Hidden diversity in the Brazilian Atlantic rainforest: the discovery of Jurasaidae, a new beetle family (Coleoptera, Elateroidea) with neotenic females

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Beetles are the most species-rich animal radiation and are among the historically most intensively studied insect groups. Consequently, the vast majority of their higher-level taxa had already been described about a century ago. In the 21st century, thus far, only three beetle families have been described de novo based on newly collected material. Here, we report the discovery of a completely new lineage of soft-bodied neotenic beetles from the Brazilian Atlantic rainforest, which is one of the most diverse and also most endangered biomes on the planet. We identified three species in two genera, which differ in morphology of all life stages and exhibit different degrees of neoteny in females. We provide a formal description of this lineage for which we propose the new family Jurasaidae. Molecular phylogeny recovered Jurasaidae within the basal grade in Elateroidea, sister to the well-sclerotized rare click beetles, Cerophytidae. This placement is supported by several larval characters including the modified mouthparts. The discovery of a new beetle family, which is due to the limited dispersal capability and cryptic lifestyle of its wingless females bound to long-term stable habitats, highlights the importance of the Brazilian Atlantic rainforest as a top priority area for nature conservation.

Coleoptera (beetles) is by far the largest insect order by number of described species. Approximately 400,000 species have been described, and many new ones are still frequently being discovered even in regions with historically high collecting activity¹. Because this group has been intensively studied since the age of early systematists, and most of the major higher-level taxa had already been described by about a century ago, discoveries of new beetle families are nowadays a rare occurrence. The vast majority of recent family descriptions is based on the already known taxa for which a new systematic placement was recovered by the use of phylogenetic analyses²–⁵. Thus, only three beetle families have been erected de novo in the 21st century based on newly collected material. Firstly, Aspidytidae were described based on two species from South Africa and China, which currently represent monotypic genera⁶,⁷. Secondly, the monotypic family Meruidae was created for a hitherto undescribed lineage from Venezuela⁸. Finally, the new monogeneric family Iberobaeniidae was described based on three species from the southern part of the Iberian Peninsula⁹,¹⁰. Whereas both Aspidytidae and Meruidae represent semiaquatic families within the suborder Adephaga, Iberobaeniidae were recovered as a deep branch in the polyphagan superfamily Elateroidea⁹.

Elateroidea (the click-beetles, soldier-beetles, fireflies and relatives), the largest superfamily of the series Elateriformia, form one of the major and oldest polyphagan lineages¹¹,¹². Currently, they contain approximately 25,000 described species classified into 13 families¹³–¹⁶. The elateroid beetles exhibit extraordinary morphological diversity. Some lineages contain representatives with a fully sclerotized, compact body and presence of a defensive pro-mesothoracic clicking mechanism, while other groups are soft-bodied, with loosely connected abdominal
segments and often with variously modified or reduced body parts\textsuperscript{13,17}. Such morphological modifications are a result of neoteny, a phenomenon in which animals retain larval characters while attaining reproductive maturity\textsuperscript{18–20}. In Elateroidea, females are usually much more affected by neoteny and exhibit various degrees of incomplete metamorphosis; from possession of a larva-like abdomen only, through having a larva-like body excluding the head, to a completely larviform body\textsuperscript{16,21–23}. Since developmental modifications lead to similar morphological traits, these groups were usually considered a monophylum (i.e., former Cantharoidea)\textsuperscript{17,21}. However, recent molecular-based research showed the multiple origin of soft-bodiedness and neoteny in Elateroidea\textsuperscript{9,13,16,22,23}.

Here, we report the discovery of a completely new elateroid neotenic lineage from the Brazilian Atlantic forest ecoregion, which is one of the Earth’s most endangered biodiversity hotspots with high levels of diversity and endemism\textsuperscript{24,25}. The story of a discovery began on 24 May 2016, when an expedition of the first author (SPR) and her undergraduate students was organized to the Municipal Biological Reserve Serra dos Toledos in the Serra da Mantiqueira mountain chain as part of research on its Lampyridae fauna\textsuperscript{26}. During that expedition, the first two larvae of the here described new elateroid lineage were collected in soil of a ravine along the main trail of the reserve (Fig. 1). Based on their bizzare morphology, including the remarkable mouthparts which resembled a beak, collectors were not able to assign the specimens to any known beetle family, although it was obvious they belong to Elateroidea. A few weeks later the larvae metamorphosed into pupae and later to a soft-bodied adult male and a larviform female. Because they represented an apparently undescribed elateroid lineage, a great effort was made to collect more larvae during every expedition to Serra dos Toledos. In 2017, adult males of a second species were found among samples from Malaise traps in Serra dos Toledos, and Luiz F. L. Silveira (Western Carolina University, USA) sent us adult males of a third species collected using Malaise traps in the Serra dos Órgãos National Park, a part of the Serra do Mar mountain chain. In this study, we conducted a detailed evaluation of the morphological characters of all life stages and investigated the phylogenetic position of the newly discovered lineage using a multi-marker molecular phylogeny. All available evidence demonstrates the need to establish a new family within the Elateroidea for three new species in two new genera.

