# Microstructure and mechanical behavior of the acorn weevil exoskeleton

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We report novel modifications to the composite miscrostructure of the exoskeleton in the snout of acorn wees vils (Coleoptera: Curculionidae) belonging to the genus Curculio Linnaeus, 1756.

As a weevil (snout beetle), members of the genus Curculio are typified by the presence of a highly elongate structure on the head, called the rostrum (snout). This structure is a hollow, strongly curved (over 90°), cylindrical extension of the exoskeleton of the otherwise nearlyspherical head, which bears at its apex the terminal chewing mouthparts. The space inside of the rostrum contains the esophagus, various muscles and tendons used for feeding, and hemolymph that serves as a rough equivalent to blood in insects. By contrast, the solid shell of the rostrum is comprised entirely of cuticle, which can be considered a laminate composite consisting of various arrangements of chitin fibers embedded in a protein matrix (see section I). Acorn weevils use this structure to excavate sites for egg-laying (oviposition) and feeding on a variety of fruits, including acorns, Japanese camellia, hazelnuts, pecans, chestnuts, and chinquapins.

During oviposition, a female engages in a unique drilling" behavior that causes significant, apparently elastic, deformation of the rostrum. The female will insert the snout into an incision made with the mandibles, eating the material as she proceeds, while rotating her head and body around the perimeter of the bore-hole. Once the apex of the snout is fully inserted, she will push up and forward with her front legs, forcing the rostrum to bend until it is nearly straight. The female will maintain tension on the rostrum in this position, continuing to ingest the substrate and rotate around the bore-hole, while slowly inserting the rostrum further into the excavated channel. Once the rostrum is fully inserted, usually up to the eyes, she will pull her snout from the bore-42 hole and deposit several eggs into the site. By maintain-43 ing constant tension on the rostrum and rotating around 44 the bore-hole, the female is able to flex the snout into

<sup>45</sup> a near-perfectly straight configuration and thereby pro-<sup>46</sup> duce a linear channel into the fruit. While this behavior <sup>47</sup> has been observed in many species of *Curculio*, we have <sup>48</sup> lacked a fundamental understanding of how female *Cur*-<sup>49</sup> *culio* rostra can withstand the repeated, often extreme <sup>50</sup> bending incurred during the process of oviposition.

We have found that the composite profile of the rostrum is strongly differentiated from the head capsule and
other body parts, with modification of both the relative
layer thicknesses and fiber orientation angles of cuticle
regions (viz. exocuticle and endocuticle), which we describe in detail below. We posit that these modifications
enable the snout to be flexed until straight while remaining within the elastic limits of the material, mitigating
the risk of structural damage, and without evident alteration of the mechanical properties of the individual components of the cuticle across the structure and between
species. Thus, the flexibility and tensile strength of the
rostrum appear to be derived exclusively from modification of the composite architecture of the exoskeleton.

Support for this hypothesis has come from three lines of evidence:

- 1. Examination of the cuticle microstructure across the length of the snout has revealed consistent modification to the composite structure of the rostrum among *Curculio* species.
- 2. Tensile testing of the rostrum has demonstrated that the mechanical strength of the cuticle components are consistent along the length of the structure and between species.
- 3. Fatigue testing has shown that a highly curved rostrum is capable of flexing hundreds of thousands of times without damage to the structure, and is apparently elastic.

We additionally describe the fracture mechanics of the so snout, as pertains to both cuticle composite structure and tensile behavior, and consider how modification of the cuticle may reduce the risk of rostral fracture dursing oviposition. To our knowledge, this is the first time

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84 that a modified composite profile has been reported as 139 REFS-HERE), while the stacking angle between pairs is

low beginning with a consideration of the impact of mi- 144 between successive plies [10–12]. 90 crostructure, then we talk about mechanical testing, blah 145 92 derstanding the mechanical behavior of the snout during 147 profile of endocuticle, with an angle of approximately 30° bending.

## MICROSTRUCTURE OF THE ROSTRUM

Insect cuticle is made up of chitin and numerous un-96 characterized proteins, which effectively act as a matrix into which the chitin is embedded [3–5]. The arrangement of the embedded chitin fibers is primarily responsible for the mechanical behavior of the cuticle, although this is modulated by the degree of tanning and water retention. [1, 3]. Accordingly, cuticle is often considered as a fiber-reinforced composite material; however, in beetles, the composite arrangement of the chitin fibers varies by cuticle region.

Insect cuticle is divisible into  $_{\mathrm{three}}$ regions (Figs. INSERT-REFS-HERE), including (1) endocuticle, which is the most compliant and innermost region; and (3) exocuticle, which is the stiffest, hardest, and outermost region. Between these lies (2) mesocu- 165 transition zone [1–3].

