A new, giant ricinuleid (Arachnida, Ricinulei), from the Pennsylvanian of Illinois, and the identification of a new, ontogenetically stable, diagnostic character

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Abstract.—A new fossil ricinuleid, Curculioideal bohemondi n. sp., from the Pennsylvanian Energy Shale of Illinois is described from a single specimen. It is the largest ricinuleid species yet described, living or extinct. The Energy Shale represents a new geographic locale for fossil ricinuleids, a sparsely distributed group. The species is distinguished from other members within the genus by the possession of very large (0.09 mm) carapace tubercles at a very low (30 mm⁻²) density. Statistical analyses are performed on extant and fossil ricinuleids to determine how their tubercles change throughout ontogeny, culminating in the recovery of a new ontogenetically stable diagnostic character: the tubercle coefficient (a measure of the size of the tubercles relative to body size).

UUID: http://zoobank.org/aa9f2de5-c49d-4f70-bba5-db12fdee406f.

Introduction

The order Ricinulei is an enigmatic group of arachnids indigenous in modern times to West Africa, South America, and Central America (Selden, 1992). The order was originally described from fossil material found in England by William Buckland, who misinterpreted the specimen as a curculionid beetle (Buckland, 1837). A year later, the first living specimen was discovered, and classified as an opilionid (Guérin-Méneville, 1838a, b). From the outset, this peculiar group of arachnids was little understood due to their limited ranges, rarity, and secretive nature (Finnegan, 1935; Selden, 1992). It was not until 1904 that ricinuleids were separated from Opiliones and elevated to order Ricinulei Thorell, 1876 (Hansen and Sørensen, 1904). To this day they remain understudied, with much of their natural history unknown.

The position of Ricinulei within Arachnida has been a subject of debate (Garwood and Dunlop, 2014). Due to the suite of conflicting characters possessed by the order, several contradictory relationships to different arachnid groups have been proposed. Early classification schemes suggested a close relationship to the mites (Acari) primarily on the basis that both orders exhibit six-legged larvae that develop into eight-legged adults (Shultz, 2007). The locking mechanism that joins the carapace and opisthosoma and the divided opisthosomal tergites suggest their classification as sister to the order Trigonotarbida, an extinct group of arachnids that are among the earliest terrestrial organisms (Dunlop, 2010). Most recently, genetic evidence suggests ricinuleids may be most closely related to the order Xiphosura (horseshoe crabs) (Ballesteros and Sharma, 2019).

Extant ricinuleids are small, blind predators that inhabit caves and the litter of rainforest floors, seeking prey with their elongate sensory second leg pair (Adis et al., 1989). A paucity of observations has left many questions regarding their ecological niche, including their diet in the wild and their population structure (Platnick and Pass, 1982). Ricinuleids are often found in large congregations, though the behavioral purpose of this grouping behavior is speculative and remains untested (Mitchell, 1969). Despite their relatively low diversity (extant ricinuleids form a single family with only three genera), ricinuleids occupy a surprisingly large number of diverse niches (Selden, 1992). Extant species occupy one of only two major ecotypes (soil dwellers or troglodytic). Nonetheless, they exhibit diverse adaptations to these niches, from the modified setae of Cryptocellus adisi Platnick, 1988, which create an air pocket around themselves to survive periodic inundations of the soil, to the elongate legs of the troglodytic Pseudocellus krejcae Cokendolpher and Enríquez, 2004, which allow it to walk over the surface of subterranean pools (Adis et al., 1999; Cokendolpher and Enríquez, 2004). Hence, despite being relatively low in taxonomic diversity, ricinuleids nonetheless represent a myriad of interesting adaptations and ecologies that demand further study.

In contrast to the low taxonomic diversity of extant ricinuleids, the fossil record presents a veritable cornucopia of ricinuleid biodiversity. Four families and six genera contain 21 identified fossil species, yet the evolutionary history of the group remains ambiguous due to large spatial and temporal gaps in the ricinuleid fossil record (Selden, 1992; Wunderlich, 2015). Fossil ricinuleids are known only from the Pennsylvanian
and the Cretaceous, and from only a few localities in Myanmar, England, Germany, and the United States (Selden, 1992; Wunderlich, 2017). The mid-Cretaceous (99 Ma) specimens (two genera, five species) come from Burmese amber (Shi et al., 2012; Selden and Ren, 2017). The majority of fossil ricinuleids, however, originate from the Atokan and Desmoinesian (Moscovian) stages of the Pennsylvanian (ca. 308–315 Ma) (Selden, 1992; Wagner and Winkler Prins, 2016). The North American specimens occur primarily in the Pennsylvanian Fossil-Lagerstätte of Mazon Creek, Grundy Co., Illinois, apart from a single specimen from Morris, Oklahoma (Selden, 1992). The Mazon Creek fossils occur in the Francis Creek Shale, which is part of the Carbondale Formation (Willman et al., 1975). The Francis Creek Shale directly overlies the Colchester No. 2 Coal; it is composed of shales and fluvial sandstones (BaIRD, 1979).

Here, we describe a new fossil ricinuleid from the Pennsylvanian of Illinois (Fig. 1), the first Pennsylvanian ricinuleid described since 1992 (Selden, 1992). The fossil comes from the Energy Shale in Vermilion County, a new stratigraphic and geographic locale for fossil ricinuleids. The fossil is referred to a new species in the genus *Curculioidea* Buckland, 1837, *C. bohemondi* n. sp. The specimen is notable for its numerous, large, well-preserved tubercles. At 21.77 mm body length, it is the largest ricinuleid known, living or extinct. We also take the opportunity to illustrate the Oklahoma ricinuleid, *Amarixys gracilis* (Petrunkevitch, 1945), mentioned by Selden (1992) but not previously figured (Fig. 2).

