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The acorn weevil snout exhibits remarkable flexibility and toughness that are derived from from the microarchitecture of its exoskeleton. Here we characterize modifications to the composite profile of the rostral cuticle that simultaneously enhance the flexibility and fracture toughness of the distal portion of the snout. Using Classical Laminate Plate Theory (CLPT) we estimate the effect of these modifications on the elastic behavior of the exoskeleton. Additionally, changes in the relative layer thicknesses and orientation angles of layers in the exoskeleton are related to the tensile behavior of the snout in six species of diverse morphology, and we demonstrate that a highly curved rostrum can be completely straightened without structural damage. We demonstrate that increased endocuticle thickness is strongly correlated with increased tensile strength in the snout. Consequently, snout stiffness is shown to be inversely correlated with fracture toughness. Finally, we identify exocuticle rich invaginations both as a likely site of crack initiation in tensile failure and as a source of morphological constraint on the evolution of the snout.

Although α -chitin is brittle and strongly anisotropic, beetle cuticle is simultaneously rigid and tough due to its unique laminate microstructure (see kamp, vincent for 15 review), which we characterize in detail below. Impact-16 prone areas and exaggerated structures in arthropods 17 generally exhibit cuticle organization that resists deformation and fracture; however, acorn weevils in the genus Curculio Linnaeus, 1758¹ instead exhibit unusual distal flexibility in an elongate extension of the head called the rostrum (snout). The rostrum is a hollow, strongly curved (over 90° in some species), cylindrical, exoskeletal extension of the otherwise nearly-spherical head, which bears at its apex the terminal chewing mouthparts. This structure is used by the female to feed and to excavate sites for egg-laying (oviposition, see Fig. 1c), and can be repeatedly bent without evident damage despite being composed of the same material as other rigid body parts. By maintaining constant pressure on the snout and rotating around the bore-hole, females are able to flex the rostrum into a straightened configuration and produce a linear channel into the fruit; a single female may prepare hundreds of such sites as an adult.

While this behavior has been observed in many species of Curculio, it was unclear how the rostrum of female acorn weevils can withstand the repeated, often extreme

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The exoskeleton of Coleoptera (beetles) is a 37 bending incurred during the process of egg-chamber ex-8 hierarchically-structured fibrous composite characterized 38 cavation. In this study we characterize the composite 9 by variously arranged α-chitin (N-acetylglucosamine) 39 profile of the rostral cuticle to account for the observed 10 nanofibrils embedded in a heterogeneous protein matrix. 40 flexibility of the snout. We show that the relative layer 41 thicknesses and fiber orientation angles of the exocuticle 42 and endocuticle of the rostrum are strongly differenti-43 ated from the head capsule and other body parts, and 44 we estimate the effect of these differences on the elas-45 ticity of the cuticle using Classical Laminate Plate The-46 ory (CLPT). Because recent studies have shown that the 47 yield strength of the beetle exoskeleton is lower in ten-48 sion than compression, we perform a comparative analy-49 sis of the ultimate tensile strength of the rostrum across 50 species and snout morphotypes; we also report the re-51 sults of displacement-controlled load cycling of the snout 52 in a species with strongly curved morphology. We relate 53 an observed increase in the volume fraction of endocuti-54 cle in the rostrum to higher tensile strength at the ros-55 tral apex in all tested species, and find that a strongly 56 curved rostrum can be flexed repeatedly without harm 57 to the structure.

> We additionally describe the fracture mechanics of the 59 snout and consider how modification of the cuticle may 60 prevent crack formation during oviposition. Based on 61 our findings, we posit that the composite profile of the 62 rostral apex enables the snout to undergo repeated dis-63 tal flexion while remaining within the elastic limits of the 64 material, mitigating the risk of structural damage, and 65 without evident alteration of the mechanical properties 66 of the individual components of the cuticle across the 67 structure and between species. Thus, the flexibility and 68 tensile strength of the rostrum appear to be derived ex-69 clusively from modification of the composite architecture 70 of the exoskeleton. To our knowledge, this is the first 71 time that a modified composite profile has been reported $_{\rm 72}$ as a means of enhancing structural elasticity in the insect 73 exoskeleton.



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.

MICROSTRUCTURE OF THE CURCULIO ROSTRUM

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In arthropods (including beetles), the exocuticle is other, forming a quasi-isotropic laminate known as the Bouligand structure [7–9]. This layout effectively mitigates the strong anisotropy of α-chitin to yield a versatile building material for the exoskeleton. Beetle endocuticle, however, is unique among arthropods and is 86 comprised of large (1-5 µm diameter) unidirectional bun-87 dles of chitin, called macrofibers. Chitin macrofibers 88 are orthotropic (axial: $E_1 = 8.5 \,\mathrm{GPa}$, transverse: cally, adjacent macrofiber laminae are paired and pseudoorthogonal (i.e., angled approx. 90° to each other, see Figs. REF), with a constant stacking angle between pairs, although other configurations have been observed [11, 12]. This geometric sequence of the macrofiber laminae yields an approximately transversely isotropic composite, similar to the Bouligand structure. Notably, the 98 resulting laminate is less rigid than the exocuticle, but exhibits greater toughness because the pseudo-orthogonal plies effectively inhibit crack formation and propagation between successive layers [11–13]. 101