115 always evident, and is sometimes considered part of the 171 increasing the flexibility of the resulting composite. Adtrix. These layers are stacked at a more or less constant 176 pletely straight without fracture. angle to each other in a helicoidal arrangement, referred 177 strong anisotropy of  $\alpha$ -chitin.

127 take advantage of this anisotropy to improve fracture re- 183 [? ? ]. This composite profile has now been uncovered OD) unidirectional bundles of chitin, called macrofibers. 186 a genus-wide trait. 134 are aligned in layers, as in Figs INSERT-FIGS-HERE 190 no difference in total cuticle thickness between the head 136 of macrofibers, with the adjacent layers forming pairs. 192 thicknesses. Below we demonstrate that that the endocu-137 The layers within each pair are pseudo-orthogonal (i.e., 193 ticle does not vary in tensile strength across the rostrum,

a means of enhancing structural elasticity in the insect 140 typically acute, although other configurations have been observed [10, 11]. It is thought that this arrangement of MIGHT BE GOOD TO SIGNPOST HERE... we dis- 142 the macrofibers contributes to the toughness of beetle excuss each of these lines of evidence in the sections be- 143 oskeletons by inhibiting crack formation and propagation

We have found that in *Curculio*, the head capsule blah. Emphasize that this is key to predicting and un- 146 (which is similar to the rest of the body) fits the general 148 between successive pseudo-orthogonal plies. Addition-149 ally, in the head capsule, the thickness of the exocuticle 150 in cross section is nearly equal to that of the endocuti-151 cle; compared to the snout, this part of the head is fairly

> By contrast, the region beyond the scrobe (antennal channel) is quite flexible, even in fully desiccated specimens. Serial thin sectioning of the snout has demon-156 strated that the cuticle in this region has a different com-157 posite structure (see Fig INSERT-REFS-HERE) than 158 the head capsule, namely:

- 1. The exocuticle is reduced to a thin shell, with the endocuticle thickened to offset this reduction.
- 2. The endocuticular macrofibers exhibit no rotation between successive pseudo-orthogonal plies, which are all oriented at approx.  $\pm 45^{\circ}$  to the longitudinal axis of the snout.

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We believe that each of these modifications contributes ticle, which is similar in microstructure to exocuticle, 166 to the observed flexibility of the snout, but in differbut less sclerotized (tanned), usually acting as a thin 167 ent ways. Because the exocuticle is generally more rigid 168 (and brittle) than the endocuticle due to its transverse In general, these regions are confluent, and not nec- 169 isotropy and microstructure, a reduction in the relative essarily sharply defined; for instance, mesocuticle is not 170 thickness of this region would likely have the effect of exocuticle. In beetles, both exocuticle and mesocuticle 172 ditionally, aligning the layers of endocuticle as a 45 deg (when present) are laminate, and have numerous lam- 173 cross ply would likely reduce the bending moment of the inae of unidirectional chitin fibers, each layer a single 174 structure. The combined effect of these modifications is fiber thick (2-4nm), embedded in a proteinaceous ma- 175 that the apex of the rostrum is able to bend until com-

Correspondingly, the portion of the snout between the to as the Bouligand structure [6–8]. This microstructure 178 head capsule and apex of the scrobe exhibits a gradual allows exocuticle and mesocuticle to exhibit transverse 179 transition in composite profile along an anterior-posterior isotropy, or complete isotropy in some cases, despite the 180 gradient. In previous work we identified these modifica-181 tions to the composite structure of the cuticle, but only Conversely, the endocuticle of Coleoptera appears to 182 within a single species, C. longinasus Chittenden, 1927 sistance of the exoskeleton. Beetles in particular have a 184 in the rostral apex of six additional, phylogenetically dishighly modified endocuticle comprised of large (5-20 µm 185 parate, species (listed below), indicating that this is likely

The macrofibers are strongly anisotropic, with an elastic 187 Notably, we could find no evidence of resilin, as indimodulus of 8-9GPa along their longitudinal axis, but 188 cated by florescent microscopy, anywhere in the cuticle only 1-2 GPa along the transverse plane. Macrofibers 189 of the head (including the rostrum). We also observed [11, 12]. Typically, endocuticle contains several layers 191 capsule and rostral apex, only differences in relative layer 138 angled approx. 90° to each other, see Figs. INSERT- 194 making it unlikely that differences in sclerotization or its cuticle.

#### TENSILE TESTING OF THE ROSTRUM II.

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To better characterize the mechanical behavior of the 201 202 snout as an integrated whole, we performed tensile testing on the snouts of six species in the genus Curculio, representing a mixture of closely and distantly related taxa. Although the heads were rehydrated by immersion in de-ionized water for 24 hours, we observed comparatively brittle fracture (but see below), in contrast to GIVE EXAMPLES HERE. The tensile behavior for the cuticle of these weevils is a result of its microstructure. which lacks pores, etc (give reasons). THIS IS ALSO 260 A GOOD SPOT TO SIGNPOST WHAT I FOUND IN 212 GENERAL.

