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Original article

Understanding the diversification and functional radiation of *Aporrectodea* (Crassiclitellata, Lumbricidae) through molecular phylogenetics of its endemic species

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ABSTRACT

The genus Aporrectodea includes some of the most conspicuous earthworm species, but its taxonomic history is among the most complex within the family Lumbricidae. Molecular phylogenetic studies have produced some advances by assigning former Aporrectodea species to other monophyletic clades and by detecting species level lineages within the cosmopolitan caliginosa-trapezoides complex. However, little attention has been devoted to endemic taxa of Aporrectodea such as Ap. rubra, Ap. arverna, Ap. gogna, Ap. balisa, Ap. velox, Ap. giardi voconca and Ap. longa ripicola. These earthworms (and additional populations of Ap. longa and Ap. nocturna) were included in a molecular phylogenetic framework in order to reconstruct the ancestral range of the genus, as well as to help understand its diversification within its native range and to perform a systematic revision. Species delimitation, ancestral area reconstruction and Bayesian inference of the phylogenetic relationships were performed using a large gene sequence (COI) dataset and a narrower dataset composed of 5 mitochondrial and nuclear markers. Phylogenetic position and species delimitation indicated that Ap. giardi voconca and Ap. longa ripicola constitute species-level entities not closely related to Ap. giardi or Ap. longa, and they were thus redescribed as Aporrectodea voconca stat. nov. and Aporrectodea ripicola stat. nov. Ancestral area reconstruction enabled location of the origin of Aporrectodea in the Auvergne-Rhône-Alps, in Southeastern France. The study findings provide some insight into the evolution of functional traits in this ecologically successful genus. Ap. rubra and Ap. arverna (small, reddish, epigeic/epianecic) and Ap. gogna (very large, dark, anecic) were recovered as the earliest branching taxa, suggesting a complex evolution of functional traits within this genus.

1. Introduction

Aporrectodea Orley, 1885 is one of the earthworm genera best known by non-specialized biologists and naturalists. This may be partly because some of the species (e.g. Aporrectodea caliginosa (Savigny, 1826), Aporrectodea trapezoides (Duges, 1828) and Aporrectodea rosea (Savigny, 1826), are amongst the most frequent and abundant in managed (such as crops, orchards and pastures) and natural habitats on a large range of soil types and climates around the world. Even though the origin of the genus is recognized to be in Europe, it includes cosmopolitan species (such as the aforementioned ones) which have been introduced in several countries and become invasive in temperate and mountain soils. Either in their native or non-native range, they play an important role in soil functioning as demonstrated by numerous studies [1–3].

Aporrectodea has a particularly conflictive taxonomic history, inextricably linked over the last 140 years to that of Allolobophora Eisen, 1873, another remarkable genus in the family Lumbricidae. Initially, Örley [4] included Enterion chloroticum Savigny, 1826 and Lumbricus trapezoides Dugès, 1828 within Aporrectodea, without explicitly establishing a type species. The later choice [5] of Enterion chloroticum as the type species of Allolobophora (due to no original type species designation) led to the automatic establishment of Lumbricus trapezoides as the

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type species of Aporrectodea. Since then, several species have been transferred from one genus to the other, perhaps as a result of the imprecision of each of their original diagnoses. Additionally, Marcel Bouché [6] created the genus Nicodrilus Bouché, 1972 for Enterion caliginosum Savigny, 1826 (synonymous to Ap. trapezoides according to Bouché) and its most closely related species. This was justified by the fact that Orley's (not Dugès') description of Ap. trapezoides was incomplete and probably based on Hungarian material, thus arguably making it an unsuitable type species [7]. However, this argument was not accepted by many taxonomists (e.g. Gates [8], Csuzdi and Zicsi [9] and Reynolds [10]), who considered Aporrectodea a valid genus and Nicodrilus to be its junior synonym. Nonetheless, Marcel Bouche's intuition was sharp, and he grouped a very homogeneous set of species within Nicodrilus, while leaving other species within Allolobophora sensu lato. Later, he transferred some of these species to the new genus Koinodrilus Qiu and Bouché, 1998, the most conspicuous being Ap. rosea. However, the differences in the diagnoses of Nicodrilus and Koinodrilus (the position of the nephridial pores) are so small that Koinodrilus is also not widely accepted as a valid genus. To further complicate the issue, the genus Eiseniona Omodeo, 1956 was created to include several Central/Eastern European species such as Eiseniona handlirschi (Rosa, 1897) and Eiseniona sineporis (Omodeo, 1952); however, authors such as Blakemore [11] considered this genus a junior synonym of Aporrectodea and retained the species within the latter.

As with other systematic conundrums, the advent of molecular phylogenetics contributed to disentangling the Aporrectodea/Allolobophora knot. Domínguez et al. [12] recovered a well-supported clade including Ap. caliginosa, Ap. trapezoides and other species assigned by Marcel Bouché to Nicodrilus, while other species (Ap. rosea, Aporrectodea georgii (Michaelsen, 1890), Aporrectodea jassyiensis (Michaelsen, 1891)) were found to be unrelated. On the other hand, an Allolobophora clade including Allolobophora chlorotica (Savigny, 1826), Allolobophora molleri Rosa, 1889, Allolobophora moebii Michaelsen, 1895 and Aporrectodea dubiosa Orley, 1881 was also strongly supported. Martínez-Navarro et al. [13] found that Aporrectodea icterica (Savigny, 1826) also belonged to a redefined Allolobophora, and that Aporrectodea pseudoantipai Qiu and Bouché, 1998 was closely related to Eiseniona handlirschi: the latter two appeared closely related to Aporrectodea sensu stricto, but it was unclear whether they should be considered a separate genus (which would correspond to Eiseniona).