Serial thin sectioning and scanning electron microscopy of fractured Curculio specimens revealed that endocuticle in the head capsule fits this general profile, with an angle of approximately 30° between successive pairs of pseudoorthogonal plies. Additionally, in the head capsule, the thickness of the exocuticle and endocuticle in cross section are nearly equal (typically between 20-30 µm). However, we found that the cuticle composite lay-up of the rostral apex is strongly differentiated from the head cap- 166 sule (see Fig. REF). Distally the exocuticle is reduced ¹⁶⁷ to a thin shell (ca. 5 µm), with the endocuticle thickened to offset this reduction and maintain a constant 168 119 laminate). We previously identified these modifications 174 controlled, uniaxial loading to fracture at a constant ₁₂₀ to the composite structure of the cuticle within a sin-₁₇₅ stress rate of 1.0 gf·s⁻¹ (detailed in Methods). In gen-121 gle species, C. longinasus Chittenden, 1927 [1?]; how- 176 eral, the specimens exhibited a non-linear viscoelastic re-122 ever this composite profile has now been uncovered in 177 sponse curve characterized by a sharp increase in stress at scrobe exhibits a gradual transition in composite profile 182 more directly with increasing strain. 128 along an anterior-posterior gradient.

131 Young's moduli of the cuticle in the rostral apex and 132 head capsule using Classical Laminate Plate Theory 133 (CLPT), as detailed in our methods. The effective 134 elastic constants of the cuticle regions of C. longina-77 comprised of numerous unidirectional laminae of chitin 135 sus (estimated previously, see REF) were used to connanofibrils; each layer is the thickness of a single fiber 136 struct constitutive equations for the entire cuticle of (2-4 nm) embedded in a proteinaceous matrix. These lay- 137 that species. The cuticle of the head capsule was ers are stacked at a more or less constant angle to each 138 estimated to have membrane and flexural moduli of $E_m = 4.77 \,\text{GPa}$ and $E_f = 6.04 \,\text{GPa}$, respectively; 140 however, in the rostral apex we found that these values 141 were reduced by approximately 72% and 60%, respec-142 tively $(E_m = 1.36 \,\text{GPa}, E_f = 2.44 \,\text{GPa})$. Two hypo- $_{143}$ thetical cuticle lay-ups were also modeled to individually 144 assess the contributions of either modified layer thickness 145 or stacking angle sequence to cuticle flexibility. The ef- $_{146}$ fective moduli of a configuration with the angle stacking $_{89}$ E_2 = E_3 = 0.52 GPa [1]) and arranged in uni- $_{147}$ sequence of typical cuticle (i.e., in the head capsule) but 90 directional plies, seen in Figs. REF [12, 13]. Typi- 148 possessing the layer thicknesses of the rostral apex were calculated as $E_m = 3.73 \,\mathrm{GPa}, E_f = 4.31 \,\mathrm{GPa}$, repre- $_{150}$ senting 22% and 29% decreases from unmodified cuticle, 151 respectively. Similarly, a hypothetical cuticle with the 152 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) had effective elastic moduli of $E_m = 3.77 \,\text{GPa}, E_f = 5.76 \,\text{GPa}, \text{ representing } 21\%$ and 4.7% decreases from unmodified cuticle, respectively. $_{157}$ Each of the cuticle modifications noted in the rostral apex 158 individually decreased the elastic moduli of the cuticle; 159 however, they appear to have a synergistic combined ef- $_{160}$ fect on cuticle elasticity, rather than a simple additive 161 effect. This result suggests that both modifications are 162 necessary in order for the snout to function properly in 163 the living animal, where the combined effect allows the 164 rostrum to bend until completely straight without frac-

FORCE-CONTROLLED LOADING TO **FRACTURE**

To better characterize the failure behavior of the roscuticle thickness (ca. $50\,\mu\mathrm{m}$ total) throughout its length. $_{169}$ trum, we performed tensile testing on the snouts of six Moreover, the endocuticular macrofibers exhibit no ro- 170 Curculio species that representing a mixture of closely tation between successive pseudo-orthogonal plies, and 171 and distantly related taxa (see Methods). Each speciare oriented at approximately $\pm 45^{\circ}$ to the longitudinal $_{172}$ men was immersed in di-H₂O for 24 hours, to simulate axis of the snout (i.e., an antisymmetric [±45°] angle-ply 173 the living tissue (see REFS), then subjected to forcethe rostrum of six additional, phylogenetically disparate, 178 higher strains, terminating in brittle fracture. We postuspecies (detailed in methods), indicating that this is likely 179 late that strain hardening occurs as the longitudinal axis a genus-wide trait. In all examined species, the portion 180 of the macrofibers becomes more closely aligned to the of the snout between the head capsule and apex of the 181 cylindrical axis of the rostrum, thereby resisting tension

We examined the correspondence between composite 183 We estimated the effect of differential cuticle orga- 184 structure and mechanical behavior of the snout using 130 nization on uni-axial membrane and transverse-flexural 185 phylogenetic linear mixed effects models (PGLMM) to 186 account for phylogenetic non-independence in residual 241 minimal off-axis deflection of the un-curved section. The 193 specimen and the ratio of exocuticle to endocuticle cross- 248 totic minimum. The rostrum behaved viscoelastically, 195 predicts a positive association between the proportion of 250 during each cycle. Strain amplitude decreased logarithanterior-posterior gradient, it is not possible to correlate 256 turned to its original shape. measurements from the fracture surface to the properties 257 of the entire snout.