Ricinuleid ontogeny consists of five post-egg life stages: larva, protonymph, deutonymph, tritonymph, and adult (Pittard and Mitchell, 1972). Determining the life stage of fossil specimens has often proven problematic. Definitive ontogenetic conclusions cannot generally be made because the majority of ricinuleid fossils fail to preserve the most-recognizable defining feature of each life stage—the number of terminal tarsomeres on each leg (Pittard and Mitchell, 1972). With the aim of identifying a potential life stage that the specimen could represent, we examine ontogenetic trends in extant ricinuleids, with a primary focus on a character often found in fossil specimens: the tubercles. Tubercles are raised, roughly hemispherical bumps covering the cuticle surface of both extinct and extant ricinuleids. The utility of tubercles in diagnosing adult members of ricinuleid species has previously been documented, but the changes these structures may undergo across life stages have not been explored (Selden, 1992). To determine if these potential changes correlate with life stage, tubercles were examined throughout the ontogeny of representative species from all three modern ricinuleid genera.

### Geological setting

The Kewanee Group lies in the middle of the Pennsylvanian strata of Illinois, situated between the underlying McCormick Group and the overlying McLeansboro Group (Willman et al., 1975). The Kewanee Group contains the highest concentration of coals of the three units and exhibits the first appearance of widespread, uniformly thick limestones in the Illinois Pennsylvanian (Kosanke et al., 1960). The ricinuleid fossil was discovered in the Carbondale Formation of the Kewanee Group, a unit notable for including among its most distinguished members the Francis Creek Shale: stratigraphic home to the Mazon Creek biota, which has previously produced a number of ricinuleid specimens (Willman et al., 1975; Selden, 1992).

In Vermilion County, where the new specimen was discovered, the Carbondale Formation averages 200 ft. thick and is composed primarily of gray shales with associated sandstone channels, thin argillaceous limestones, and numerous coals (Jacobson et al., 1980). The fossil was found in the Energy Shale Member of the Carbondale Formation, a gray siltstone that can reach up to 100 ft. in thickness (Jacobson et al., 1980). The Energy Shale is located between the underlying Herrin No. 6 Coal and the overlying Anna Shale and exhibits a noteworthy drop of 3–5% sulfur content from the neighboring rocks (Jacobson et al., 1980). Sulfurs in coals are thought to originate from exposure to seawater, hence this low-sulfur content suggests that the Energy Shale represents a freshwater system (Neilson, 1983).

The Energy Shale is primarily interpreted as a series of crevasse-splay deposits; the source river is thought to be preserved in the form of the Walshville Channel, an enormous (one mile wide, 60–80 ft. thick) channel sandstone cutting through the upper part of the Herrin No. 6 Coal and the Energy Shale (Burk et al., 1987; Hatch and Affolter, 2002). The deposition of these fluvial features is considered to have been contemporaneous with the deposition of peat, which became the Herrin No. 6 Coal (Treworgy and Jacobson, 1985). The Energy Shale dates to the Desmoinesian Stage (ca. upper Moscovian), fixing it firmly in the middle of the Pennsylvanian, 308–306 Ma (Willman et al., 1975). The paleoclimate would have been hotter and drier than the preceding stages of the Pennsylvanian, but still wet enough to form swamps during frequent humid, rain-heavy periods (Cecil, 1990).

The basal 5 m of the Energy Shale contain numerous siderite concretions that preserve various plants as well as terrestrial and marine fauna (Burk et al., 1987; Baird et al., 1985). Concretions occur in a faintly laminar gray, silty mudstone and contain diverse organisms, such as branchiopods, freshwater shark egg capsules, syncarid shrimps, *Euproops*, aquatic bivalves, and, on rare occasions, flying insects (Baird et al., 1985). Later transgression of the sea from the southwest flooded the Walshville Channel river system, resulting in the overlying brackish–marine Anna Shale and Bereton Limestone members (Treworgy and Jacobson, 1985; Burk et al., 1987).

### Morphological interpretation

Both part and counterpart (Fig. 1) represent the dorsal surface, with the part representing an external mold and the counterpart preserving a combination of internal and external molds. The ventral surface is not preserved. Tubercles cover the entire body and are unimodal in size. Tubercles lack any sort of surface sculpturing or ornamentation; if such fine features were originally present, they were likely lost during fossilization. The rounded apex and overall circular morphology of the well-defined opisthosoma tubercles suggest they most closely resembled the small dome-like tubercles of extant ricinuleids; conversely, the shallow relief exhibited by the less well-defined
carapace tubercles suggests a more flattened apex typical of iridescent tubercles (Salvatierra and Tourinho, 2016; Fig. 3). Due to the lack of any sculpted surfaces on these tubercles however, such identifications remain tenuous. At 113% the diameter and 60% the density of the next largest and sparsely distributed ricinuleid tubercles (Fig. 4), the tubercles of this specimen are greater in size and distributed more sparsely than those of any other known fossil species (Selden, 1992).

The cucullus is partially preserved as a scrap anterior to the carapace on both the part and the counterpart. Some weak tuberculation is present on the part, however much is obscured by heavy mineralization, including its margin with the carapace. No tuberculation is visible on the counterpart. It appears to have been preserved in an open position based on its significant anterior projection. A small depression is visible running longitudinally near the lateral edge of the cucullus in the part, similar

Figure 1.  *Curculioidea bohemondi* n. sp., holotype and only known specimen, ACCN2015-104, Pennsylvanian (Desmoinesian) Energy Shale Member, Carbondale Formation, Kewanee Group, near Little Vermilion River, Georgetown, Vermilion County, Illinois. (1) Photograph of part; (2) explanatory drawing of part; (3) photograph of counterpart; (4) explanatory drawing of counterpart. I–IV = walking legs 1–4; cuc = cucullus; fe = femur; m line = median line; mg = medial groove; mt = metatarsus; pa = patella; pm = plant material; pyg = pygidium; ti = tibia; tr, trochanter. Gray areas represent kaolinite; other thin lines are fractures. Scale bars=1 mm.
to the observed longitudinal grooves in the cucullus of *Curculioides mcluckiei* Selden, 1992.