**Results**

Based on their divergent morphology (Figs. 2–5, S1–S12) and the results obtained from the molecular phylogeny (Figs. 6, S13), we describe here *Jurasai digitusdei* gen. et sp. nov., *J. itajubense* gen. et sp. nov. and *Tujamita plenala-tum* gen. et sp. nov., for which we erect a new family Jurasaidae fam. nov. Below we provide remarks on their biology, ecology and behavior, as well as concise diagnostic descriptions for all new taxa based primarily on salient diagnostic characters. Detailed morphological diagnoses and descriptions are given in the Supplementary Text.

**Observations of behavior of the immature stages and adults.** Immature stages of two species (out of three) representing two different genera are known to date. Larvae and pupae of *J. itajubense* and *T. plenala-tum* occur sympatriically in the Municipal Biological Reserve Serra dos Toledos. They are usually found in the

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**Figure 1.** Habitat of *Jurasai itajubense* gen. et sp. nov. and *Tujamita plenalatum* gen. et sp. nov. in the Biological Reserve of Serra dos Toledos: (a) main trail; (b) ravine along the main trail; (c) soil with roots where most larvae were collected; (d) larva of *J. itajubense* near the root.
5–20-cm-thick layer of soil just under the leaf litter (Figs. 1c, 2a–d, 3). Although both species share the same microhabitat, *T. plenalatum* is much more rare; of the total 87 larvae collected, only four belonged to that species. The feeding habits of both larvae and adults are not known. Despite our offering the larvae various potential food items in the laboratory, no movement of the mouthparts or signs of pumping into the esophagus were observed. However, since many larvae were found between roots, we hypothesize that they feed on juices of fungal hyphae.

Both larvae and adults usually stay in the soil. When unearthed and disturbed they become active and soon bury themselves again into the substrate. Bioluminescence is present in several elateroid lineages in Brazil, i.e., Lampyridae (fireflies), Phengodidae (glowworms) and Elateridae: Pyrophorini (luminous click-beetles)^22,27,28^.

However, it was not detected in Jurasaidae, neither by the naked eye nor by using long exposure photography. In both species, the pupal stage lasts 22–24 days in males and 8–9 days in females. Whilst a male pupa can move only its abdomen, a female pupa is capable of walking, although it usually stays burrowed in the substrate or resting on the surface in a C-shaped position (Fig. 2d). The adult female is wingless and resembles a larva in general

Figure 2. Live specimens of Jurasaidae fam. nov. (a) larvae of *Tujamita plenalatum* gen. et sp. nov. (upper specimen) and *Jurasai itajubense* gen. et sp. nov. (lower specimen); (b) larva of *T. plenalatum*; (c–e) *J. itajubense*: larva, female pupa and adult female, respectively; (f) adult female of *T. plenalatum*.
appearance, with only certain parts of the head adult-like (Jurasaia, Fig. 2e) or with the head, prothorax and legs adult-like, though different in appearance from the corresponding parts in the male (Tujamita, Fig. 2f) (Table S1).

About a week after eclosion, the male actively searches for a female and then copulates several times each for a short period of time (Fig. 4, Supplementary Video). More detailed information on field and laboratory observations are given in the Supplementary Text.

Systematics. Jurasaia gen. nov. (Figs. 1d, 2a,c–e, 3a,b, 4, 5a–d, S1–S4, S6a–i, S8a,b, S10a–i, S11a,b, S12a–h).

Type species. Jurasaia itajubense sp. nov.; by present designation.