214 is that longer snouts may have a longer transition gradi- 269 for the material to completely relax after deformation, the 215 ent from basal to apical profile to reinforce the junction 270 specimen would slowly accumulate strain and deform vis-216 between the rostrum and head capsule against buckling. 271 coelastically (REF-stress history dependence paper). Afthat the gross elastic behavior of the cuticle is consis- 275 strain. tent across the genus, and that a single mechanism (i.e. 276 227 than the exocuticle, likely because of its organization into 282 ing. We therefore expect that under normal conditions 228 large bundles of aligned, anisotropic fibers, leading to a 283 in life, repeated bending of the snout does not exceed the 229 trade-off between rigidity and toughness. Consequently, 284 yield strength of the cuticle. 230 the altered composite profile of the cuticle in the rostral 231 apex makes the rostrum simultaneously more flexible and 232 fracture resistant.

III. LOAD CYCLING OF CURCULIO CARYAE

a typical female specimen of Curculio caryae, a species 292 describe here. 239 specimen was aligned so that uniaxial tension would in- 294 sented a nearly continuous fracture surface, character-240 duce elongation of the distal portion of the rostrum with 295 istic of comparatively brittle failure, presumably due to

variance, with species membership included as a random 242 strain per cycle was fixed at an amplitude sufficient to effect. Our models show that the maximum force sus- 243 completely elongate the snout and generate a tensile load tained at the site of failure was strongly correlated with 244 of 1.0 N in the straightened configuration (ca. 20% ultithe cross-sectional area of the endocuticle, and not the 245 mate strength) at a frequency of 0.33 Hz. The test was exocuticle, at that site. Accordingly, there was a nega- 246 terminated after a period of two weeks (ca. 400K cycles) tive correlation between ultimate tensile strength of the 247 when the stress amplitude appeared to reach an asympsectional area at the site of fracture. Although CLPT 249 indicated by histeresis in the stress-strain relationship exocuticle and stiffness of a generalized cuticle, we found 251 mically with cycle number, and the specimen initially no correspondence between the cross-sectional properties 252 appeared to have deformed plastically during the test. of the fracture site and the gross behavior of the entire 253 We believed that this indicated damage to the specimen; rostrum. Because the cross-sectional areas of the cuti- 254 however, after cleaning the specimen in a 24 hour wash cle regions vary across the length of the head along an 255 with ethanol and water, we observed that rostrum re-

We cannot fully account for the stress relaxation of 258 the rostrum after testing, but we speculate that it arose Instead, we found that the uniaxial elastic modulus 259 from the general mechanism associated with cuticle vis-(low strain: E_{low}) and secant modulus at failure (E_{sec}) 250 coelasticity. The endocuticle is made of aligned α -chitin were inversely correlated with ultimate strain and frac- 261 nanofibrils whose crystalline structure is enforced by hyture toughness, implying that stiffer specimens, and by 262 drogen bonds between individual chitin chains along their extension, stiffer cuticle profiles, are generally more brit- 263 length; macroscopic viscoelastic behavior results from tle. We also observed a moderate, stiffening size-effect 264 slippage between these chains as the hydrogen bonds with respect to rostral length, contrary to our expecta- 265 break and reform in response to shearing between the tion that a longer, more strongly curved (REFS) snout 266 chitin molecules. We posit that repeated strain may have would require increased flexibility to avoid fracture dur- 267 caused such slippage in the endocuticle of the rostral apex ing oviposition. One possible explanation for this trend 268 during the fatigue test; however, without sufficient time Thus, Young's modulus of the rostrum would be com- 272 ter immersion in ethanol and water, the hydrogen bonds paratively higher in these species because of the higher 273 would relax sufficiently to allow the specimen to return volume fraction of exocuticle. From these tests, we infer 274 to its original configuration, dissipating the accumulated

The specimen did not show any evidence of fractures, the modified composite profile) confers increased flexibil- 277 micro-tears, or shear cusps anywhere in the surface of ity and tensile strength to the rostral apex in the genus 278 the exocuticle, and, furthermore, the tensile strength of Curculio. in agreement with our current understanding 279 the specimen was consistent with other members of its of cuticle mechanobiology. In addition, the endocuticle 280 species ($F_m ax = 5.02 \,\mathrm{N}$). Given this surprising result, demonstrably contributes more to rostral tensile strength 281 it appears that the specimen was undamaged by the test-

FRACTOGRAPHY OF TEST SPECIMENS

Although it was not always possible to identify void 287 nucleation and crack initiation sites given the complex 288 failure modes evident in the fractured specimens, we ob-To confirm that repeated, proscribed straightening of 289 served several patterns characteristic of both the microthe rostrum does not result in damage to the cuticle, 290 scale behavior of the cuticle and the meso-scale behavior we performed displacement-controlled fatigue testing on 291 of the rostrum during uniaxial tensile failure, which we

that exhibits extreme $(80-90^{\circ})$ rostral curvature. The 293 In transverse view, the exocuticle consistently pre-

296 the relatively homogeneous arrangement of α-chitin lam- 354 in some specimens, the contralateral crack fronts were typically appeared to fracture at lower strains than the 356 a large dorsal section of cuticle. endocuticle, with shear-cusp formation evident both at 357 From these patterns we hypothesize that the ticle laminae, implying that ply-splitting occurred via 368 tension only along the ventral surface of the rostrum. mode II fracture between macrofibers at high strain. We Hypothesize that intra-laminar extension-shear cou-313 pling also yielded off-axis, in-plane resultant forces as 314 a function of lamina orientation angle; mode III shear-315 ing then occurred between laminae with opposing in-316 plane resultant forces, causing the observed inter-ply delamination. Tensile failure of the macrofiber laminae would ultimately occur via mixed-mode I/II (transverse 319 tension/intra-laminar shear) fracture due to an increase in applied stress caused by ply-splitting in adjacent laminae. 321

At the meso-scale, most specimens fractured along a 322 323 single plane across and between the occipital sulci, which are cuticle invaginations that traverse the entire length of the rostrum. These sulci increase the volume fraction of exocuticle in the ventral part of the snout and contain large interfaces ideal for void nucleation. The exocuticle of the occipital sulci usually displayed shear cusps oriented outward from the center of the invagination, continuing dorsolaterally and ventromedially. Ventrally the cusps converged toward a prominent scarp where the crack fronts joined, although in specimens with large cross-sectional areas of endocuticle, this was often obscured by delaminated endocuticular macrofibers.