The carapace exhibits a straight anterior margin with the cucullus. It then broadens posteriorly until its termination with the gently procurved posterior margin. A single median longitudinal groove runs down the midline of the carapace, visible most clearly as an external mold on the part; the counterpart has fractured along this groove. No change in tubercle distribution is evident nearer to the groove. It extends about two-thirds the length of the carapace before fading out. The median groove divides this anterior region of the carapace into two raised areas, forming a pair of raised lobes most visible as an external mold on the part. The median groove runs directly into the topographically lower posterior region of the carapace, forming a triangular spandrel-like architecture between the two lobes. Although this precise morphological arrangement of raised areas is unique, raised carapace areas have previously been observed in the Pennsylvanian ricinuleid *Amarixys stellaris* Selden, 1992. No posterior depressions (or the characteristic smaller tubercles normally contained therein) are evident. Extensive tuberculation is present across the carapace on both the part and the counterpart. A crack running latitudinally through the carapace on both the part and counterpart, and the glue used to repair it, partly obscures the area where eyes likely may have been originally present. Because no evidence of eyes can be observed on the margins of the glued area, it appears they may have been reduced in size, if indeed they were present at all. With such a large area obscured, no definite conclusion can be made.

The opisthosoma is better preserved on the part, but also possesses areas of mineralization and fracturing. Its margin with the carapace is defined by an untuberculated groove, which is visible as an external mold on the part; the counterpart has fractured along this groove. The line appears to have been somewhat warped by taphonomic processes and is only well defined close to the carapace and near the distal end of the body; it can be seen most clearly on the counterpart. Tubercles appear almost ordered on the opisthosoma, substituting the random distribution found across the rest of the body for a series of curved rows and columns, mirroring the curve of the carapace-opisthosoma boundary. Several particularly well-preserved tubercles are
exhibited on the counterpart directly adjacent to this boundary. At the distal end of the opisthosoma, the pygidium appears relatively well defined, though neither the second nor the third segments appear to be extended. The pygidium curves slightly off the midline, presumably warped by taphonomic processes.

The appendages are preserved in varying degrees of detail. No appendage is preserved completely; preserved material invariably terminates before reaching the tarsus. The pedipalps are present neither in the part nor the counterpart. It is possible that they were folded beneath the carapace or in some way torn off near the coxae. Legs I and II are best preserved, though detail is lost distally from the body. A dark, organically stained area extending from the right leg I in the part is interpreted as a continuation of the leg, including the patella, tibia, and part of the metatarsus. Unlike the femur of this appendage, these segments lack tuberculation or any form of structure more delicate than their gross morphology. This is most likely due to taphonomic processes and is not a true representation of the original condition of the limb. The femora are the best-preserved podomeres, with leg II providing a particularly exceptional example on the left side of the part and right side of the counterpart. This segment retains much of the original tuberculation but, unlike leg I, lacks any further distal segments. Interpretation of the segments is generally based on small areas where preservation is weaker and appear to line up with where segments would presumably be divided. It is likely these liminal regions of the limb, being less robust than the adjacent segments, were lost during fossilization. The femur of leg II, compared to that of leg I, does not appear to be enlarged. Leg III appears to be the most poorly preserved leg, retaining only lobe-shaped structures extending out from the body midway down the carapace. These distorted shapes are interpreted as originally being the first and second trochanters of these limbs, now damaged and contorted. It is unlikely that these areas represent the femur given they lack an elongate morphology and appear roughly as long as they are wide. This width is notably greater than that of the trochanter of any other appendage. Leg IV is the longest preserved appendage, although, as in the case of leg I, significant detail is lost in the distal segments, so their identity is somewhat dubious. The first two trochanters of leg IV are preserved as very small scraps that lack much detail, which are followed by a better preserved elongate structure interpreted as the femur. The femur pinches out distally, followed by a small patch slightly longer than it is wide, interpreted as the patella. Finally, the tibia is the most poorly preserved and the most distal segment the specimen exhibits from leg IV; it is present only as a scrap some distance away from, but still in line with, the terminus of the patella.

Materials and methods

The specimen (Fig. 1) is preserved in a brown, fine-grained concretion, which is split horizontally through the specimen, creating part and counterpart. Several fragmentary plant remains are preserved in longitudinal section on the same plane as the specimen. It was studied and photographed dry, in both direct and low-angle light, using a Leica M205C stereomicroscope, and photographed with a Canon EOS 5D MK II camera mounted on the microscope. Illustrations were drawn using a camera lucida attachment to the microscope at 11× magnification. The pencil drawing was then traced in Adobe Illustrator. All measurements are in mm.

Due to the large size of the new specimen, we posited that it might represent the adult form of a previously described fossil species. In order to investigate this possibility, ontogenetic analyses of both extant and extinct ricinuleids were performed involving carapace length and tubercle size (two characters that are reliably ubiquitous in fossil specimens), in the expectation that these criteria might identify juvenile stages (Selden, 1992). Observations were made on ricinuleid species for which one or more nymphs have been identified and described. The species included in this analysis included all three extant ricinuleid genera: Ricinoïdes karschii Hansen and Sørensen, 1904, R. hansenii Legg, 1976, R. sp., Pseudocellus olmeca Valdez-Mondragón, Francke, and Botero-Trujillo, 2018, Palpocellass conori Tourinho and Saturnino, 2010, C. pelaezi Pittard and Mitchell, 1972, and C. magnus Ewing, 1929. Measurements were made directly from the literature for all species, excepting P. pearsei, C. magnus, R. karschii, and R. sp. These species were analyzed instead from a collection of previously undescribed nymphal stages in the collections of the American Museum of Natural History (AMNH), and the California Academy of Sciences (CAS).

