mathrm{and}$ <sup>185</sup> 4.7% decreases from unmodified cuticle, respectively.

> Each of the cuticle modifications noted in the rostral 187 apex individually decreased the elastic moduli of the cuti-188 cle; however, they appear to have a synergistic combined 189 effect on cuticle elasticity, rather than a simple additive 190 effect. This result suggests that both modifications are 191 necessary in order for the snout to function properly in 192 the living animal, where the combined effect allows the 193 rostrum to bend until completely straight without frac-

## FORCE-CONTROLLED LOADING TO **FRACTURE**

To better characterize the mechanical behavior of the tation between successive pseudo-orthogonal plies, which 198 rostrum at failure, we performed tensile testing on the  $\pm 4$  are all oriented at approximately  $\pm 45^{\circ}$  to the longitudinal personants of six Curculio species, representing a mixture axis of the snout (i.e., an antisymmetric [±45°] angle-ply 200 of closely and distantly related taxa. Heads of female 201 specimens of each species were rehydrated by immersion In previous work we identified these modifications to 202 in de-ionized water for 24 hours to simulate the con203 dition of the living tissue (but see Klocke and Schmitz 261 are responsible for the mechanical behavior of the rosindependence in the residual variances of the models.

coelastic response curve characterized by a sharp increase 270 trum simultaneously more flexible and fracture resistant. in stress at higher strains, terminating in apparently brittle fracture after a period of strain hardening. We believe that strain hardening prior to fracture is a consequence of 271 III. stiff longitudinal axis of the macrofibers becoming more closely aligned to the longitudinal axis of the rostrum, thereby resisting tension more directly at higher strains. We observed that the maximum force sustained at the site of failure was strongly correlated with the crosssectional area of the endocuticle, and not the exocuticle, at that site. Consequently, there was a positive correlation between ultimate tensile strength of the specimen and the ratio of endocuticle to exocuticle cross-sectional area at the site of fracture.

Although CLPT predicts a positive association between the proportion of exocuticle and stiffness of a generalized cuticle, we found no correspondence between the cross-sectional properties of the fracture site and the gross behavior of the entire rostrum. Because the 285 peared to reach an asymptotic minimum. cross-sectional areas of the cuticle regions vary across the length of the head along an anterior-posterior grasnout under low strain (E-low) was inversely correlated with the ultimate strain and toughness of the snout (as was the secant modulus, E-sec) These associations were all found to be statistically significant and independent of species membership.

242 effect with respect to rostral length. This observation was 297 from the same mechanism as the viscoelastic behavior initially quite puzzling, as we had expected that a longer 298 observed in tensile testing. The endocuticle is made of (and typically more strongly curved [insert ecomorph pa-  $_{299}$  aligned  $\alpha$ -chitin nanofibrils whose crystalline structure is per ref here]) snout would need to be more flexible to 300 enforced by hydrogen bonds between individual chitin avoid fracture during oviposition. Based on preliminary 301 chains along their length. The viscoelasticity of the cuconfocal microscopy data (not presented herein), we spec- 302 ticle is thought, in part, to come from slippage between ulate that this may be the result of a longer transition  $_{303}$  these chains as the hydrogen bonds break and reform in gradient from the basal profile to the apical profile in 304 response to shearing between the chitin molecules. We longer rostra. We hypothesize that a decrease in exocu- 305 believe that repeated strain may have caused such slipticle thickness along a longer portion of the base might 306 page in the endocuticle of the rostral apex during the reinforce the snout against buckling; however Young's 307 fatigue test; however, without sufficient time for the mamodulus of the rostrum would be comparatively higher 308 terial to completely relax after deformation, the specimen in these species because of the greater volume fraction of 309 would slowly accumulate strain and consequently deform exocuticle.