#### Tensile behavior

We observed that the maximum force sustained at the 266 (name some). 215 site of the break was strongly correlated with the cross- 267 sectional area of the endocuticle, and not the exocuticle. 268 a smooth, nearly continuous fracture surface, indicative fracture, indicating that as the proportion of exocuticle 272 hand exhibited severe delamination, ply-splitting, and 221 increased, tensile strength decreased. In other words, 273 fiber pulling, consistent with viscoelastic/plastic behavof the endocuticle across species. These associations were 275 ical consideration of the microstructure of this material 224 found to be statistically significant and independent of 276 (name something here). These patterns indicate that the 225 species membership, rostrum length, and location on the 277 endocuticle is probably less brittle than the exocuticle, snout. 226

These data have three important implications:

- 1. There is very little variation in the gross elastic behavior of the cuticle across the genus, in agreement with our current understanding of cuticle mechanobiology. (singh 2016, jansen 2016)
- 2. The endocuticle contributes more to the tensile strength of the rostrum than the exocuticle, possibly because the endocuticle is organized in large bundles of aligned, anisotropic fibers.
- 3. Thus, in addition to making the cuticle more flexible, the altered apical composite profile makes it less likely to break, suggesting a possible means by breakage, maybe say this in conclusions).

242 in specimens with a longer snout. This observation was 297 243 initially quite puzzling, as we had expected that a longer 298 tic failure mode, based on the pattern of fiber dislocation

195 chitin composition within the cuticle are responsible for 244 (and typically more strongly curved [insert ecomorph pathe mechanical behavior of the rostrum. The available 245 per ref here]) would need to be more flexible to avoid evidence therefore suggests that the relative flexibility of 246 fracture during oviposition. Based on preliminary confothe snout is solely derived from the composite profile of 247 cal microscopy data, we speculate that this phenomenon 248 may be the result of a longer scrobe/transition/gradient <sup>249</sup> from the basal profile to the apical profile in longer rostra. 250 A decrease in exocuticle thickness along a longer portion <sub>251</sub> of the base might reinforce the snout against buckling; 252 however Young's modulus of the rostrum would be com-<sup>253</sup> paratively higher in these species because of the greater volume fraction of exocuticle. [OR SAY MORE CON-CISELY THAT: based on prelim CLSM data we believe 256 that the snout is reinforced against bucking with thicker 257 exocuticle in the base of these species; as a result, the 258 higher volume fraction of exocuticle in the snout increases 259 young's modulus for such species

#### В. Fractography

Examination of the fracture surfaces and adjacent cu-262 ticle of tensile testing specimens revealed that although fracture was comparatively brittle, the fracture mechan-264 ics of the exocuticle and endocuticle differed according to 265 their microstructure, in agreement with previous studies

In cross-section, the exocuticle consistently presented Consequently, there was a negative correlation between 269 of relatively brittle fracture, likely due to the microarutlimate tensile strenth of the specimen and the ratio of 270 chitecture and resultant transverse isotropy of the Bouliexocuticle to endocuticle cross-sectional area at the site of 271 gand structure (explain this). Endocuticle, on the other UTS is strongly correlated with the cross-sectional area 274 ior shown in previous studies and congruent with theoret-278 most likely due to the alignment of the  $\alpha$ -chitin fibers in 279 each macrofiber.

In addition, the exocuticle typically appears to fracture before the endocuticle, with shear-cusp formation 282 evident over unbroken endocuticle. We note, however, 283 that the exocuticle of weevils/beetles is anchored to the 284 endocuticle by cross-linking fibers in a transition zone de-285 scribed by Kamp et al. (see refs.). The presence of shearcusps therefore indicates that the fibers of the endocuti-287 cle are shearing past each other within each ply along 288 the radial/normal plane (type II shearing???). Further-289 more, given the delamination observed between plies, the 290 layers of endocuticle are liekly shearing past each other (type III shearing) along the transverse plane. We therefore infer that, under tension, the endocuticle tends to which the system evolved (via negative selection of 293 deform visco-elastically and plastically along the longi-294 tudinal axes of the macrofibers, while the overlying exo-<sup>295</sup> cuticle exhibits brittle fracture due to shearing between Finally, we observed that the elastic modulus is higher 296 the stretching endocuticle fibers to which it is anchored.

Additionally, the fracture surfaces show a characteris-

299 in the plies of the endocuticle. (copy description from 335 other members of its species. Given this surprising result, DATA PRESENTATION, for figure 5B).