Tubercle diameter was divided by the length of the carapace to determine the “tubercle coefficient” (TC), a measure of the size of the tubercles compared to the size of the organism. Carapace length, as opposed to the total body length, was chosen because it has been shown to vary less between individuals than total body length (Adis et al., 1989). In keeping with previous assessments of tubercle diameter, measurements were taken from the carapace only (Selden, 1992). Measurements were determined using ImageJ (imagej.net); 20 tubercles, the highest number that could reliably and consistently be found in fossil
specimens, were measured from each individual. In the case of asymmetric tubercles, diameters were taken along their longest axis. When tubercles appeared in clumps, as in the case of the Ricinoides specimens, diameters were taken from the individual tubercles making up these clumps. All five life stages were sampled in six of the 10 species; measurements for P. quetzalcoatl exclude the protonymph, while P. olmeca and C. conori do not include the protonymph or deutonymph. Tubercles appeared to exhibit consistent overall morphologies between life stages (i.e., clumped tubercles in the protonymph of R. karschii developed into clumped tubercles in the adults), although no intensive examination of their morphologies was attempted.

Two-way analyses of variance (ANOVA) were performed in R (r-project.org) to determine whether TC varies significantly between life stages (R Core Team, 2019). The data for each species was divided into two columns: TC, calculated individually for each of the 20 diameters, and life stage (LS). Normality was confirmed using the diagnostic plots in R. Linear models were constructed from the data, treating LS as the independent variable and TC as the dependent variable. A null linear model was constructed for each species assuming no influence of LS on TC, and an alternative linear model was constructed assuming LS did influence TC. ANOVA tests were performed to determine if any significant difference (p < 0.05) was present between the fitted data of these two models.

Two one-sided t-tests (TOST) were utilized to test for statistical equivalence (p < 0.05) of TC between life stages. Unlike ANOVA tests, TOST analyses assume the samples are statistically different as the null hypothesis, with the alternative hypothesis indicating the two samples are equivalent (Limentani et al., 2005). TOST analyses are primarily used for equivalence testing in the pharmaceutical industry, but more recently have been employed to test biological hypotheses (Rose et al., 2018). Due to the nature of the test, only two samples can be compared at a time (Limentani et al., 2005). As such, three TOST analyses (R-package ‘equivalence’), performed for each species to assess the equivalence of all stages to each other (Robinson, 2016). The first compared the TC values of the protonymph and deutonymph; the second compared the TC values of the tri- and adult; and the third compared the combined TC values of the protonymph and deutonymph to the combined TC values of the protonymph and adult. When a life stage from a species was not sampled, comparisons were made only between the life stages available. The average standard deviation of the samples being compared was used as the theta value (referred to as “epsilon” in the r-code), the variable representing the amount of variation the means of two samples may exhibit while remaining statistically equivalent (Limentani et al., 2005). The standard deviation is a measure of the variability of a sample that roughly represents the average distance between the data points making up the sample and the mean (McKenzie, 2014). If two means are equivalent, they would be expected to fall within less than a standard deviation from each other; implying each is within the expected average variability of the other. Taking the average standard deviation from both samples mitigates the potential issue of a single large standard deviation from one of the samples guaranteeing equivalence.

ANOVA tests were again performed to determine if TC varies between species in extant genera. Data were divided into two columns, TC and species. One test was performed for each genus. Species was treated as the independent variable and TC as the dependent variable. A null model was constructed that assumed no influence of species on TC, as well as an alternative model that assumed species did influence TC. The fit of the data to these models was compared using an ANOVA test. An additional ANOVA test was performed on the same parameters of the two largest fossil Curculioides species, C. gigas Selden, 1992 and C. bohemondi n. sp., to determine whether this character was species specific in the genus.

Repositories and institutional abbreviations.—The new specimen is held in the Research and Collections Center, Illinois State Museum, under the designation ACCN2015-104. The Oklahoma Amarixys gracilis (Petrunkevitch, 1945) is held in the Field Museum of Natural History with number PE56942.

Systematic paleontology

Order Ricinulei Thorell, 1876
Suborder Posterioricinulei Wunderlich, 2015
Infraorder Palaeoricinulei Selden, 1992
Family Curculioididae Cockerell, 1916
Genus Curculioides Buckland, 1837

Type species.—Curculioides ansticii Buckland, 1837.

Diagnosis.—As in Selden (1992).

Remarks.—The specimen can be placed in Ricinulei on the basis of its possession of a cucullus, which is a synapomorphy for the order. The specimen is referred to the family Curculioididae because of the presence of a median line or sulcus on the dorsal opisthosoma. The new fossil has a median line, not a sulcus, and thus belongs in Curculioides rather than Amarixys Selden, 1992.

Curculioides bohemondi new species

Figures 1, 3.1–3.3

Holotype.—One specimen, ACCN2015-104, from the middle Pennsylvanian (Desmoinesian, 308–306 Ma) Energy Shale Member of the Carbondale Formation, Kewanee Group, near Little Vermilion River, Georgetown, Vermilion County, east-central Illinois (Baird et al., 1985).

