Microstructure-derived strength in 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 13 structure on the head, called the rostrum (snout). This 14 structure is a hollow, strongly curved (over 90°), cylindri-15 cal 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 26 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 41 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

45 a near-perfectly straight configuration and thereby pro-46 duce a linear channel into the fruit. While this behavior 47 has been observed in many species of *Curculio*, we have 48 lacked a fundamental understanding of how female *Cur-*49 *culio* rostra can withstand the repeated, often extreme 50 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 this arrangement of the macrofibers contributes to the

MIGHT BE GOOD TO SIGNPOST HERE... we dis- 142 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 intothree regions 106 (Figs. INSERT-REFS-HERE), including (1) endo- 162 transition zone [1–3].

115 always evident, and is sometimes considered part of the 171 a genus-wide trait. exocuticle. In beetles, both exocuticle and mesocuticle 172 strong anisotropy of α -chitin.

127 take advantage of this anisotropy to improve fracture re- 183 the snout is solely derived from the composite profile of sistance of the exoskeleton. Beetles in particular have a 184 its cuticle. highly modified endocuticle comprised of large $(5-20\mu m)$ OD) unidirectional bundles of chitin, called macrofibers. Macrofibers are aligned in layers, as in Figs INSERT-FIGS-HERE [11, 12]. Typically, endocuticle contains 133 several layers of macrofibers, with the adjacent layers forming pairs. The layers within each pair are pseudo- 187 135 orthogonal (i.e., angled approx. 90° to each other, see 188 the cuticle, it is possible to make several predictions 136 Figs. INSERT-REFS-HERE), while the stacking angle 189 about the mechanical behavior of the snout. First, we 137 between pairs is typically acute, although other configu- 190 can predict that the endocuticle will contribute more to 138 rations have been observed [10, 11]. It is thought that 191 the tensile strength of the rostrum than the exocuticle,

a means of enhancing structural elasticity in the insect 140 toughness of beetle exoskeletons by inhibiting crack formation and propagation between successive plies [10–12].

We have found that in Curculio, the head capsule cuss each of these lines of evidence in the sections be- 143 (which is similar to the rest of the body) fits the general low beginning with a consideration of the impact of mi- 144 profile of endocuticle, with an angle of approximately 30° of crostructure, then we talk about mechanical testing, blah 145 between successive pseudo-orthogonal plies. Additionblah. Emphasize that this is key to predicting and un- 146 ally, in the head capsule, the thickness of the exocuticle 92 derstanding the mechanical behavior of the snout during 147 in cross section is nearly equal to that of the endocuti-148 cle; compared to the snout, this part of the head is fairly 149 rigid.

> By contrast, the region beyond the scrobe (antennal 151 channel) is quite flexible, even in fully desiccated spec-152 imens. Serial thin sectioning of the snout has demon-153 strated that the cuticle in this region has a different composite structure (see Fig INSERT-REFS-HERE) than 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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Correspondingly, the portion of the snout between the cuticle, which is the most compliant and innermost 163 head capsule and apex of the scrobe exhibits a gradual region; and (3) exocuticle, which is the stiffest, hardest, 164 transition in composite profile along an anterior-posterior and outermost region. Between these lies (2) mesocu- 165 gradient. In previous work we identified these modificaticle, which is similar in microstructure to exocuticle, 166 tions to the composite structure of the cuticle, but only but less sclerotized (tanned), usually acting as a thin 167 within a single species, C. longinasus Chittenden, 1927 168 [? ?]. This composite profile has now been uncovered In general, these regions are confluent, and not nec- 169 in the rostral apex of six additional, phylogenetically disessarily sharply defined; for instance, mesocuticle is not 170 parate, species (listed below), indicating that this is likely

Notably, we could find no evidence of resilin, as indi-(when present) are laminate, and have numerous lam- 173 cated by florescent microscopy, anywhere in the cuticle inae of unidirectional chitin fibers, each layer a single 174 of the head (including the rostrum). We also observed fiber thick (2-4nm), embedded in a proteinaceous ma- 175 no difference in total cuticle thickness between the head trix. These layers are stacked at a more or less constant 176 capsule and rostral apex, only differences in relative layer angle to each other in a helicoidal arrangement, referred 177 thicknesses. Below we demonstrate that that the endocuto as the Bouligand structure [6–8]. This microstructure 178 ticle does not vary in tensile strength across the rostrum, allows exocuticle and mesocuticle to exhibit transverse 179 making it unlikely that differences in sclerotization or isotropy, or complete isotropy in some cases, despite the 100 chitin composition within the cuticle are responsible for 181 the mechanical behavior of the rostrum. The available Conversely, the endocuticle of Coleoptera appears to 182 evidence therefore suggests that the relative flexibility of

TENSILE TESTING AND FRACTURE **MECHANICS**

Based on our understanding of the microstructure of

192 because the endocuticle is organized in large bundles of 201
193 aligned, anisotropic fibers. Second The predicted behav194 ior of the snout is borne out by the data, where UTS is
195 strongly correlated with the cross-sectional area of the 203
196 endocuticle across species. In addition to making the
197 cuticle flexible, this profile makes it less likely to break, 204
198 suggesting a possible means by which the system evolved
199 (negative selection of breakage, maybe say this in con200 clusions).

III. FATIGUE TESTING OF CURCULIO CARYAE

IV. CONCLUSION

V. METHODS

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

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AUTHOR CONTRIBUTIONS

- Andrew Jansen: Conducted sectioning and staining,
 microscopy and imaging, tensile and fatigue
 testing, statistical analysis, and participated in
 manuscript preparation.
- Jason Williams: Conducted tensile and fatigue testing,
 participated in manuscript preparation.
- Nikhilesh Chawla: Facilitated microscopy, tensile and fatigue testing, and participated in manuscript preparation.
- Nico Franz: Facilitated specimen acquisition and imaging, participated in manuscript preparation.

ADDITIONAL INFORMATION

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

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

METHODS

Histological sectioning

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Tensile and fatigue testing

Specimen imaging and microscopy

Statistical analysis

General approach To explore the relationships between the composite structure and mechanical properties of the cuticle, we fit phylogenetic linear mixed-effects models to the data using maximum likelihood estimation. In order to control for phylogenetic non-independence in the data, we included the species of each specimen as a random effect in all models. We also allowed for correlation in the error term of the models, as specified by variance-covariance matrix generated from a Brownian motion model of trait evolution along the phylogeny. Response variables and covariates were natural-log transformed, as needed, to ensure model residuals were normally distributed and homoscedastic. In all models, we tested whether the inclusion of phylogenetic correlation ²⁸⁷ in the model error produced significantly better model 288 fit, using a likelihood-ratio test and R_{σ}^2 -difference test $_{289}$ between the fully-specified model and a model lacking 290 the phylogenetic effect.

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

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- 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-302 sectional area of endocuticle and exocuticle at the site 303 of fracture as fixed effects, including an interaction term, 304 and with maximum tensile force sustained prior to frac-305 ture as a response variable. This model was then com-306 pared to models with only cross-sectional area of either 307 endocuticle or exocuticle as the sole fixed effect in the 308 model. We then tested the first hypothesis by using 309 likelihood-ratio tests and $R_{\beta*}^2$ -difference tests between 310 each of the three models.

The hypothesis that

Model selection and fitting

Estimating phylogenetic signal

Code availability

Data availability

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