342 shearing, while macrofibers oriented against the direc- 397 genus, and that rostral flexibility is achieved exclusively 343 tion of crack propagation primarily displayed fracture 398 through a modified cuticle lay-up in all Curculio species. 344 by transverse shear along the plane of ply-splitting in 399 This conclusion raises the intriguing possibility that a 345 adjacent laminae. Because the laminae form a cylin- 400 single ancestral shift in cuticle organization at the ros-347 posing fracture modes. In addition, the ventrolateral 402 – opened a large region of morphospace for exploration, 348 surfaces often exhibited extensive inter-ply delamination 403 resulting in present day diversity of the genus. and fiber de-bonding in scarp-like prominences, likely due 404 to a combination of tensile failure and shearing along 405 we infer that the exocuticle exhibits brittle fracture at the dorsally-radiating crack front. Dorsally, the coa- 406 a comparatively low strain, due to shearing between the 352 lescent crack fronts often caused significant de-bonding 407 endocuticle macrofibers to which it is anchored; these

inae in the Bouligand structure (REF). The exocuticle 355 out of plane and coalesced via transverse shear through

the fracture surface and across exocuticle (but with en- 358 exocuticle-rich occipital sulci are the most likely site for docuticle intact) adjacent to the plane of fracture. Con- 359 the initiation of void nucleation and catastrophic failure versely, the endocuticle exhibited severe delamination, 360 of the integrated rostral cuticle in cross section, as illusoff-axis ply-splitting, and fiber-pulling away from the 361 trated in Fig REF. According to this model, structural fracture surface, indicating the relatively high tough- 362 failure would take place as cracks propagate through the ness of the unidirectional α -chitin organization within the 363 endocuticle from these sutures, which penetrate the enmacrofibers (REF). Because the exocuticle of weevils is 364 tire thickness of the laminate. Although other, more anchored to the endocuticle by cross-linking fibers, ex- 365 complex failure modes have been observed, we posit that ocuticular shear-cusp formation in uniaxial tension sug- 366 in live specimens this is the most likely mechanism of gests extension-shear coupling within individual endocu- 367 tensile failure because typical bending behavior generates

CONCLUSIONS

The rostrum of Curculio is characterized by a discon-371 tinuous composite profile, wherein the cuticle is strongly 372 differentiated in terms of relative layer thicknesses and 373 orientation angles along an anterior-posterior gradient. 374 These modifications are sufficient to achieve marked re-375 duction in the effective membrane and flexural moduli 376 of the cuticle (72% and 60%, respectively) in constitu-377 tive models based on CLPT, thereby accounting for the $_{\rm 378}$ observed flexibility of the rostral apex in live specimens. 379 However, these reductions can only be realized with both 380 modifications to the cuticle, which have a non-additive 381 effect on cuticle elasticity, implying that these weevils 382 require both modifications to function properly during 383 oviposition. Likewise, tensile and fatigue testing have $_{384}$ revealed a trade-off between stiffness and fracture resis-385 tance (as measured by ultimate strain and toughness), 386 mediated by the relative proportion of endocuticle in the 387 laminate. The altered composite profile of the cuticle in 388 the rostral apex makes the rostrum simultaneously more 389 flexible and fracture resistant, permitting the structure The first layer of endocuticle usually fractured along 390 to be flexed without exceeding the elastic limits of the the same plane as the exocuticle, and ventrally the en- 391 cuticle. To our knowledge, this is the first time that docuticle laminae typically converged toward a scarp-like 392 the composite profile of the cuticle has been related to region characterized by severe delamination and numer- 393 a gradient in elasticity and tensile performance across a ous de-bonded macrofibers. Additionally, macrofibers 394 cuticular structure in arthropods. Because these associaligned with the direction of crack propagation exhib- 395 ations were independent of species membership, we posit ited extensive ply-splitting with intermittent transverse 396 that the behavior of the cuticle is consistent across the der, contralateral fibers in the same lamina display op- 401 tral apex - yielding higher flexibility and tensile strength

Based on fractographic analysis of the test-specimens, 353 and ply-splitting followed by broom-like tensile failure; 408 macrofibers fail at higher strain as a result of mixed-

409 mode shearing and tensile fracture within and between 426 of flexibility. In particular, the cuticle is invaginated in 410 laminae. This outcome is consistent with behavior shown 427 precisely the portion of the snout that experiences the 411 in previous studies and is congruent with theoretical con- 428 greatest degree of tension during antero-dorsal flexion; 412 sideration of cuticle microstructure in CLPT, which pre- 429 the doubly-thick exocuticle in the invagination thus credicts extension shear-coupling $(A_{16}, A_{26} \neq 0)$ for individ-430 ates an unavoidable, brittle weak-point in the otherwise 414 ual off-axis macrofiber laminae. These results imply that 431 endocuticle-dominated rostral apex. This constraint, as 415 fracture initiation occurs in the comparatively brittle ex-432 well as the minimization of exocuticle thickness in the 416 ocuticle, and that the reduction in exocuticle thickness 433 rostral apex, and the increased toughness derived from a 417 in the snout apex might serve to mitigate crack forma- 434 thickened endocuticle, lead us to consider the avoidance 418 tion in rostral bending. Based on this pattern of frac- 435 of catastrophic structural failure as a driving selective 419 ture behavior, we identified the exocuticle-rich occipital 436 pressure in the evolution of the rostrum. 420 suture as a common point of void nucleation and crack 421 initiation. From a biological perspective, these findings 422 reveal an unexpected morphological source of evolution- 437 423 ary constraint on rostral flexibility, raising the intriguing 424 possibility that this system evolved primarily via nega-438 425 tive selection of fracture, rather than positive selection 439 any associated accession codes and references, are avail-

VI. METHODS

Methods, including statements of data availability and 440 able in the online version of this paper.