Diagnosis.—Curculioides with cuticular tubercles >0.090 mm in diameter and a density of ~30 mm⁻². All other species of Curculioides have smaller tubercles (<0.085 mm) and a much greater density (>60 mm⁻²) (Selden, 1992, table 1); lacking transverse opisthosomal lines seen in C. gigas; raised areas on anterior carapace divided by median longitudinal groove.

Description.—Body length 23.59 mm. Tubercles distributed evenly over entire body, diameter 0.09 mm, density 30 mm⁻².
Leg IV tibia length 4.46 mm, width 1.48 mm. Leg IV patella length 2.30 mm, width 1.7 mm. Trochanter II length 2.24 mm. Leg IV femur length 5.54 mm, width 1.75 mm. Leg III trochanter II length 2.63 mm, width 1.76 mm. Leg II femur length 8.44 mm, width 1.11 mm. Leg III trochanter I length 1.81 mm, width 1.75 mm. Leg III trochanter II length 2.63 mm, width 2.48 mm. Leg IV trochanter I length 1.33 mm. Leg IV trochanter II length 2.24 mm. Leg IV femur length 5.54 mm, width 1.36 mm. Leg IV patella length 2.30 mm, width 1.7 mm. Leg IV tibia length 4.46 mm, width 1.48 mm.

Etymology.—Named for the eleventh century crusader Marc Guiscard, who received the nickname Bohemond (after a mythical giant) due to his immense stature, in recognition of the large size of the fossil specimen.

Remarks.—Tuberculation is one of the main characters used to distinguish the fossil ricinuleids (Selden, 1992). Figure 4 shows a plot of tubercle diameter against density for Curculioides species. Curculioides bohemondi n. sp. plots far from the other species in this genus. The tubercles of Curculioides bohemondi n. sp. are much larger (>0.090) than those of other Curculioides species (see Selden, 1992, table 1). The smallest are found in Curculioides pococki Selden, 1992 (<0.055), in Curculioides granulatus Petrunkevitch, 1949 they are <0.065, and in Curculioides mcluckiei Selden, 1992, Curculioides eltringhami Petrunkevitch, 1949, and Curculioides scaber Scudder, 1890 they are 0.065–0.085 mm in diameter. In Curculioides gigas Selden, 1992 the tubercles are bimodal in size, with the larger being ~0.073 mm in diameter. There are also differences in tubercle density. The greatest density occurs in Curculioides pococki (>95 mm^2), with a high density (~90 mm^2) also found in Curculioides adompha Brauckmann, 1987 from the Namurian (approximately Serpukhovian–Bashkirian) of Germany. The density in Curculioides scaber is >85 mm^2; it is 65–85 mm^2 in Curculioides granulatus and Curculioides mcluckiei, and <65 mm^2 in Curculioides eltringhami. However, in Curculioides bohemondi n. sp. the density is sparse at ~30 mm^2.

Other characters used to distinguish Curculioides species are the presence of a pair of longitudinal ventral sulci in Curculioides ansticci and the presence of transverse lines on the dorsal opisthosoma in Curculioides gigas. Curculioides scaber possesses a triangular sclerite at the anterior terminus of the opisthosomal median longitudinal line. Paired pits posterior and lateral to the median longitudinal groove of the carapace are seen in most Curculioides species. None of these characters is present in the new specimen (although the presence of ventral sulci cannot be known because the ventral surface is not visible). Curculioides bohemondi n. sp. is the only known species within Curculioides that possesses raised carapace areas.

Genus Amarixys Selden, 1992

Type species.—Amarixys sulcata (Melander, 1903).

Amarixys gracilis (Petrunkevitch, 1945)

Holotype.—Curculioides gracilis Petrunkevitch, 1945 (p. 68–70, text-fig. 34, pl. 11, figs 8–10), ISM 14862, part and counterpart, Illinois State Museum, Springfield, IL; from Westphalian D of Mazon Creek, Illinois.

Diagnosis.—As in Selden (1992).

Remarks.—The specimen of this species, from a strip mine tip in Westphalian rocks, north-west of Morris, Okmulgee County, Oklahoma, which was formerly numbered MCP 548 and held in the collections of the Mazon Creek Project, Northeastern Illinois University, Chicago, was described by Selden (1992), but not figured. It has now been accessioned to the Field Museum in Chicago as PES6942, and the opportunity is taken here to figure it (Fig. 2). Notice that this curculionid genus is easily separated from Curculioides by the median sulcus on the dorsal opisthosoma.

Discussion

Ontogeny.—The incomplete nature of many fossil ricinuleid specimens renders it difficult to determine the life stage to which the individual belonged. This causes ambiguity in species diagnoses because conspecifics in different stages of development may be mistaken for different species. To alleviate this issue, some general growth trends in modern ricinuleids, which may be applicable to fossil specimens, are described and applied to the new specimen.

Modern adult male and female ricinuleids can readily be distinguished by the terminal segments of leg III, modified in the male to form a conspicuous copulatory apparatus (Pittard and Mitchell, 1972). Due to the incomplete preservation of fossil specimens, this specific trait is not often available. Remaining potentially identifiable in the fossil record however is the sexually dimorphic leg thickening exhibited in many adult male
ricinuleids. This limb thickening is characterized by wider basal segments of leg II compared to the other limbs of the organism, although males of some species (e.g., Cryptocellus becki Platnick and Shadah, 1977 and C. iaci Tourinho, Lo Man-Hung, and Bonaldo, 2010) instead exhibit this thickening on the basal segments of leg III (Adis et al., 1989; Tourinho et al., 2010; Salvatierra et al., 2013). The inflated limbs of the adult males appear to form only at maturity; they are not generally observed in tritonymphs. While the degree of thickening may differ between species and individual males, no thickening is observed in females.