302 rich gular sutures are the most likely site for the initiation 338 in life, repeated bending of the snout does exceed the 303 of void nucleation and failure of the integrated rostral cu- 339 yield/plastic limit of the cuticle, and the bending strain ticle in cross section. Structural failure would take place 340 is purely elastic/visco-elastic. 305 as cracks propagate through the endocuticle from these 341 306 sutures, which penetrate the entire thickness of the lami- 342 ulate that the microstructure of the endocuticle is recross-sectional profile of *C. caryae* is flattened ventrally Flattening this region may reduce tensile-strain across the gular sutures when the snout is bent dorsally, thus 311 reducing the risk of fracture in the elongate, strongly-312 curved rostrum in this species.

# FATIGUE TESTING OF CURCULIO CARYAE

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315 longinasus, which exhibits the most extreme degree of 359 any damage to the specimen. bending in the sis species examined.

400K cycles, complete elongation, rehydrated, coated 321 322 in grease to prevent loss of moisture and stiffening of the specimen. We observed visco-elastic behavior in the specimen, as indicated by histeresis in the stress-strain 361 initially believed that this indicated damage to the specimen; however, after cleaning the specimen in a 24 hour wash with ethanol and water, we observed that the specimen returned to its original shape. The specimen did 332 not show any evidence of fractures or shear cusps any- 366 334 the tensile strength of the specimen was consistent with 368 able in the online version of this paper.

336 it appears that the specimen was undamaged by the test-From this pattern we hypothesize that the exocuticle- 337 ing. We therefore believe that under normal conditions

We cannot fully account for these results, but we specnate. We speculate that this could be the reason why the 343 sposible for what we observed. The endocuticle is made  $_{344}$  of aligned  $\alpha$ -chitin nanofibrils whose crystaline structure is enforced by hydrogen bonds between individual chitin 346 chains along their length. The viscoelasticity of the cu-347 ticle is thought, in part, to come from slippage between 348 these chains as the hydrogen bonds break and reform in 349 response to shearing between the chitin molecules. We 350 believe that repeated strain may have caused such slip-351 page in the endocuticle of the rostral apex during the 352 fatigue test, but without sufficient time for the mate-353 rial to completely relax after deformation, the specimen would slowly accumulate strain and consequently deform The shearing motion and accommodation of strain led 355 visco-elastically/visco-plastically. After 24 hours soaking us to question how the cuticle might accommodate re- 356 in ethanol and water, the hydrogen bonds would relax peated strain, as is seen in the living organism. We 357 sufficiently to allow the specimen to return to its original therefore performed fatigue testing on a female curculio 358 configuration, dissipating the accumulated strain without

#### CONCLUSIONS

Overall conclusion - the only difference between the relationship during each cycle. Fmax decreased logarith- 362 flexible part and the stiff part is the microstructure; there mically with cycle number, etc., and the specimen ap- 363 is no difference in the mechanical behavior of the endocupeared to have deformed plastically during the test. We 364 ticle at different locations on the snout and across species.

#### $\mathbf{V}$ . METHODS

Methods, including statements of data availability and 333 where in the surface of the exocuticle, and, furthermore, 367 any associated accession codes and references, are avail-

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### **AUTHOR CONTRIBUTIONS**

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

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

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

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

# ADDITIONAL INFORMATION

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

# COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

#### **METHODS**

## Histological sectioning

# Tensile and fatigue testing

## Specimen imaging and microscopy

### Statistical analysis

General approach To explore the relationships be-435 tween the composite structure and mechanical proper-436 ties of the cuticle, we fit phylogenetic linear mixed-effects 437 models to the data using maximum likelihood estimation. 438 In order to control for phylogenetic non-independence in 439 the data, we included the species of each specimen as a random effect in all models. We also allowed for corre-441 lation in the error term of the models, as specified by 442 a variance-covariance matrix generated from a Brown-443 ian motion model of trait evolution along the phylogeny. 444 Response variables and covariates were natural-log trans-445 formed, as needed, to ensure model residuals were nor-446 mally distributed and homoscedastic. In all models, we tested whether the inclusion of phylogenetic correlation in the model error produced significantly better model 449 fit, using a likelihood-ratio test and  $R_{\sigma}^2$ -difference test 450 between the fully-specified model and a model lacking 451 the phylogenetic effect.

Hypothesis testing The following three hypotheses 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-sectional area of endocuticle and exocuticle 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 cross-sectional area of either model. We then tested the first hypothesis by using likelihood-ratio tests and  $R_{\beta*}^2$ -difference tests between each of the three models.

The hypothesis that

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- 473 Model selection and fitting
- Estimating phylogenetic signal

475	Code availability
476	Data availability

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