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AUTHOR CONTRIBUTIONS

M.A.J. conducted sectioning and staining, microscopy 487 and imaging, tensile and fatigue testing, statistical 488 analysis, and participated in manuscript preparation. 489 J.W. conducted tensile and fatigue testing and partic-490 ipated in manuscript preparation. N.C. facilitated mi-⁴⁹¹ croscopy, tensile and fatigue testing, and participated in manuscript preparation. N.M.F. facilitated specimen 493 acquisition and imaging and participated in manuscript 494 preparation.

ADDITIONAL INFORMATION

Supplementary information is available in the online version of the paper. Reprints and permissions infor-498 mation is available online at www.nature.com/reprints. Correspondence and requests for materials should be addressed to M.A.J.

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

METHODS

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Specimen acquisition and taxon sampling

Specimens for use in tensile and fatigue testing came 506 from the Hasbrouck Insect Collection at Arizona State University [ASUC]. This set of specimens was supplemented with material housed in the following collections, using the codens of Arnett et al. (1993)(REF):

511 Ontario, Canada

512 D.C., USA 513

520 morphotypes, according to the phylogenetic hypotheses 572 specimens were then placed into individual glass vials to used herein are C. caryae (Horn, 1873), C. humeralis 574 microscopy. (Casey, 1897), C. proboscideus Fabricius, 1775, C. sul-524 catulus (Casey, 1897), C. uniformis (LeConte, 1857), and 525 C. victoriensis (Chittenden, 1904).

Histological sectioning

529 ducted on exemplary female specimens of C. humeralis 581 brittle specimens when gripping the ends of each head, 530 and C. longinasus Chittenden 1927. Live specimens of 582 a method was devised to create solid handles that could vation. A female specimen was selected and the rostrum 584 the cuticle. was separated from the head capsule with a fine-edged 585 For each head, four 1 cm² strips of gaffer tape were cut; removed and then discarded. The remaining portion of 587 specimen. A strip of tape would be laid flat, with a large the rostrum and the head capsule were then embedded 588 drop of cyanoacrylate glue placed upon the upturned surin EMbed812, as follows.

539 and then transferred to a 2:1 mixture of acetone to epoxy 591 trum was aligned perpendicular to the edge of the strip. 540 resin. Samples remained at 21°C for 12 hours on a shaker 592 Hardening of the cyanoacrylate effectively embedded the 542 into a 1:1 mixture of acetone to resin, following by a 594 straight section of the snout (from the base to a point 543 1:2 mixture (each for 12 hours and at 21°C on a shaker 595 distad of the apex of the scrobe) for testing. A second 544 table), before finally being placed into a silicone mold 596 strip of tape was fixed over this mass with an additional surface of each sample for microtomy.

550 used to expose a cross-section (transverse plane) of the 602 handles for testing. ₅₅₁ apical and basal portions of the rostrum and to remove ₆₀₃ Prior to testing, each specimen was placed in de-552 excess material. Semi-thin sections (0.5 µm thick) were 604 ionized water for 24 hours to allow full saturation of

553 kept and stained with toluidine-blue-borax for light mi-554 croscopy and imaging.

Cold-fracture of specimens

Two pinned female specimens of each species were se-557 lected at random and retained for cold-fracturing of the 558 rostrum. The heads of the specimens were removed and 559 cleaned using a 95% ethanol solution and a thin paint-CMNC: Canadian Museum of Nature Collection, Ottawa, 560 brush. Any muscles protruding from the occipital fora-561 men were removed with a fine-edged razor blade. The ⁵⁶² antennae were removed directly using forceps to pull the USNM: National Museum of Natural History, Washington, 563 scape (basal section of antenna) from the antennal in-₅₆₄ sertion. Cleaned specimens were stored at -80° C for 24 565 hours, then fractured using forceps over a chilled alu-Cold fracture, semi-thin sectioning, and tensile testing 566 minum block. To fracture each specimen, the head capwere conducted on randomly chosen female specimens 567 sule and rostrum were each gripped firmly in a pair of belonging to six species obtainable through field work in 568 forceps; the forceps were then sharply rotated to fracture the southwestern USA and northwestern Mexico. Taxon 569 the base of the snout via dorsal flexion. The rostrum sampling was targeted to represent a mixture of dis- 570 was fractured a second time, after separation from the parate radiations and sister taxa with a variety of rostral 571 head capsule, using the same procedure. The segmented of authors(REF & REF). These six species of Curculio 573 protect the fracture surfaces from contamination prior to