This same leg thickening may be exhibited in the specimen of Curculioioides bohemondi n. sp. The basal segments of leg III appear roughly as long as they are wide, giving them an inflated appearance in comparison to the other legs. If the trend observed in extant species is an indication, this could suggest the specimen represents an adult male. This region of the fossil is not very well preserved in general however, so a definitive conclusion is difficult to make.

Another trend throughout development is the increasing depth and definition of the carapace median longitudinal groove. In species that exhibited a median longitudinal groove, the most well-defined groove was found in the adult, a slightly less-defined groove in the tritonymph, and, if present at all, a very vaguely defined groove in the deutonymph of a few species (Cryptocellus magnus, C. iaci, P. pearsei). This was generally associated with a change in tubercle distribution as well, with a sparser tubercle density closer to the groove, becoming more apparent up to maturity.

The well-defined median longitudinal groove of Curculioioides bohemondi n. sp. would suggest the specimen represents an adult or possibly a tritonymph. Although this character does not provide a precise life stage determination, it does provide utility for comparative analysis. Curculioioides mcluckiei is another Pennsylvania ricinuleid from Illinois with morphological similarities to C. bohemondi n. sp., however the median groove confirms that these two do not represent the same species. The C. mcluckiei holotype exhibits a median longitudinal groove at least as well defined as that of C. bohemondi n. sp., suggesting the specimen was near maturity despite being significantly smaller than C. bohemondi n. sp. in overall size. This would imply that C. bohemondi n. sp. is in fact a different species with a larger adult size.

Further diagnostic information was revealed through the ontogenetic analyses of tubercle size. ANOVA tests examining how the tubercle coefficient changed through maturation found no significant (p < 0.05) difference in TC between life stages in any of the species examined (Table 1). TOST analyses found significant (p < 0.05) results in all species examined, suggesting that TC remains equivalent between life stages (Table 2). Further ANOVA tests examining how TC differs between species found significant (p < 0.05) results in all tested genera, suggesting that the null hypothesis of species having no influence on TC be rejected (Table 2). These tests demonstrate that TC is an ontogenetically stable character that can be used to differentiate species regardless of life stage.

The consistency of the tubercle coefficient between life stages implies isometric growth of the tubercles with regard to the carapace. The ratio of tubercle diameter to carapace length remained the same regardless of the life stage examined in all tested modern species. Because the incomplete preservation of Pennsylvania specimen limbs prevents life stage from being empirically determined, the presence of this character in the Paleozoic must be inferred. The modern ubiquity of this character, present in all extant genera, would heavily imply its ancestral nature. It is therefore very likely that the Pennsylvania ricinuleids exhibited this isometric growth pattern as well.

To examine if the character is useful in differentiating previously described Pennsylvania ricinuleids from the new specimen, the tubercle coefficient of Curculioioides gigas, a geographic compatriot and contemporary of C. bohemondi n. sp., was compared to that of C. bohemondi n. sp. using an ANOVA test. The two (C. bohemondi n. sp. with a mean TC of 1.5e-2 ± 1.5e-3 and C. gigas with a mean TC of 8.0e-3 ± 8.6e-4) were found to be significantly (p < 0.05) different, supporting the conclusion that the two are different species. Although the lack of variability in TC through ontogeny effectively disqualifies it for use in determining life stage, the species-specific nature of the character suggests its utility in diagnosing both modern and fossil species. This has notable significance to the fossil ricinuleids, among which, if life stage cannot be determined, cryptic conspecifics may more easily remain undetected. The use of the tubercle coefficient in species diagnosis, consistent across the entire lifespan of the organism, circumvents this issue.

**Taphonomy and geologic significance.**—The presence of a terrestrial ricinuleid in an assemblage of primarily aquatic organisms is explained by the interpretation of the Energy Shale as a crevasse splay deposit (Baird et al., 1985). The levee breakage involved in the deposition of a crevasse splay unleashes water from the river onto the surrounding floodplain, catching terrestrial organisms such as ricinuleids off guard and depositing them among aquatic organisms and

### Table 2. Results of TOST analyses testing for equivalence (p < 0.05) of tubercle coefficient between life stages. Dashes (–) indicate comparisons that were not performed in this study. *PD-TA* represents the combined protonymph and deutonymph values compared to the combined tritonymph and adult values.

<table>
<thead>
<tr>
<th>Species</th>
<th>Protonymph-Deutonymph</th>
<th>Tritonymph-Adult</th>
<th>PD-TA*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ricinoides karschi</td>
<td>2.73e-2</td>
<td>2.08e-2</td>
<td>3.85e-4</td>
</tr>
<tr>
<td>R. hanseni</td>
<td>3.59e-3</td>
<td>2.82e-3</td>
<td>2.68e-2</td>
</tr>
<tr>
<td>R. sp.</td>
<td>7.44e-3</td>
<td>1.55e-3</td>
<td>3.24e-2</td>
</tr>
<tr>
<td>Cryptocellus magnus</td>
<td>1.00e-2</td>
<td>8.24e-3</td>
<td>2.07e-2</td>
</tr>
<tr>
<td>C. pelaezii</td>
<td>3.84e-2</td>
<td>4.80e-3</td>
<td>2.85e-5</td>
</tr>
<tr>
<td>C. conori</td>
<td>–</td>
<td>1.06e-2</td>
<td></td>
</tr>
<tr>
<td>Pseudocellus pearsei</td>
<td>6.94e-3</td>
<td>3.73e-2</td>
<td>2.19e-3</td>
</tr>
<tr>
<td>P. quetzalcoatl</td>
<td>–</td>
<td>1.55e-3</td>
<td>4.79e-4</td>
</tr>
<tr>
<td>P. olmeca</td>
<td>–</td>
<td>1.20e-2</td>
<td>–</td>
</tr>
</tbody>
</table>

### Table 3. Results of ANOVA tests measuring the significance (p < 0.05) of the effect of species on the tubercle coefficient.