Diagnostic description. Adult male (Figs. 4, 5a–d, S1, S2a–h, S4, S6a–i, S8a,b). Body length from frons to apex of abdomen 2.5–3.0 mm, from frons to apex of wing 3.5–5.0 mm. Head with antenna approximately as long as distance between head and elytral apex; labrum 1.8–4.0 times as wide as long; maxillary palpus 4- or 5-segmented; apical labial palpomere 2–5 times longer than basal palpomere. Pronotum (Figs. S2b, S4b,d, S6e) widest at
anterior half, posterior half narrowed posteriad; without lateral carinae; posterior edge smooth, not marginated; mesoventrite (Figs. S4f–h, S6e) 1.1 times longer than wide, with anterior margin deeply arcuate; mesepimeron with anterior part indistinct in lateral view, separated from mesanepisternum by weakly impressed suture; mesocoxal cavities separated at middle by 1.3 times mesocoxal cavity width; metaventrite (Fig. S4f) 1.3 times longer than wide, 2.5 times longer than mesoventrite, widest at anterior 1/3. Elytra (Figs. 5a,c, S1a,f) shorter than abdomen, tapered apically, with median edges separated and divergent apicad, lateral edges sinuate, apices swollen; hind wing (Fig. S8a,b) surpassing elytral apex by 0.5–0.7 times elytral length, apical field long (0.5–0.6 times as long as total wing length). Tarsomere IV (Fig. S1e) evenly sclerotized, truncate apically. Abdomen (Fig. S1h) narrow, with sides subparallel; phallus and parameres (Figs. S2f,g, S4m–o) together 1.6 times wider than long; endophallus emerging from dorso-apical elongate notch. **Adult female** (Figs. 2e, 4, S2i,j, S3) (based on *J. itajubense* only). Abdomen and thorax almost equal to those of larvae; head with a pair of pigmented stemmata; mandible falcate; labrum free; maxilla and labium separated, labium not channeled to fit mandibles; antenna with four antennomeres; leg with tibia and tarsus separated, tarsus 1–1–1, with a pair of claws. **Mature larva** (Figs. 1d, 2a,c, S10a–i, S11a,b, S12a–h) (based on *J. itajubense* only). Body 4–7 mm long, slender, 9–10 times longer than wide; clypeolabrum (Fig. S12c) with two long setae. Mandibles and labium (Fig. S10b) forming elongate and sharpened beak-like mouthpart; mandible (Figs. S11b, S12e,f) with apex gradually sharpened apicad, parallel-sided, base elliptical; hypopharyngeal bracon hyaline. Pronotum (Fig. S12a) with four setae between lateral edge and lateral sclerotized stripe (one pair anteriorly and one pair posteriorly), two setae between stripes anteriorly; mesothorax,
metathorax, and abdominal segments I–VIII (Fig. S1a,b) with one pair of laterodorsal setae near anterior margin, one pair of laterodorsal setae near posterior margin, and a pair of lateroventral setae near posterior margin. Legs (Fig. S10g) separated by 5–6 times diameter of coxa.

**Etymology.** From Tupi-Guarani language; Jura = mouth; saí = minuscule, thin; allusion to the larval mouth. Gender: neuter.

**Composition and distribution.** Jurasaí itajubense (Brazil: Minas Gerais) and J. digitusdei (Brazil: Rio de Janeiro).

**Jurasaí itajubense sp. nov.** (Figs. 1d, 2a–c, 3a, 4, 5a,b, S1a–e, S2, S3, S6a–e, S8a, S10a–i, S11a,b, S12a–h).

**Type material.** Holotype, male, “Brazil, Minas Gerais state, Itajubá municipality, Biological Reserve of Serra dos Toledos (22°25′21.3″S 45°22′06.2″W), 1,358 m, soil ravine, collected as pupa on 7.VI.2018 (adult on 5.VII.2018, died on 8.VIII.2018), Rosa S.P, Barbosa T. & Paiva J. leg.” (deposited at MZUSP). For information on paratypes and other material examined see the Supplementary Text.

**Diagnostic description.** Adult male (Fig. 5a,b). Labrum with anterior margin emarginate; maxillary palpus 4-segmented; elytra strongly tapered; hind wing venation with only RA1+2, MP1+2 veins; abdominal sternite VIII partly exposed; parameres with apices tapered and curved inwards. For more details see the generic diagnostic description and the Supplementary Text.

**Etymology.** From the type locality, Itajubá, in Minas Gerais state, Brazil.

**Jurasaí digitusdei sp. nov.** (Figs. 5c,d, S1–i, S4, S6f–i, S8b).

**Type material.** Holotype, male, “Brazil, Rio de Janeiro State: Teresópolis, Parque Nacional da Serra dos Órgãos, malaise trap, PVE 6B (22°28′11″S 43°0′5.3″W, 868 m), VI.2015, Silveira & Khattar leg.” (deposited at DZRJ). For information on paratypes and other material examined see the Supplementary Text.

**Diagnostic description.** Adult male (Fig. 5c,d). Labrum with anterior margin rounded; maxillary palpus 5-segmented; elytra weakly tapered; hind wing venation with RA1+2, R3, RA1+4, RM loop and medial field veins; abdominal sternite VIII concealed; parameres with apices sausage-like, directed posteroverrad. For more details see the generic diagnostic description and the Supplementary Text.

**Etymology.** From Latin; digitus = finger, dei = of god; allusion to “Dedo de Deus”, a mountain peak near the type locality in Serra dos Órgãos National Park, whose shape resembles a hand pointing up towards the sky.