Tensile and fatigue testing

Force-controlled loading to failure Five female spec-577 imens of each species were randomly allocated for use 578 in tensile testing. The head of each specimen was re-To illustrate the relative proportions of the cuticle re- 579 moved, cleaned, and prepared as described above in the gions in cross-section, serial semi-thin sectioning was con- 580 cold-fracture protocol. To avoid destroying the delicate, both species were collected into 95% ethanol for preser- 583 be clamped tightly into grips without risk of damage to

razor blade. The apical 1/4th of the rostrum was also 586 these were used as gripping and mounting points for the 589 face. The curved portion of the snout was then placed The cuticle was first immersed in acetone for 24 hours, 590 into the drop such that the straight portion of the rostable to prevent hardening. These were then transferred 593 curved portion of the snout in a solid mass, isolating a with pure resin. The mold was placed into an oven heated 597 layer of cyanoacrylate to provide a dorsal gripping surto 38°C, and the resin was allowed to cure for 24 hours. 598 face for the mass, and a small mark was made to indicate The resulting blocks were machined to prepare the apical 599 the extent of the head inside the mass. This embedding 600 procedure was likewise repeated for the head capsule, re-A Leica Ultracut R Microtome and diamond knife were 601 sulting in a finished specimen with anterior and posterior

605 the cuticle, simulating the condition of live tissue (see 660 microscopy and SEM to image the fracture surfaces of 609 rostrum and the posterior margin of the head capsule 604 using a Leica M205 C stereomicroscope and attached 610 The exposed section of the snout was coated in petroleum 665 computer running the software Leica Application Suite 611 jelly using a cotton swab to prevent loss of moisture and 666 (LAS), as well as a Visionary Digital Passport II system 612 stiffening of the specimen during the test. Specimens 667 using a Canon EOS Mark 5D II camera outfitted with 613 were loaded in a Tryton 250 Microforce Testing System 668 interchangeable microptic lenses. Specimen length, layer 614 equipped with a 5N load cell and mechanical clamp grip. 669 thicknesses, macrofiber orientation angles, and cross-₆₁₇ pling interval of 0.1 s. Engineering stresses and strains ₆₇₂ ments multiplied by a scaling factor for the image. were reported only for specimens that did not fracture due to strain accumulation at the interface between the rostrum and the cyanoacrylate handles.

Displacement-controlled cyclic loading To confirm 622 that repeated, complete extension of a strongly curved 623 rostrum would not result in fracture of the cuticle, a rep-624 resentative female specimen of C. carvae was allocated 625 for fatigue testing. The head capsule of the specimen was fixed to a push-pin using cyanoacrylate glue; this served as a pedestal and gripping location for the posterior portion of the specimen. The apex of the rostrum was fixed to a strip of ripstop nylon fabric equal in length to the head using cyanoacrylate glue. As with tensile testing, the specimen was placed in de-ionized water for 24 hours, then coated in petroleum jelly using a cotton swab immediately prior to load cycling.

The end of the fabric gripped and used to elongate the 635 snout in tension; in this way, the rostrum would return to its original configuration in a spring-like manner, as in living specimens, rather than being forced to return to the initial 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. The specimen was subjected to displacementcontrolled loading sufficient to fully extend the snout and generate a load stress of 1 N, or approximately 20% of the tensile strength of the species average. Load cycling took 645 place at a rate of 0.33Hz, and was continued for 14 days (ca. 400K cycles), until the tensile stress in the sample 647 approached an asymptotic minimum.

Once the test was concluded, the specimen was placed 649 in a 50% ethanol solution for 24 hours to clean the 650 petroleum jelly from the rostrum. The specimen was examined for surface fractures and micro-tears, then sub-652 jected to tensile testing via the same protocol as the other ₆₅₃ specimens to assess whether the cuticle had begun to fa-654 tigue.

Specimen imaging and microscopy

655

659 trum. Fracture behavior was assessed using both light 714 laminate.

also REF). Once removed for testing, the specimen was 661 the specimens in transverse view. Electron microscopy gripped using the cyanoacrylate handles at the marked 662 was conducted using a JEOL JSM6300 scanning eleclocations immediately beyond the anterior margin of the 663 tron microscope, while light microscopy was conducted All specimens were subjected to force-controlled uniaxial 670 sectional areas were assessed in the Leica Application tension at a rate of $1.0\,\mathrm{gf\cdot s^{-1}}$ until failure, with a sam- $_{671}$ Suite and in Adobe Illustrator using pixel-wise measure-

Constitutive modeling of the cuticle

General Approach The effective uni-axial membrane 675 and transverse flexural elastic moduli of idealized cuti-676 cle organizations representing both the rostral apex and 677 head capsule were estimated using Classical Laminate 678 Plate Theory (CLPT). The composite profiles of both 679 types of cuticle were idealized using the layer thicknesses $_{680}$ and stacking sequences observed in C. longinasus. This 681 particular species was chosen because we derived the ef-682 fective elastic constants of the individual components of 683 the cuticle in previous work. In addition, C. longinasus 684 exhibits a profile that is typical and representative for the 685 genus Curculio, based on examination of the six species used for tensile testing, as described previously.

For C. longinasus the total thickness of the cuticle 688 in the head and rostrum is roughly 50 µm, as in most 689 specimens of the other examined species. In the head 690 capsule, the exocuticle occupies between 30-50\% of 691 the through-thickness of the laminate, with the re-692 maining thickness nearly evenly divided between 12 693 layers of endocuticle. We use the maximum (50%)694 of through-thickness, or 25 µm) for the model, since 695 the cuticle appears to deviate from this value only 696 in sulci (grooves), pores, and other scattered features 697 of surface sculpture. The macrofiber laminae of the 698 endocuticle were assigned a stacking sequence sequence of $699 \ 0^{\circ}, 90^{\circ}, 30^{\circ}, -60^{\circ}, 60^{\circ}, -30^{\circ}, 90^{\circ}, 0^{\circ}, -60^{\circ}, 30^{\circ}, -30^{\circ}, 60^{\circ},$ 700 thereby representing pairs of orthogonal plies stacked at 701 a constant rotation angle of 30°, in approximation of the 702 living tissue.