<table>
<thead>
<tr>
<th>Genus</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ricinoides</td>
<td>4.39e-14</td>
</tr>
<tr>
<td>Cryptocellus</td>
<td>&gt;2.2e-16</td>
</tr>
<tr>
<td>Pseudocellus</td>
<td>&gt;2.2e-16</td>
</tr>
</tbody>
</table>
sediments from the river (Treworgy and Jacobson, 1985). Indeed, the ricinuleid very likely drowned based on the position of its limbs, spread out as if turgid from the surrounding water (Cooke, 1967). The visible unumbrellated groove on the anterior portion of the specimen’s opisthosoma is part of the ricinuleid carapace-opisthosoma locking mechanism—it is not visible when the two body segments are in the usual locked position (Pittard and Mitchell, 1972). The visibility of this groove would suggest the carapace and opisthosoma are in the “unlocked” position, perhaps another result of the specimen’s turbulent final moments.

The allochthonous deposition of the ricinuleid makes interpretation of the specimen’s original paleoenvironment difficult, but not impossible. The Herrin No. 6 Coal, representing a heavily forested environment, likely represents the kind of ecosystem the ricinuleid would have called home (Treworgy and Jacobson, 1985). Deposition of the Herrin No. 6 Coal is considered contemporaneous with the Energy Shale, and in places is even interbedded with it (Burk et al., 1987). Additionally, the presence of in-situ tree stumps within the Energy Shale directly demonstrates how material from the Energy Shale crevasse splay would have covered these active swamps and forests with sediment (Nelson, 1983). Most likely the ricinuleid was originally a resident of one of these forests and was swept up in a sudden flooding event and deposited within the crevasse splay.

The siderite concretion itself is indicative of a freshwater environment. Only an aquatic system provides the water chemistry necessary for the formation of such a feature (Woodland and Stenstrom, 1979). Likely the mechanics of concretion formation would mirror that displayed in the Mazon Creek, where, after initial burial, a small amount of aerobic decomposition created a microenvironment around the organism ideal for the precipitation of siderite; with enough time the concretion forms, preserving the specimen (Woodland and Stenstrom, 1979).

Comparison to the Francis Creek Shale.—The Energy Shale is a new locality for fossil ricinuleids, but remains in close geographic and temporal proximity to the more renowned Mazon Creek of the Francis Creek Shale where the majority of North American fossil ricinuleids have been found (Selden, 1992). As such, a brief discussion comparing the two strata is warranted.

The Mazon Creek fauna belongs to the Francis Creek Shale, found in Grundy County Illinois, located northwest of the locality for C. bohemondi n. sp. in Vermilion County Illinois (Baird, 1979). Neither location offers a continuous stratigraphic column featuring both members. The Energy Shale is found mostly in southern and eastern Illinois, while the Francis Creek Shale is found in northern Illinois (Baird, 1979; Burk et al., 1987). However, the lateral continuity of the Herrin No. 6 Coal provides a benchmark, suggesting the Energy Shale overlying it is somewhat younger than the Francis Creek Shale that, where it is deposited, is significantly beneath the Herrin (Willman et al., 1975).

Both the Energy Shale and the Francis Creek Shale are thought to broadly represent river delta systems, but there are some important differences (Burk et al., 1987; Clements et al., 2018). Lithologically, the Energy Shale is composed primarily of siltstones and silty shales that decrease in grain size with increasing distance from the Walshville Channel (Burk et al., 1987). The Francis Creek Shale is composed of shales and siltstones that grade laterally into crossbedded sandstones in multiple areas (Baird, 1979). Although the lithologies are similar, their distribution suggests a difference in the morphology of the deltaic system. Whereas the Energy Shale was fed primarily by the singular Walshville Channel, the Francis Creek Shale was apparently sourced by numerous smaller rivers (Treworgy and Jacobson, 1985).

Another important difference is in the interpreted salinity of the two deposits. The Energy Shale is largely a freshwater deposit, as signaled by its aquatic biota and the low sulfur content of its contemporaneous underlying coal (Nelson, 1983; Baird et al., 1985). Only at the farthest extent from the Walshville Channel does the unit appear to exhibit a marine influence, where euryhaline bivalves indicate a brackish bay system (Burk et al., 1987). The Francis Creek Shale, however, features marine influence to different degrees throughout the unit (Clements et al., 2018). Brackish and marine biota in addition to aquatic and terrestrial organisms, as well as the ubiquitous pinstriped bedding indicative of tidal rhythms all suggest a sizable marine influence (Clements et al., 2018). Siderite concretions, which require a freshwater system in which to precipitate, are explained by the frequent but periodic recession of these tides, allowing the concretion to form in a temporary freshwater environment (Woodland and Stenstrom, 1979). However, it appears that the nonmarine organisms preserved within these Francis Creek Shale concretions are not necessarily contemporaneous with the deposition of the unit—leading to another important difference between the two members (Clements et al., 2018).

Both members overlie a coal bed—the Energy Shale being situated above the Herrin No. 6 Coal while the Francis Creek Shale is above the Colchester No. 2 Coal (Willman et al., 1975; Jacobson et al., 1980). The relationship between these shales and their underlying coals differs in timing of deposition (Baird, 1979; Jacobson et al., 1980). Whereas the Energy Shale is thought to have been deposited concurrently with the peat deposition of the Herrin No. 6 Coal, the Francis Creek Shale is thought to have been deposited after the deposition of the Colchester Coal (Burk et al., 1987; Clements et al., 2018). Nonmarine and terrestrial biota preserved in the Francis Creek Shale are thought to have been integrated into the member from already inundated swamps in the deltaic system, flooded by the transgressing ocean (Clements et al., 2018). Concretions are thought to have been formed around already-dead organisms at the bottom of these flooded brackish swamps (Clements et al., 2018). The result is a system that, although paleontologically and lithologically similar, is actually fairly disparate in terms of the intricacies governing fossil formation within it.