**Tujamita gen. nov.** (Figs. 2a,b,f, 3c, 5e,f, S5, S6–l, S7, S8c, S9, S10–m, S11c–e, S12i–l).

**Type species.** Tujamita plenalatum sp. nov.; by present designation.

**Diagnostic description.** Adult male (Figs. 5e,f, S5, S6–l, S7, S8c). Body length from frons to apex of abdomen 3.1–4.5 mm, from frons to apex of wing 3.2–4.6 mm. Head with antenna approximately 2/3 as long as distance between head and elytral apex; labrum four times as wide as long; maxillary palpus 5-segmented; apical labial palpomere 6–8 times longer than basal palpomere. Pronotum (Figs. S5d, S6j) weakly narrowed posteriorly, with lateral margins subparallel and carinate; posterior edge marginated; mesoventrite (Figs. S6k, S7–h) 1.3 times wider than long, with anterior margin weakly arcuate; meseviperomer with anterior part distinct in lateral view, separated from mesaneusternum by groove suture; mesocoxal cavities separated at middle by 0.7 times mesocoxal cavity width; metaventrite (Fig. S7g) 1.1 times wider than long, 2.8–2.9 times longer than mesoventrite, widest at midlength. Elytra (Figs. 5e, S5a) slightly shorter or as long as abdomen, parallel-sided, median edges contiguous to apex; apical lobe flattened. Hind wing (Fig. S8c) surpassing elytral apex by 0.2 times elytral length, apical field short (0.3–0.4 times as long as total wing length). Tarsomere IV (Fig. S5e) deeply notched. Abdomen (Fig. S5f) wide, with sides rounded and tapered apically; phallus and parameres (Fig. S7o,p) together 1.1 times wider than long; endophallus emerging from apical oval orifice on dorsal surface of phallus. Adult female (Figs. 2f, S9). Meso-, metathorax and abdomen almost equal to those of larva; head, pronotum and leg (Fig. S9a–g) adult-like but different from those of male: compound eyes very small, flat, not protruded; posterior tentorial pits absent; antenna moniliform, with nine antennomeres; pronotal carinae almost absent, protrochanter reduced to very narrow sclerotized strip; tarsi 4–4–4.

**Mature larva** (Figs. 2a,b, S10–m, S11c–e, S12i–l). Body 4–6 mm long, stout, 5–6 times longer than wide; clypeolabrum (Figs S10j, S12i) with four long setae; mandibles and labium (Figs. S10j,l, S11c–e) forming short, stout, beak-like mouthpart; mandible (Figs. S11d, S12l) with apex abruptly sharpened apically, convergent anteriad, base triangular; hypopharyngeal brachon sclerotized. Pronotum with five setae between lateral edge and lateral stripe (one pair anteriorly, one pair posteriorly, and one setae at midlength), four setae between stripes (one pair anteriorly and one pair posteriorly); mesothorax, metathorax, and abdominal segments I–VIII with six setae at midlength (one pair lateral, one pair laterodorsal and one pair parasagittal); ventral surface of abdominal segments I–VIII with six setae at midlength (two pairs ventrolateral and one pair parasagittal). Legs separated by 8–12 times diameter of coxa.

**Etymology.** From Tupi-Guarani language; Tuja = adult, mitã = child; allusion to neoteny. Gender: neuter.

**Composition and distribution.** Only: T. plenalatum (Brazil: Minas Gerais).
Jurásaidae fam. nov. (Figs. 2–5, S1–S12).

Type genus. Jurasa gen. nov.