havior of the cuticle is consistent across the genus, in 313 allow the specimen to return to its original configuration, agreement with our current understanding of cuticle 314 dissipating the accumulated strain. <sub>259</sub> mechanobiology. It is therefore unlikely that differences <sub>315</sub> The specimen did not show any evidence of fractures,

2009). Each specimen was subjected to force-controlled, 262 trum in different species; instead, a single mechanism uniaxial loading to fracture at a constant stress rate of 263 (i.e. the modified composite profile) confers increased  $0.05\,\mathrm{N\cdot s^{-1}}$  (detailed in Methods). We then examined 264 flexibility and tensile strength to the rostral apex in the the correspondence between composite structure and me- 265 genus Curculio. he endocuticle demonstrably contributes chanical behavior of the snout, using phylogenetic linear 266 more to rostral tensile strength than the exocuticle, likely mixed effects models to account for phylogenetic non- 267 because of its organization into large bundles of aligned, 268 anisotropic fibers. Consequently, the altered composite In general, the specimens exhibited a non-linear vis- 269 profile of the cuticle in the rostral apex makes the ros-

## LOAD CYCLING OF CURCULIO CARYAE

To confirm that repeated, proscribed straightening of 273 the rostrum does not result in damage to the cuticle, 274 we performed displacement-controlled fatigue testing on 275 a typical female specimen of Curculio caryae, a species 276 that exhibits extreme  $(80 - 90^{\circ})$  rostral curvature. The 277 specimen was then aligned so that uniaxial tension would 278 induce elongation of the apical portion of the rostrum. 279 The strain per cycle was fixed at an amplitude sufficient 280 to completely elongate the snout and generate a tensile 281 load equal to 0.2× the average tensile strength of con-282 specifics in the straightened configuration, at a frequency 283 of 0.33 Hz. The test was terminated after a period of two <sup>284</sup> weeks (ca. 400K cycles) when the strain amplitude ap-

We observed viscoelastic behavior in the rostrum, as 287 indicated by histeresis in the stress-strain relationship dient, it is not possible to correlate measurements from 288 during each cycle. Strain amplitude decreased logariththe fracture surface to the properties of the entire snout. 289 mically with cycle number, and the specimen initially Instead, we found that the uniaxial elastic modulus of the 290 appeared to have deformed plastically during the test. <sup>291</sup> We believed that this indicated damage to the specimen; 292 however, after cleaning the specimen in a 24 hour wash 293 with ethanol and water, we observed that rostrum re-294 turned to its original shape.

We cannot fully account for the stress relaxation of Additionally, we observed a moderate, stiffening size- 296 the rostrum after testing, but we speculate that it arose 310 viscoelastically (REF). After 24 hours soaking in ethanol From these tests, we infer that the gross elastic be- 312 and water, the hydrogen bonds would relax sufficiently to

260 in sclerotization or chitin composition within the cuticle 316 micro-tears, or shear cusps anywhere in the surface of

paper

319 species ( $F_max = 5.02 \,\mathrm{N}$ ). Given this surprising result, 374 cross-sectional areas of endocuticle, this was often ob-320 it appears that the specimen was undamaged by the test- 375 scured by delaminated endocuticular macrofibers. 321 ing. We therefore expect that under normal conditions 376 The first layer of endocuticle usually fractured along 322 in life, repeated bending of the snout does not exceed 377 the same plane as the exocuticle, and ventrally the en-323 the yield strength of the cuticle, and the bending strain 378 docuticle laminae typically converged toward a scarp-like 324 is purely elastic.

## FRACTOGRAPHY OF TEST SPECIMENS

Due to the complex failure modes evident upon examination of the fractured specimens, 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 333 here.