In the rostral apex, the exocuticle is reduced to a thin 5 µm in thickness, or 10% of the total 705 cuticle thickness. The endocuticle displays a more com-706 plex pattern of layer thicknesses in the rostral apex than 707 in the head capsule. Each of the eight outermost lay-708 ers are of nearly equal thickness to the exocuticle (5 μm), 709 while the four innermost layers have a combined thick-710 ness equal to that of the exocuticle or a single layer of The fracture surfaces of cold-fractured specimens were $_{711}$ outer endocuticle ($h_{outer} = 5 \,\mu\text{m}, h_{inner} = 1.25 \,\mu\text{m}$). The examined using scanning electron microscopy to charac- 712 stacking sequence with respect to the longitudinal axis terize the composite profile and microstructure of the ros-

716 and stacking angle sequence to cuticle flexibility in the 750 described above, we calculate the extensional stiffness $_{717}$ rostral apex, two hypothetical cuticle lay-ups were also $_{751}$ matrix [A], bending stiffness matrix [D], and bending- $_{718}$ modeled, each with only one of the modifications present $_{752}$ extension coupling matrix [B] for each laminate consist-720 has the layer thicknesses of the rostral apex, but fiber 754 plane. The elements of these matrices can be found ac-721 orientations of the head capsule, while the second has 755 cording to: the fiber orientations of the rostral apex, but the layer thicknesses of the head capsule.

Because these laminates are not symmetric, there exists for each a bending-extension coupling matrix [B]populated with non-zero terms, thus complicating the calculation of effective in-plane elastic moduli. To circumvent this difficulty and enable meaningful comparison between each laminate, all of the layups are reflected 730 about their inner surface. This effectively doubles their thickness while producing a balanced, symmetric lami-732 nate with no coupling between bending and extension 733 (i.e., $[B] = 0_{3,3}$). Estimation of in-plane elastic constants from the extension([A]) and bending ([D]) matrices is de-735 scribed in detail below; Matlab R2018b was used to nu-736 merically evaluate the final values of the effective elastic 737

Classical Laminate Plate Theory We begin by calcu-738 lating the 2D reduced stiffness matrix for each part of the 740 cuticle. For orthotropic materials with the principal axes $_{741}$ parallel to the ply edges, the reduced stiffness matrix is 742 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: 743

$$Q_{11} = \frac{E_1}{1 - \nu_{12}\nu_{21}},$$

$$Q_{12} = \frac{E_1\nu_{21}}{1 - \nu_{12}\nu_{21}} = Q_{21},$$

$$Q_{21} = \frac{E_2\nu_{12}}{1 - \nu_{12}\nu_{21}} = Q_{12},$$

$$Q_{22} = \frac{E_2}{1 - \nu_{12}\nu_{21}},$$

$$Q_{66} = G_{12}.$$
(2)

For each layer k, the reduced stiffness matrix is transformed to account for the layer orientation angle θ within the laminate coordinate system, yielding a reduced transformed stiffness matrix according to:

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

where the transformation matrix [T] is defined as:

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

To assess the individual contributions of layer thickness 749 Using the lay-ups specified for each type of cuticle in the cuticle of the rostral apex. The first of these models $_{753}$ ing of n layers at a distance z from the laminate mid-

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

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

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

These stiffness matrices relate vectors of resultant 757 forces $\{N\}$ and bending moments $\{M\}$ to mid-surface 758 strains and curvatures $\{\epsilon^{\circ}\}$ and $\{\kappa\}$, respectively, in the 759 laminate according to the following relationship:

For symmetric laminates, $[B] = 0_{3,3}$, and therefore:

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

$$(7)$$

or, in expanded form:

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

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

If we make the simplifying assumptions (see ref.) that 764 (1) the laminate experiences pure axial loading and 765 transverse bending (i.e., $N_{yy}=N_{xy}=0$ and $M_{yy}=$ 766 $M_{xy}=0$, respectively) and (2) the laminate is a beam 767 of sufficiently high aspect ratio to minimize the Poisson 768 effect and anisotropic shear coupling (i.e., below we ef-769 fectively let $A_{12}^* = A_{16}^* = 0$ and $D_{12}^* = D_{16}^* = 0$), then 770 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 773 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:

776 substituting $A_{12}^*=A_{16}^*=0$ based on the assumptions 801 exploration and fitting was conducted in R version 777 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 779 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 781 laminate, we first specify the moment-curvature relation 782 of an Euler-Bernoulli beam:

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

Along the x-axis, the second moment of area for a rect-784 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-786 ment along the x-axis is related to the moment of the 830 this study. 787 beam by:

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

790 ticity of the laminate can be found by making Eq. 13 spe- 838 R_{σ}^2 , and $R_{\beta*}^2$ [R:'r2glmm':r2beta] (see Supplementary 791 cific to transverse flexure of the x-axis and rearranging 839 Data). R_{σ}^2 is the proportion of generalized variance ex-792 the terms:

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

794 to:

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

Model selection and fitting In order to explore the 797 relationships between the composite structure and me-798 chanical properties of the cuticle, we fit phylogenetic lin-799 ear mixed-effects models (PGLMM) to the tensile testand, by inverting this equation (let $A^* = A^{-1}$) and 800 ing data using maximum likelihood estimation. Model 802 3.5.1 (2018-07-02) -- 'Feather Spray' using the **nlme' and 'ape' packages. Response variables and 804 covariates were natural-log transformed, as needed, to (11) 805 ensure that the normalized model residuals were nor-806 mally distributed [R:shapiro.test] and homoscedastic [R:levene.test], using numerical and graphical analysis (see Supplementary Data). In order to control for phy-809 logenetic non-independence in the data, we included the 810 species of each specimen as a random effect in all models. 811 We also allowed for correlation in the error term of the 812 models, as specified by a variance-covariance matrix gen-813 erated from a Brownian motion [R: 'ape':corBrownian] 814 model of trait evolution along a phylogeny.