Diagnostic description. Male. Body soft (Fig. 5); head (Figs. S2a, S4a, S5d, S6a–c,f,g, S7a,b,d) declivous, fronto- clypeal suture absent; labrum sclerotized, free, separated from head capsule by membrane; antennal insertions elevated and visible from above; antennal filiform, with 11 antennomeres. Mandible falcate, unidentate; maxillar- lary cardo, stipes, galea and lacinia indistinct, last palpomere shorter than all remaining palptome; gular sutures separated; two posterior tentorial pits present. Thorax (Figs. S2b, S4b–h, S5d, S6e,j,k, S7f–h) with pronotum wider than long, narrower than elytral base; proepisternum keel-shaped, strongly convex medially, prosternoal process not extending beyond coxae; procoxal cavities open internally and externally, widely separated by prosternal process; mesoventrite evenly sclerotized with anterior margin arcuate, separated from metaventrite by feebie suture; leg (Figs. S1e,g, S5b,e) with meso- and metatibia oblique, pro- and mesotrochantin exposed; procoxa, mesocoxa and mesal part of metacoxa conical and strongly projecting, tibial spurs long, tarsi 5–5–5, without lamellae or pulvilli. Elytra (Figs. 5, S1a,f, S5a) soft, irregularly punctate, epipleura gradually narrowed posteriorly; wings (Fig. S8) folded longitudinally in resting position, longer than elytra, hiding the abdomen in dorsal view, veins reduced or blurred, cells and transverse cross veins absent, apical field with three triangular selerotizations and a notch between to two most apical ones. Abdomen (Figs. S1h, S4h, S5f) with five free ventrites (i.e., sternites III–VII); sternites II and VIII largely membranous and concealed; sternite IX (Figs. S2e, S4l, S5g, S7m) with apex bilobed, densely pilose and projecting above phallus and parameres; punctures surrounded by usually 2–3 campaniform sensilla in a dog’s-paw pattern (Fig. S1i). Aedeagus (Figs. S2f–h, S4m–o, S5g–o) trilobate, not entirely retractable into abdomen, phallobase 2.0–2.7 times as long as wide, 1.4–2.6 times as long as parameres, sheathed by a tubular membrane which opens anteriorly into a pair of large balloon-like membranous vesicles; parameres and phallus together 1.1–1.6 times wider than long, basal struts of phallus absent; endophallus emerging from dorso-apical opening, flagellum absent. Female (Fig. 2e,f). Body elongate, wingless, with varying degree of neoteny, but always with at least meso-/metathorax and abdomen larva-like (except for ooporus in posterior margin of sternite VIII) (Fig. S3j,k) (Table S1); leg short, with paired claws; ooporus with pair of membranous lobe-like valves, each with supporting sclerotized plate. Larva (Figs. 1d, 2a–c, S10, S11, S12a–f, i–l). Body cream or milky-white, cylindrical, with few setae; head sclerotized, wedge-like; epicranial sutures and endocarina absent; clupeolabrum triangular, translucent; gula, maxillae and labium fused to each other and to head capsule ventrally, anterior part of labium projected with pair of channels dorsally, fitting apical half of strongly sharpened mandibles, forming a beak-like mouthpart; basal half of mandibles elongate and retracted into anterior 2/3 of head, linked to inner sclerotized rod that extends into prothorax; pronotum with pair of parasagittal selerotized stripes; sternum with strongly sclerotized median longitudinal rod; leg short, 5-segmented; trochanter ring-shaped, pretarsus glabrous.

Composition. Jurasa (two species) and Tujamita (monotypic).

Distribution. Brazil (Minas Gerais, Rio de Janeiro).

Phylogenetic analyses. All phylogenetic analyses placed Jurásaidae in Elateroidea and showed an identical backbone topology for the superfamily, with the main clades branching off in the following order: Artematopodidae + Omethidae (incl. Telegeusinae), Throscidae, Eucnemidae, Cerophytidae + Jurásaidae, and the terminal clade of “higher elateroids” sensu Kundrat et al.13 (i.e., Lycidae, Iberobaeniidae, Lampyridae, Cantharid, Elateridae, Omalisidae, Phengodidae and Rhagophthalmidae). The maximum likelihood (ML) phylogenetic tree of 251 terminals with collapsed branches and Elateroidea families highlighted is given in Fig. 6; the full-resolution tree is in Fig. S13. Jurásaidae were always placed as a sister group of Cerophytum elateroides (Cerophytidae), with 90–95% bootstrap support in the ML analyses and 100% posterior probabilities in the Bayesian (BI) analyses. This clade was sister to the “higher elateroids” with robust statistical support in the BI analyses and weaker support in the ML analyses (Figs. 6, S13). Jurásaidae formed a maximally supported monophylum, with Tujamita sister to a clade formed by two species of Jurasa. We associated the different developmental stages and sexes using molecular markers to confirm the field and laboratory observations (Fig. S14). The uncorrected pairwise genetic distance between coxl sequences for J. itajubense and T. plenulatum was 28.5% (coxl was not available for J. digitusdei).
each one with a supporting sclerotized plate (Fig. S3,j,k), and the larvae are characterized by the following unique characters: fused ventral mouthparts (i.e., maxillae and labium), labial channels located dorsally, and the basal half of the mandibles retracted deeply into the head capsule and linked to an inner sclerotized rod that extends into the prothorax (Figs. S10b,d,e, S11, S12c–e–f, g–i–l). Apart from the above-mentioned unique morphological features, Jurasaidae differ from each of the other soft-bodied elateroid groups in the Neotropics, i.e., Phengodidae, Lampyridae, Cantharidae, Lycidae, and Omethidae: Telegeusinae, by a number of additional features (see the Supplementary Text).