In transverse view, the exocuticle consistently presented a nearly continuous fracture surface, indicative of comparatively brittle failure, due to the helicoidal ar- $_{337}$  rangement of  $\alpha$ -chitin laminae and resultant transverse isotropy of the Bouligand structure (REF). The exocuticle typically appeared to fracture at lower strains than the endocuticle, with shear-cusp formation evident both at the fracture surface and across unbroken endocuticle on the rostral surface adjacent to the plane of fracture. Conversely, the endocuticle exhibited severe delamination, off-axis ply-splitting, and fiber-pulling away from the fracture surface, indicating that endocuticular macrofibers are less brittle than the exocuticle, likely as a consequence of the unidirectional alignment of their  $\alpha$ chitin nanofibrils (REF). Because the exocuticle of weevils is anchored to the endocuticle by cross-linking fibers, the presence of exocuticular shear-cusp formation in uniaxial tension suggests extension-shear coupling within individual endocuticle laminae, which caused ply-splitting via mode II fracture between macrofibers at high strain. This intra-laminar extension-shear coupling also yielded off-axis, in-plane resultant forces as a function of lamina orientation angle; mode III shearing then occurred be-360 mode I/II (transverse tension/intra-laminar shear) frac- 415 exhibits marked reduction in relative thickness of the and splitting in adjacent laminae. 362

370 oriented outward from the center of the invagination, 425 modifications to the cuticle, which have a non-additive

317 the exocuticle, and, furthermore, the tensile strength of 372 the cusps converged toward a prominent scarp where the the specimen was consistent with other members of its 373 crack fronts joined, although in specimens with large

> 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 383 shearing, while macrofibers oriented against the direction of crack propagation primarily displayed fracture by transverse shear along the plane of ply-splitting in 386 adjacent laminae. Because the laminae form a cylinder, contralateral fibers in the same lamina display opposing fracture modes. In addition, the ventrolateral 389 surfaces often exhibited extensive inter-ply delamination 390 and fiber de-bonding in scarp-like prominences, likely due 391 to a combination of tensile failure and shearing along 392 the dorsally-radiating crack front. Dorsally, the coa-393 lescent crack fronts often caused significant de-bonding 394 and ply-splitting followed by broom-like tensile failure; 395 in some specimens, the contralateral crack fronts were 396 out of plane and coalesced via transverse shear through <sup>397</sup> a large dorsal section of cuticle.

> From these patterns we hypothesize that the 399 exocuticle-rich occipital sulci are the most likely site for 400 the initiation of void nucleation and catastrophic failure 401 of the integrated rostral cuticle in cross section, as illus-402 trated in Fig REF. According to this model, structural 403 failure would take place as cracks propagate through the endocuticle from these sutures, which penetrate the en-405 tire thickness of the laminate. Although other, more 406 complex failure modes have been observed, we posit that 407 in live specimens this is the most likely mechanism of 408 tensile failure because typical bending behavior generates 409 tension only along the ventral surface of the rostrum.

#### CONCLUSIONS

The rostrum of *Curculio* is characterized by a discontween laminae with opposing in-plane resultant forces, 412 tinuous composite profile, wherein the cuticle is strongly causing inter-ply delamination. Tensile failure of the 413 differentiated along an anterior-posterior gradient. In macrofiber laminae would ultimately occur via mixed- 414 particular, the cuticle in the apical portion of the rostrum ture due to an increase in applied stress caused by ply- 416 in stacking-angle between pseudo-orthogonal plies of the 417 endocuticle, which are aligned at  $\pm 45^{\circ}$  to the longitudinal At the meso-scale, most specimens fractured along a 418 axis of the snout. Constitutive models demonstrated that single plane across and between the occipital sulci, which 419 an idealized apical cuticle could theoretically achieve 72% are cuticle invaginations that traverse the entire length 420 and 60% reductions in the effective longitudinal memof the rostrum. These sulci increase the volume fraction 421 brane and flexural moduli, respectively, compared to the of exocuticle in the ventral part of the snout and contain 422 cuticle of the head capsule, thus accounting for the oblarge interfaces ideal for void nucleation. The exocuti- 423 served flexibility of the rostral apex in live specimens. cle of the occipital sulci usually displayed shear cusps 424 However, these reductions can only be realized with both 371 continuing dorsolaterally and ventromedially. Ventrally 426 effect on cuticle elasticity, implying that these weevils 427 require both modifications to function properly during 456 ture in CLPT, which predicts extension shear-coupling oviposition.