The preferred phylogeny for North American Curculio 816 is that of (REF-Bonal et al 2016); this phylogeny was 817 generated using Maximum-Likelihood methods and is un-818 trametric, but has uniform internal branch lengths. The 819 preferred tree was pruned to include only those species 820 used in tensile testing, with a polytomy at each species 821 to represent the individual specimens examined for that 822 species. Because the branch lengths of the tree were not specified, all branch lengths were set equal to 1. In gen-824 eral, phylogenetic regression is very robust with respect 825 to missing or incomplete branch length data; underesti-(14) 826 mation of branch length causes overestimation of phy-827 logenetic signal, which was not found to be statistically 828 significant for any of our models, and we therefore be-829 lieve that this procedure does not affect the findings of

Other models of trait evolution were considered dur-832 ing model exploration, including Ornstein-Uhlenbeck 833 [R:'ape':corMartins] and variable rate (AC/DC) (15) 834 [R: 'ape':corBlomberg] models. However, neither of 835 these produced a significantly better fit to the data Thus, given the assumption that $D_{12}^* = D_{16}^* = 0$, 836 than the Brownian motion model for any comparison, as Young's modulus for the effective transverse flexural elas- 837 measured by likelihood score and residual variance, viz. 840 plained by fixed effects, and is generally used to compare between covariance structures (REF). $R_{\beta_*}^2$ measures the 842 multivariate association between the outcome and the 843 fixed effects within a given correlation structure, and is generally used to compare between fixed effects (REF). 845 We additionally estimated the phylogenetic signal in the From inversion of Eq. 7 (let $D^* = D^{-1}$) this reduces 846 residual variance of each model using two measures of spatial autocorrelation (Abouheif's C and Moran's I) 848 and two measures of phylogenetic signal (Pagel's λ and 849 Blomberg's κ), as detailed below.

850 In all models, we tested whether the inclusion 851 of phylogenetic correlation in the model error pro852 duced significantly better model fit, using a likelihood- 890 For our second aim we examined the relationship beeach cofactor was significantly different from zero.

Hypothesis testing Our objectives for hypothesis test- 901 men stiffness as a fixed effect. ing using the PGLMM's were threefold: (1) assess whether the altered ratio of exocuticle to endocuticle in the rostral apex has an effect on the tensile strength of the snout; (2) test whether a trade-off exists between specimen stiffness and resistance to fracture; and (3) examine whether snout length and flexibility are correlated.

To test the relationship between relative layer thick-871 nesses and tensile strength, we fitted a fully-specified $_{872}$ model with the cross-sectional area of exocuticle and endocuticle at the site 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 compared to models with only crosssectional area of either endocuticle or exocuticle, but not both, as the sole fixed effect in the model. We then tested the whether one or both regions was significantly correlated to the maximum force sustained in tension by using likelihood-ratio tests and $R^2_{\beta*}$ -difference tests between each of the three models. We found that only the cross-sectional area of endocuticle was a significant fixed 915 889 strength as the response variable.

ratio test [R:'1mtest':1rtest] and R_{σ}^2 -difference so tween specimen stiffness and resistance to fracture. Spectest [R:'r2glmm':r2beta(method='sgv')] between the 892 imen stiffness was characterized using a low strain elasfully-specified model and a model lacking the phyloge- 893 tic modulus (averaged across the first 33% of the stressnetic effect. Models that incorporated covariance due to 894 strain curve) and the secant modulus at failure. Re-Brownian motion consistently produced a higher likeli- 895 sistance to fracture was quantified in terms of ultimate hood score and fit to the data; however, no model ex- 896 strain and fracture toughness (measured as area under hibited statistically significant phylogenetic signal in any 897 the stress-strain curve). Four models were fitted, two of of the variables. Once models were fitted to the data, 898 which had ultimate strain as the response variable and T and F-statistics were calculated to determine whether 899 two with fracture toughness as response variable. Each model then used one of the different measures of speci-

> Finally, in the third aim we explored whether a size-903 effect might exist in the Curculio rostrum, particularly 904 if longer (and typically more curved) rostra were more 905 flexible than shorter, straighter rostra. We generated two 906 models: the low strain and secant elastic moduli individ-907 ually served as response variables, with specimen length 908 as the fixed effect in both.

Code availability

R, Python, and Matlab scripts used to manipulate and 911 analyze the raw data (as well as their outputs), produce 912 figures, and estimate effective elastic constants is avail-913 able from the corresponding author upon request.

Data availability

Stress-strain curves for all tensile and fatigue-tested effect, and therefore we elected to remove cross-sectional 916 specimens are provided as PDFs (Supplement 1). Diagarea of exocuticle as a fixed effect to make the model 917 nostic plots for all PGLMMs are provided as PDFs (Sup-886 more parsimonious. The final model generated for our 918 plement 2). PGLMM terms and output are provided in 887 first aim featured the ratio of exocuticle to endocuticle 919 a PDF (Supplement 3). Raw and processed data will be in cross-section as a fixed effect, with ultimate tensile 920 provided by the corresponding author upon reasonable 921 request.

REFERENCES

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