Due to the parallel origins of soft-bodiedness, body miniaturization and the characters connected with prematurely terminated metamorphosis in Elateroidea9,14,16,22,23,29, a detailed phylogenetic placement of Jurasaiidae based on morphology alone would be precarious. Indeed, a preliminary examination of J. itajubense led the discoverers of this species to place it tentatively in Penicilliphorini, a small group of uncertain position, either in Phengodidae or Omethidae: Telegeusinae10,31. However, our detailed morphological investigation revealed the considerable differences between Jurasaiidae and Penicilliphorini (see the Supplementary Text) and the molecular phylogenetic analyses placed Jurasaiidae unambiguously to basal elateroid splits as a sister group of Cerophytidae, far from Phengodidae, Telegeusinae and other soft-bodied lineages (Figs. 6, S13). Although Jurasaiidae and Cerophytidae adults are superficially different (i.e., both sexes in Cerophytidae are completely metamorphosed, with a fully sclerotized body and functional clicking mechanism), their larvae share the cylindrical, white and grub-like body, very small wedge-shaped head without dorsal and ventral epicranial sutures, non-opposable, flattened and channelled mandibles, labium with channels to fit the mandibular apices, prostermum with a median sclerotized endocarinate rod, and short legs. Remarkably, many of these features are also shared with Eucnemididae and Throscidae, and were used to define a monophyletic group formed by Eucnemididae, Throscidae and Cerophytidae22,33. On the other hand, the mandibular apices are an unambiguous synapomorphy of Jurasaiidae + Jurasaiidae. Since the phylogenetic relationships among the basal elateroids including the above-mentioned families have not yet been satisfactorily resolved (and this is also beyond the scope of this paper), more effort should be made to test the monophyly of Eucnemidae, Throscidae, Cerophytidae and Jurasaiidae using more robust datasets and analytical approaches11–13,32,33. This would be crucial for understanding the evolution of modified larval morphology associated with the adaptation to burrowing in soil or rotten wood and feeding on juices of fungal hyphae.

The discovery of Jurasaiidae and their placement in the basal Elateroidea shed new light on the evolution of neoteny in this beetle superfamily. Neoteny in Elateroidea has long been a widely studied phenomenon19,21, and recent studies repeatedly confirmed that it originated several times not only within the superfamily but also within several distantly related families14,20,22,23,29. Elateroidea are divided into the “basal grade” with mostly hard-bodied groups with only Omethidae and Jurasaiidae having a soft body, and the robustly supported clade of “higher elateroids” which contains predominantly fully sclerotized click-beetles and the vast majority of soft-bodied lineages14. It is hypothesized that soft-bodiedness in both adult sexes is the initial stage of ontogenetic modifcations leading to development of highly modified larviform females20. Until now, the neotenic females with variously modified morphology were known almost exclusively within terminal higher elateroids, and Omethidae: Telegeusinae were the only basal lineage with unknown but supposedly neotenic females. Jurasaiidae represent the second loss of sclerotization among basal elateroid lineages, and are the first proven case of highly modified neotenic females outside the terminal clade of higher elateroids. Interestingly, the females of both genera in Jurasaiidae exhibit different degrees of neoteny, with Jurasai being almost completely larviform and Tujamita having the head, prothorax and legs partly adult-like, although more reduced compared to an adult male (Supplementary Text, Table S1). Similar cases of different levels of morphological modifications in females of different genera were reported also for Lampyridae19, Lycidae20, Rhagophthalmidae34 and Elateridae23. Not only females, but also males in Jurasaiidae, present interesting morphological modifications connected with soft-bodiedness and neoteny. In Elateroidea, divergence in the abdominal morphology represents a continuous scale from five ventrites of which all or at least some are connate (the majority of the well-sclerotized elateroids) through various intermediate stages to seven or eight ventrites, which are all free and connected by extensive intersegmental membranes (majority of the soft-bodied lineages)39. Jurasaiidae males represent an intermediate form; their abdomen is soft and with five free ventrites (Figs. S1h, S5f). A similar condition is only known in the otherwise morphologically dissimilar Brachysceptidae. Moreover, similar to females, Jurasai males are morphologically more affected by ontogenetic modifications than Tujamita males, having a pronotum without strengthening structures (pronotum with lateral carinae in Tujamita), elytra with median edges divergent posteriorly and with separately tapered apices (elytra parallel-sided, with sutural edges contiguous to apex in Tujamita), and with the abdomen narrower and more exposed (abdomen wider and shorter, usually covered by elytra in Tujamita) (Figs. 5, S1h, S5f, S6e–k). Finally, the two species of Jurasai exhibit different degrees of morphological modifications, with J. itajubense being more neotenic than J. digitusdei, having a reduced number of maxillary palpomeres, less compact and relatively narrower elytra, and much more reduced hind wing venation (Figs. 5a–d, 5a,b,e–g, S6c,g, S8a,b). Our present findings therefore represent additional evidence for the complex and gradual morphological modifications caused by the various degrees of incomplete metamorphosis in elateroid beetles.