of the cuticle, while increased stiffness was associated 460 void nucleation and crack initiation. with lower failure strains and reduced fracture toughness. 461 433 Additionally, fatigue testing of a highly-curved rostrum 462 unexpected morphological source of evolutionary con-434 confirmed that the apical portion of the snout can be sub- 463 straint on rostral flexibility, raising the intriguing pos-435 jected to repeated antero-dorsal flexion without evidence 464 sibility that this system evolved primarily via negative 436 of damage. We therefore conclude that the altered com- 465 selection of fracture, rather than positive selection of 442 ations were independent of species membership, we posit 471 endocuticle-dominated rostral apex. This constraint, as  $_{444}$  genus, and that rostral flexibility is achieved exclusively  $_{473}$  rostral apex, and the increased toughness derived from  $_{447}$  we infer that the endocuticle deforms viscoelastically  $_{476}$  in the evolution of the rostrum. 448 along the longitudinal axes of the macrofibers under 449 uniaxial tension, causing ply-splitting and delamination 450 at high strain. However, the overlying exocuticle ex-451 hibits brittle fracture at a comparatively lower strain, due to shearing between the stretching endocuticle fibers to which it is anchored. This outcome is consistent 478

 $(A_{16}, A_{26} \neq 0)$  for individual off-axis macrofiber laminae. Our tensile testing data revealed that increased endo- 458 Based on this pattern of fracture behavior, we identified cuticle thickness correlates with greater fracture strength 459 the exocuticle-rich occipital suture as a common point of

From a biological perspective, these findings reveal an posite profile of the cuticle in the rostral apex makes the 466 flexibility. In particular, the cuticle is invaginated in rostrum simultaneously more flexible and fracture resis- 467 precisely the portion of the snout that experiences the tant, and that this profile allows the rostrum to be flexed 468 greatest degree of tension during antero-dorsal flexion; into a completely straight configuration without exceed- 469 the doubly-thick exocuticle in the invagination thus creing the elastic limits of the cuticle. Because these associ-  $_{470}$  ates an unavoidable, brittle weak-point in the otherwise that the behavior of the cuticle is consistent across the 472 well as the minimization of exocuticle thickness in the through a modified cuticle lay-up in all Curculio species. 474 a thickened endocuticle, all point to avoidance of catas-Based on fractographic analysis of the test-specimens, 475 trophic structural failure as a driving selective pressure

## **METHODS**

Methods, including statements of data availability and with behavior shown in previous studies and is congru- 479 any associated accession codes and references, are avail-455 ent with theoretical consideration of cuticle microstruc- 480 able in the online version of this paper.

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## **AUTHOR CONTRIBUTIONS**

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

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Jason Williams: Conducted tensile and fatigue testing, 530 participated in manuscript preparation. 531

Nikhilesh Chawla: Facilitated microscopy, tensile and 547 532 533 preparation. 534

Nico Franz: Facilitated specimen acquisition and imaging, participated in manuscript preparation. 536

#### ADDITIONAL INFORMATION

Supplementary information is available in the online 538 version of the paper. Reprints and permissions infor-540 mation is available online at www.nature.com/reprints. 556 Correspondence and requests for materials should be addressed to Andrew Jansen.

#### COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests. 544

#### **METHODS**

## Specimen acquisition and taxon sampling

Specimens for use in tensile and fatigue testing came fatigue testing, and participated in manuscript 548 from the Hasbrouck Insect Collection at Arizona State 549 University [ASUC]. This set of specimens was supple-550 mented with material housed in the following collections, using the codens of Arnett et al. (1993)(REF):

> CMNC: Canadian Museum of Nature Collection, Ottawa, Ontario, Canada

> USNM: National Museum of Natural History, Washington, D.C., USA

Cold fracture, semi-thin sectioning, and tensile testing 557 were conducted on randomly chosen female specimens belonging to six species obtainable through field work in the southwestern USA and northwestern Mexico. Taxon sampling was targeted to represent a mixture of disparate radiations and sister taxa with a variety of rostral morphotypes, according to the phylogenetic hypotheses of authors (REF & REF). These six species of Curculio used herein are C. caryae (Horn, 1873), C. humeralis (Casey, 1897), C. proboscideus Fabricius, 1775, C. sulcatulus (Casey, 1897), C. uniformis (LeConte, 1857), and 567 C. victoriensis (Chittenden, 1904).

## Histological sectioning

To illustrate the relative proportions of the cuticle regions in cross-section, serial semi-thin sectioning was conducted on exemplary female specimens of C. humeralis and C. longinasus Chittenden 1927. Live specimens of both species were collected into 95% ethanol for preservation. A female specimen was selected and the rostrum was separated from the head capsule with a fine-edged razor blade. The apical 1/4th of the rostrum was also removed and then discarded. The remaining portion of the rostrum and the head capsule were then embedded in EMbed812, as follows.

The cuticle was first immersed in acetone for 24 hours. and then transferred to a 2:1 mixture of acetone to epoxy resin. Samples remained at 21°C for 12 hours on a shaker table to prevent hardening. These were then transferred into a 1:1 mixture of acetone to resin, following by a 585 1:2 mixture (each for 12 hours and at 21°C on a shaker table), before finally being placed into a silicone mold with pure resin. The mold was placed into an oven heated to 38°C, and the resin was allowed to cure for 24 hours. The resulting blocks were machined to prepare the apical surface of each sample for microtomy.

A Leica Ultracut R Microtome and diamond knife were used to expose a cross-section (transverse plane) of the 593 apical and basal portions of the rostrum and to remove excess material. Semi-thin sections (0.5 µm thick) were

595 kept and stained with toluidine-blue-borax for light mi- 647 the cuticle, simulating the condition of live tissue (see 596 croscopy and imaging.

## Cold-fracture of specimens

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Two pinned female specimens of each species were selected at random and retained for cold-fracturing of the rostrum. The heads of the specimens were removed and cleaned using a 95% ethanol solution and a thin paintbrush. Any muscles protruding from the occipital foramen were removed with a fine-edged razor blade. The antennae were removed directly using forceps to pull the scape (basal section of antenna) from the antennal in-606 sertion. Cleaned specimens were stored at  $-80^{\circ}$ C for 24 607 hours, then fractured using forceps over a chilled alu-608 minum block. To fracture each specimen, the head cap-609 sule and rostrum were each gripped firmly in a pair of forceps; the forceps were then sharply rotated to fracture the base of the snout via dorsal flexion. The rostrum was fractured a second time, after separation from the 613 head capsule, using the same procedure. The segmented 614 specimens were then placed into individual glass vials to 615 protect the fracture surfaces from contamination prior to 616 microscopy.

#### Tensile and fatigue testing

Force-controlled loading to failure Five female spec-619 imens of each species were randomly allocated for use in tensile testing. The head of each specimen was removed, cleaned, and prepared as described above in the cold-fracture protocol. To avoid destroying the delicate, brittle specimens when gripping the ends of each head, a method was devised to create solid handles that could be clamped tightly into grips without risk of damage to the cuticle.