Size.—It is widely known that many insects of the Carboniferous (Mississippian–Pennsylvanian) exhibited some form of gigantism as compared to their modern forms (Shear and Kukalová-Peck, 1990). This has been attributed to a higher oxygen content in the planet’s atmosphere at the time, allowing the organism to absorb sufficient amounts of the gas despite the smaller surface area to volume ratio implied by a larger overall size (Shear and Kukalová-Peck, 1990). The
tracheal system of insects is made more efficient by higher atmospheric oxygen partial pressure (Dudley, 1998). In an environment without a high-oxygen atmosphere, large insects must dedicate a high amount of resources to the expansion of their tracheal system simply to supply the necessary oxygen to tissues for their metabolic function (Kaiser et al., 2007).

Among Pennsylvanian ricinuleids, C. bohemondi n. sp. is the largest by a significant margin. The closest in length (excluding the cucullus and pygidium) is C. gigas, at 18 mm long, whereas C. bohemondi n. sp. is 20.98 mm long (Selden, 1992). Both species, however, dwarf even the largest known extant ricinuleid, which only measures ~6 mm long (Naskrecki, 2008). It is possible that this change in size may be due in part to the higher oxygen content of the Pennsylvanian atmosphere. Though not insects themselves, extant ricinuleids exhibit a respiratory system composed of sieve trachea, not entirely unlike the respiratory system of insects (Levi, 1967). If fossil ricinuleids exhibited this similar system, then the higher oxygen partial pressure would likely have a similar effect to that which it has on insects, allowing the organism to bypass the limitations of oxygen diffusion with a higher-efficiency system that requires less energy investment into the trachea (Levi, 1967). However, this comparative gigantism is relatively minimal compared to the larger forms of some Carboniferous arthropods, and as such, is unlikely to have been entirely due to higher atmospheric oxygen content. Differing ecologies between ricinuleids of these two disparate eras likely played an important role as well (Shear 1942; Lythgoe, 1979).

The ancestral presence of the leg-based sensory structure is suggested in both Cretaceous and Pennsylvanian ricinuleids. The Cretaceous species exhibit an elongate second leg pair, which is the typical morphology of the modern leg II sensory system (Wunderlich, 2015). Although no Pennsylvanian ricinuleid specimens exhibit preserved tarsi, the segments of the second leg are consistently the longest among the limbs in all known species, including C. bohemondi n. sp., implying an elongate second limb in the Paleozoic species as well (Selden, 1992). Similar to the antennae of modern insects, most likely the second leg pair has always served a sensory function, originally supplementing the visual information from the eyes and allowing behaviors to continue in the dark (Pelletier and McLeod, 1994). Investigation regarding the presence of sensory sensilla on the surface of fossil ricinuleid tarsi will be necessary to confirm this, however, because the possibility remains that the preexisting elongate second limbs may have been later coopted to serve a primarily sensory function. In the absence of an alternative original purpose for these disproportionately long limbs, it seems plausible that extinct ricinuleids such as C. bohemondi n. sp. would have used them to probe the lycopod-forest floor for small invertebrate prey, similar to their modern relatives (Clements et al., 2018).

Paleoecology.—Due to the incomplete nature of the specimen and the problems inherent in interpreting ecology from an isolated concretion (let alone one isolated in an environment separate from that in which it likely lived), definite conclusions regarding the ecology of the specimen are difficult to make; however, some speculation can be made. All modern ricinuleids occupy one of two overarching niches—terricolous species (living beneath the rainforest floor and into the subsoil), and cavernicolous species (living in the complete darkness of subterranean caverns) (Selden, 1992; Adis et al., 1999). It appears however that the ancestral niche of the order is not reflected in these modern ecologies. The recently discovered Cretaceous ricinuleids are preserved in amber, suggesting that they may have exhibited a scanorial lifestyle or at least frequented the bases of tree trunks—an unusual setting for modern ricinuleids (Cooke, 1967; Wunderlich, 2015).

Likewise, the ubiquity of eyes among Pennsylvanian species would also suggest they were surface dwellers. Eyes are not present in extant ricinuleids, although some species exhibit carapacial translucent areas, which may or may not serve a rudimentary photosensitive function (Botero-Trujillo and Valdez-Mondragón, 2016). It is likely their eyes were lost due to the transition to the low-light environments of the subsurface, a phenomenon occurring in many species that have made similar ecological shifts (Sumner-Rooney, 2018). Their presence in fossil species however would indicate the ancestral life habit was surficial.

Modern ricinuleids possess modified first and second leg pairs, the tarsi of which are covered with specialized sensilla and slit organs that perform thermosensory, hygrosensory, gustatory, olfactory, mechanosensory, and proprioceptive functions (Talarico et al., 2006). The long length of the second leg makes it particularly well suited for sensory purposes; modern ricinuleids tap these limbs exploratorily on the ground as they navigate their environment (Talarico et al., 2005). The apparent presence of this structure in eye-bearing fossil forms would suggest an ancestral dual sensory system, with the non-visual sensory apparatus of the legs allowing ricinuleids to thrive in a cavernicolous habitat even early into the order’s subterranean invasion. This trend of preexisting non-visual sensory structures allowing for adoption of a cavernicolous lifestyle has been previously documented in many clades, most notably catfish (Walls, 1942; Lythgoe, 1979).

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