The neotenic elateroid beetles are known for their extremely low dispersal ability, limited geographic ranges, and strong dependence on long-term climatically stable habitats including the humid tropics30,35,36. These attributes make neotenics excellent indicators of the long, uninterrupted evolutionary history of tropical forests. The Brazilian Atlantic rainforest is a complex biome formed by a network of ecosystems without clear limits, covering approximately 3,000 km of the eastern coast of Brazil37. It hosts one of the world’s most diverse tropical forest biotas with an exceptionally high level of species endemism38,39 but due to human activities leading to the intense landscape transformation during the last centuries, it is considered one of the most endangered biomes and belongs among the “hottest biodiversity hotspots” on the planet40–42. The region currently remains in constant degradation, mostly fragmented into isolated archipelagos of small forest fragments surrounded by open-habitat.
matrices, with only less than 10% of its original coverage\textsuperscript{25,41,42} and only about a third of its total extent preserved by conservation units\textsuperscript{24}. Nevertheless, even those remaining forest fragments still harbor a multitude of newly discovered animal taxa including both vertebrates\textsuperscript{43–45} and invertebrates\textsuperscript{46–51}. Currently, the largest and best-preserved portions of the Brazilian Atlantic rainforest are situated near the southern mountain ranges including Serras do Mar and da Mantiqueira\textsuperscript{42}, where all Jurasaidae known thus far were collected. Such a late discovery of a new beetle family, which is due to the cryptic lifestyle of its representatives bound to these ancient, stable habitats, further stresses the importance of the Brazilian Atlantic rainforest as a top priority area for nature conservation.

**Material and Methods**

**Taxon sampling and collecting sites.** A total of 120 specimens (21 adult males, 8 adult females, 4 pupae, 87 larvae) from two localities in the Brazilian Atlantic forest ecoregion in southeastern Brazil were studied (Supplementary Text). Larvae, pupae, adult males and females of the first two species were collected in the soil or using Malaise traps in the Serra dos Toledos reserve, and adult males of the third species were collected using Malaise traps in the Serra dos Órgãos National Park. For the detailed numbers of individuals under each species, corresponding localities and information on the collecting and rearing of specimens see the Supplementary Text. The Serra dos Toledos reserve is situated in the municipality of Itajubá, Minas Gerais state in the Mantiqueira mountain range. It comprises an area of 10.7 km\textsuperscript{2} with altitudes ranging from 1,100 to 1,800 m. The reserve was originally covered by dense ombrophilous and Araucaria forests, but nowadays it includes secondary forests and suffers from the deforestation caused by the surrounding agricultural and livestock activities\textsuperscript{52}. The Serra dos Órgãos National Park is situated in the municipality of Teresópolis, Rio de Janeiro state in the Serra do Mar mountain range. It covers an area of 105.3 km\textsuperscript{2} with altitudes ranging from 200 to 2,285 m. This conservation unit comprises one of the largest area of dense ombrophilous vegetation in Rio de Janeiro state, although approximately 45% are under anthropogenic land use, mainly pastures. Both Serra dos Toledos and Serra dos Órgãos are situated in the Atlantic Rainforest domain\textsuperscript{48,53}. The collection, maintenance, and shipping of specimens were conducted in accordance with the Brazilian environmental laws (Sisbio permits 53842–1, 53842–2, 53842–3, 43943–1, SiGen Shipping Registration R1CE59F, and authorization of the Municipal Government of Itajubá).

**Laboratory methods.** Whole genomic DNA was extracted using the E.Z.N.A. Tissue DNA Kit (Omega Bio-tek Inc. Norcross, USA) following standard protocol but with the overnight incubation and elution performed twice with 100 µl Elution buffer each. Amplifications were performed using Qiagen Multiplex PCR Plus Master Kit (Qiagen, Hilden, Germany) according to the manufacturer protocols. The PCR amplification conditions and list of primers used were reported in previous study\textsuperscript{24}. We sequenced two nuclear and two mitochondrial markers which have been widely used for Coleoptera and Elateriformia phylogenies\textsuperscript{9,13,22,55,56}, i.e., 18S rRNA (1842–1843 bp), 28S rRNA (627–629 bp), rrnL (505 bp), and cox1 mtDNA (723 bp). Selection of these markers, which are abundantly represented in GenBank, enabled us to test the position of the newly discovered lineage using extensive outgroups. The PCR products were purified using Exonuclease I and FastAP Thermosensitive Alkaline Phosphatase (Life Technologies, Darmstadt, Germany), and subsequently sent for sequencing to the commercial facility run by Macrogen Europe, Netherlands.

**Dataset assembly and phylogenetic analyses.** In addition to the field and laboratory observations, we associated the different developmental stages and sexes using the sequences from 10 specimens (three adult males, two adult females, one pupa, four larvae) representing three morphologically defined species from two localities (Table S3). We used the alignment method and the maximum likelihood analysis as described below.