For each head, four 1 cm<sup>2</sup> strips of gaffer tape were cut; these were used as gripping and mounting points for the specimen. A strip of tape would be laid flat, with a large drop of cyanoacrylate glue placed upon the upturned surface. The curved portion of the snout was then placed 632 into the drop such that the straight portion of the rostrum was aligned perpendicular to the edge of the strip. 634 Hardening of the cyanoacrylate effectively embedded the 635 curved portion of the snout in a solid mass, isolating a 636 straight section of the snout (from the base to a point 637 distad of the apex of the scrobe) for testing. A second 638 strip of tape was fixed over this mass with an additional layer of cyanoacrylate to provide a dorsal gripping sur-640 face for the mass, and a small mark was made to indicate the extent of the head inside the mass. This embedding 642 procedure was likewise repeated for the head capsule, re-643 sulting in a finished specimen with anterior and posterior 698 examined using scanning electron microscopy to charac-644 handles for testing.

648 also REF). Once removed for testing, the specimen was 649 gripped using the cyanoacrylate handles at the marked 650 locations immediately beyond the anterior margin of the 651 rostrum and the posterior margin of the head capsule 652 The exposed section of the snout was coated in petroleum 653 jelly using a cotton swab to prevent loss of moisture and stiffening of the specimen during the test. Specimens 655 were loaded in a Tryton 250 Microforce Testing System 656 equipped with a 5N load cell and mechanical clamp grip. 657 All specimens were subjected to force-controlled uniaxial tension at a rate of  $1.0 \,\mathrm{gf}\cdot\mathrm{s}^{-1}$  until failure. Results were 659 reported only for specimens that did not fracture due to 660 strain accumulation at the interface between the rostrum 661 and the cyanoacrylate handles.

Displacement-controlled cyclic loading To confirm 663 that repeated, complete extension of a strongly curved 664 rostrum would not result in fracture of the cuticle, a rep-665 resentative female specimen of C. caryae was allocated 666 for fatigue testing. The head capsule of the specimen was 667 fixed to a push-pin using cyanoacrylate glue; this served 668 as a pedestal and gripping location for the posterior por-669 tion of the specimen. The apex of the rostrum was fixed to a strip of ripstop nylon fabric equal in length to the 671 head using cyanoacrylate glue. As with tensile testing, the specimen was placed in de-ionized water for 24 hours, 673 then coated in petroleum jelly using a cotton swab im-674 mediately prior to load cycling.

The end of the fabric gripped and used to elongate the 676 snout in tension; in this way, the rostrum would return 677 to its original configuration in a springlike manner, as 678 in living specimens, rather than being forced into position. The snout was aligned such that complete elongation of the curved section would take place in tension with minimal off-axis deflection of the un-curved section. 682 The specimen was subjected to displacement-controlled 683 loading sufficient to fully extend the snout and generate 684 a load stress of 1 N, or approximately 20% of the ten-585 sile strength of the species average. Load cycling took place at a rate of 0.33Hz, and was continued for 14 days (ca. 400K cycles), until the tensile stress in the sample approached an asymptotic minimum.

Once the test was concluded, the specimen was placed 690 in a 50% ethanol solution for 24 hours to clean the 691 petroleum jelly from the rostrum. The specimen was 692 examined for surface fractures and micro-tears, then sub-<sub>693</sub> jected to tensile testing via the same protocol as the other 694 specimens to assess whether the cuticle had begun to fa-695 tigue.

#### Specimen imaging and microscopy

The fracture surfaces of cold-fractured specimens were 699 terize the composite profile and microstructure of the ros-Prior to testing, each specimen was placed in de- 700 trum. Fracture behavior was assessed using both light 646 ionized water for 24 hours to allow full saturation of 701 microscopy and SEM to image the fracture surfaces of 702 the specimens in transverse view. Electron microscopy 739 the laminate coordinate system, yielding a reduced transwas conducted using a JEOL JSM6300 scanning elec- 740 formed stiffness matrix according to: tron microscope, while light microscopy was conducted using a Leica M205 C stereomicroscope and attached computer running the software Leica Application Suite 707 (LAS), as well as a Visionary Digital Passport II system 708 using a Canon EOS Mark 5D II camera outfitted with interchangeable microptic lenses. Specimen length, layer 710 thicknesses, macrofiber orientation angles, and cross-711 sectional areas were assessed in the Leica Application 712 Suite and in Adobe Illustrator using pixel-wise measure-713 ments multiplied by a scaling factor for the image.