To investigate the position of the newly discovered lineage, we merged the sequences of individuals from the three newly discovered species (one per species) with the 251-taxon dataset from the most comprehensive phylogenetic study of Elateriformia to date\textsuperscript{56}. Scirtiformia were used as an outgroup, since they represent basal phylogenetically important lineages for which some fragments were missing (Table S3). Forward and reverse sequences were edited and assembled using Geneious 7.1.7\textsuperscript{97} and multiple sequence alignments were generated using MAFFT 7.157\textsuperscript{88} at default parameters. Alignment of the length-invariable protein-coding cox1 sequences was checked by amino acid translation. Basic statistics and the pairwise sequence divergences based on uncorrected p-distance were calculated using MEGA 6.06\textsuperscript{39}. The resulting alignment included 4780 homologous positions (2338, 1128, 591 and 723 positions for 18S, 28S, rrnL and cox1, respectively), and contained 2134 conserved, 2408 variable, and 1872 parsimony informative characters.

Model and best-fit partitioning testing was carried out using a greedy algorithm in PartitionFinder 1.1.1\textsuperscript{40} under the corrected Akaike information criterion. Subsequent phylogenetic trees were generated using maximum likelihood analysis (ML) and Bayesian inference (BI). ML was conducted using RAxML 8.2.8\textsuperscript{98} with the default settings, partitioned by genes and codons, and bootstrapped with 1000 pseudoreplicates. BI was performed using MrBayes 3.2.6\textsuperscript{68} with the GTR + I + G model for most partitions (SYM + I + G for 28S rRNA) and partitioning by genes and codon positions as recommended by PartitionFinder. Four chains were run for 40 million generations using the Markov chain Monte Carlo method. Stationary phase and convergence were checked using Tracer 1.7.1\textsuperscript{104} and the first 20% of generations were discarded as burn-in. The posterior probabilities (PP) were calculated from the remaining trees. All analyses were run on the CIPRES web server\textsuperscript{45}.

We also evaluated the occurrence of substitution saturation in our data using Xia’s nucleotide substitution saturation test\textsuperscript{46} implemented in DAMBE 5.6.14\textsuperscript{47}. We analyzed separately each non-coding gene and each position of the protein-coding cox1 mtDNA using 10,000 replicates on the fully resolved sites and with the empirical proportion of invariant sites estimated from the data. Only little saturation was detected in our data except for the 3rd codon positions of the cox1 fragment, which were substantially saturated (Table S4). Therefore, we also performed the above-described ML and BI analyses with discarded third codon positions. Additionally, to test
the effect of extensive outgroup on the alignment accuracy, we removed from the dataset all terminals other than Elateroidea and Byrrhidae (outgroup), re-aligned the sequences of individual markers, and re-analyzed the concatenated dataset of 95 terminals using both ML and BI analyses.

Morphology. Male specimens were soaked in 10% KOH for 24 hours before dissection; female abdomens were boiled in KOH for dissection of genital tract; larvae soaked in KOH were extensively damaged, so they were dissected without treatment. Specimens and body parts were mounted in slides with Hoyer’s medium and examined under a stereomicroscope and compound microscope. Live specimens were also examined under a microscope to examine chaetotaxy and mouthparts. Drawings were made using a Zeiss Discovery V8 stereomicroscope with an attached camera lucida; photos were taken with a Canon A640 digital camera and a Axiocam camera attached to a Zeiss Imager A1 microscope. For examination with the scanning electron microscope, the specimens were rinsed with soapy water, gradually dehydrated in graded ethanol (70%, 80%, 90%, 99%), dried in a critical point dryer with liquid CO2, (Bal Tec SCD-050), mounted on aluminum stubs with double-sided conductive tape, and sputtered with gold. Images were taken using a Zeiss LEO 440 scanning electron microscope. Stacked photographs were combined using Helicon Focus (Helicon Soft Ltd., Ukraine). Body width of the specimens was measured at the widest part of the body, pronotal length at midline, and pronotal width at the widest part. The examined material is deposited at the Universidade Federal de Itajubá (UNIFEI), Museu de Zoologia da Universidade de São Paulo (MZUSP), the collection of Prof. J. A. P. Dutra at the Universidade Federal do Rio de Janeiro (DZJR), and the collection of Palacky University, Olomouc, Czech Republic (UPOL). Morphological terminology follows Lawrence et al.13, with the classification of Elateroidea follows Kundrata et al.15, with the subsequent changes made by other authors14,15,30.

Data availability
All newly generated sequences have been deposited in GenBank under the accession numbers MN578248–MN578256 and MN583350–MN583375 (Table S3). The datasets analyzed during the current study are available from the corresponding author on request.

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Author contributions

S.P.R., C.C. and R.K. conceived and designed the study. S.P.R. collected the samples and carried out the experiments. S.P.R. and C.C. conducted the morphological investigation and wrote the taxonomic supplementary information. K.K. and R.K. did the molecular laboratory work and carried out the molecular phylogenetic analyses. S.P.R. and R.K. performed the literature search and wrote the manuscript. All authors discussed the results, and reviewed and approved the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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