# Constitutive modeling of the cuticle

General Approach

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Cuticle profile of model The cuticle of C. longinasus 717 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 721 and endocuticle. In the apical cuticle we assumed the 222 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 727 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 calculating the 2D reduced stiffness matrix for each part of the cuticle. For orthotropic materials with the principal axes parallel to the ply edges, the reduced stiffness matrix is 735 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:

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

<sub>738</sub> formed to account for the layer orientation angle  $\theta$  within <sub>759</sub> of sufficiently high aspect ratio to minimize the Poisson

$$[\bar{Q}] = [T]^{-1}[Q][T]^{-T},$$
 (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 permuta-743 tions, we calculate the extensional stiffness matrix [A], bending stiffness matrix [D], and bending-extension coupling matrix B for each laminate consisting of n layers at  $_{746}$  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 749 forces  $\{N\}$  and bending moments  $\{M\}$  to mid-surface 750 strains and curvatures  $\{\epsilon^{\circ}\}$  and  $\{\kappa\}$ , respectively, in the 751 laminate according to the following relationship:

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

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

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 756 (1) the laminate experiences pure axial loading and For each layer k, the reduced stiffness matrix is trans-  $M_{xy} = 0$  and  $M_{yy} = 0$  a

760 effect and anisotropic shear coupling (i.e., below we ef-761 fectively let  $A_{12}^*=A_{16}^*=0$  and  $D_{12}^*=D_{16}^*=0$ ), then 762 we can calculate the in-plane effective flexural and axial Young's moduli of the laminate along the x-axis.

For axial Young's modulus of the laminate, we first 765 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:

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

and, by inverting this equation (let  $A^* = A^{-1}$ ) and 768 substituting based on the assumptions above, we infer:

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

We therefore define Young's modulus for effective axial 770 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 808 1772 laminate, we first specify the moment-curvature relation 809 773 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-775 angular cross-section is:

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

Given the assumption that  $M_{yy} = M_{xy} = 0$ , the mo-777 ment along the x-axis is related to the moment of the 778 beam by:

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

Thus, given the assumption that  $D_{12} = D_{16} = 0$ , 824 each of the three models. 780 Young's modulus for the effective transverse flexural elas-781 ticity of the laminate can be found by making Eq. 13 spe-782 cific to transverse flexure of the x-axis and rearranging 783 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 829 784 785 to:

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

General approach To explore the relationships be-788 tween the composite structure and mechanical proper-(9) 789 ties of the cuticle, we fit phylogenetic linear mixed-effects 790 models to the data using maximum likelihood estimation. In order to control for phylogenetic non-independence in 792 the data, we included the species of each specimen as a 793 random effect in all models. We also allowed for correlation in the error term of the models, as specified by 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 nor-799 mally distributed and homoscedastic. In all models, we 800 tested whether the inclusion of phylogenetic correlation 801 in the model error produced significantly better model fit, using a likelihood-ratio test and  $R_{\sigma}^2$ -difference test between the fully-specified model and a model lacking 804 the phylogenetic effect.

Hypothesis testing The following three hypotheses 806 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-816 sectional area of endocuticle and exocuticle at the site 817 of fracture as fixed effects, including an interaction term, and with maximum tensile force sustained prior to fracture as a response variable. This model was then com-820 pared to models with only cross-sectional area of either 821 endocuticle or exocuticle as the sole fixed effect in the (15) 822 model. We then tested the first hypothesis by using <sub>823</sub> likelihood-ratio tests and  $R^2_{\beta_*}$ -difference tests between

> The hypothesis that Model selection and fitting Estimating phylogenetic signal

> > Code availability

Data availability

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# REFERENCES

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