Molecular phylogenetic studies have also focused on the *Ap. caliginosa* species complex itself [14,15]. Their results showed that, not only are *Ap. caliginosa* and *Ap. trapezoides* not the same species, but they both contain several species-level cryptic lineages. These lineages display different geographical ranges and dispersal histories [16,17]. The difficulty in genotyping old (or missing) type specimens or obtaining topotypes corresponding to a single lineage has so far hindered the assignment of the names *caliginosa* and *trapezoides* to lineages (and the taxonomic description of the others).

Study of those cosmopolitan representatives of Aporrectodea have highlighted the need to sample the native ranges of the species to pinpoint their geographic origin [17]. However, the endemic species of Aporrectodea (found within the putatively native range of the genus, France) have received little attention regarding molecular phylogenetics and phylogeography. The following species could reveal the centre of diversification of the genus and explain its evolutionary origin before human-mediated dispersal blurred the biogeographic patterns: Aporrectodea rubra (Vedovini, 1969), Aporrectodea arverna (Bouché, 1969), Aporrectodea gogna (Bouché, 1972), Aporrectodea balisa (Bouché, 1972) and Aporrectodea velox (Bouché, 1967). Interestingly, these species provide a wider outline of the functional and ecological diversity of the genus, including both relatively small, red pigmented forms (similar to Lumbricus Linnaeus, 1758) and large and strongly pigmented forms (reminiscent of Scherotheca Bouché, 1972). In addition, several endemic subspecies such as Aporrectodea rubra acidicola (Bouché 1972) (known from Provence), Aporrectodea giardi voconca (Bouché, 1972) (known

from Auvergne-Rhône-Alpes) and *Aporrectodea longa ripicola* (Bouché, 1972) (known from the Paris Basin up to the north of the Rhône Basin) have been described, although their taxonomic status remains unconfirmed.

The aim of this study was to include the aforementioned taxa for the first time in the multilocus phylogenetic framework of Lumbricidae, together with additional representatives of *Aporrectodea longa* (Ude, 1885) and *Aporrectodea nocturna* (Evans, 1946) comprising their native range. This will allow to: i) reconstruct the ancestral range of the genus *Aporrectodea*, ii) understand the diversification of the genus within its native range, and iii) perform a systematic revision of some of its taxa.

2. Material and methods

2.1. Specimens, sampling and morphological description

Specimens included in this study were collected between October and December 2021 in Auvergne-Rhône-Alpes, Bourgogne-Franche-Comté, Grand Est and Provence-Alpes-Côte d'Azur (France). The specific localities are listed in Table 1.

The earthworms were collected by digging and hand-sorting the soil, then rinsed with water and fixed in 70%. Once immobilized, the specimens were immediately transferred to 100% ethanol for further molecular analyses. Species classification and morphological diagnoses were conducted using the same set of external and internal morphological characters reported by Ref. [6].

The following main external morphological characters were considered: mean length, mean number of segments, mean weight (all three measured in adult, complete, ethanol fixed specimens), pigmentation, type of prostomium, position of papillae, position of first dorsal pore, position of spermathecal pores, position of clitellum and position of *tubercula pubertatis*. The following main internal anatomical characters were considered: position of oesophageal hearts, position and morphology of calciferous glands, position of crop, position of gizzard, type of typhlosole, shape of nephridial bladders, number and position of seminal vesicles, and number and position of spermathecae.

2.2. DNA sequencing and phylogenetic analysis

Two datasets differing in specimen and gene coverage were assembled.

Dataset 1 consisted of a phylogeographic sample of individuals from different populations of the target species (*Ap. rubra, Ap. rubra acidicola, Ap. arverna, Ap. gogna, Ap. balisa, Ap. velox, Aporrectodea giardi* (Ribaucourt, 1901), *Ap. giardi voconca, Ap. nocturna, Ap longa, Ap. longa ripicola*) from different parts of their range (Table 1, Supplementary Table 1, Supplementary Fig. 1).

For each locality sampled, we selected up to five adult individuals per species for DNA barcoding. Small ventral integument tissue samples were assembled in 96-well plates and shipped to the Centre for Biodiversity Genomics at the University of Guelph (Ontario, Canada) for processing. After the total genomic DNA was extracted using a CTABbased approach, the standard DNA barcode for animals [18] - a 658bp fragment of mitochondrial gene cytochrome c oxidase subunit 1 (COI) - was amplified using the primers C_LepFolF-C_LepFolR [19]. Sequencing reactions were carried out with the same primer pair, and the products were subjected to clean-up with PureSeq-MP (Aline Biosciences, Woburn, USA) before Sanger sequencing in a DNA sequencer (ABI 3730XL). Consensus sequences from automatically assembled contigs (subsequently reviewed and manually edited when needed) were uploaded to BOLD (www.boldsystems.org), along with trace files, specimen data and images. The DNA sequences, together with metadata are publicly available in the (to be provided) dataset.

Additional sequences for the species *Ap. giardi, Ap. nocturna,* and *Ap. longa* were downloaded from BOLD (www.boldsystems.org, Supplementary Table 1), discarding misidentifications based on their clustering

Table 1

Sa	amplin	ig local	lities of	the	species	of 1	Aporr	rectod	ea u	inder	stud	y.
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Sample ID	Species	Country	State/ Province	Latitude	Longitude
DFM- 0975 - 0979	Aporrectodea gogna	France	Auvergne- Rhône-Alpes, Ain	46.216	5.340
DFM- 0980 - 0981	Aporrectodea giardi voconca	France	Auvergne- Rhône-Alpes, Ain	46.216	5.340
DFM- 0982 - 0984	Aporrectodea nocturna	France	Auvergne- Rhône-Alpes, Ain	45.966	5.174
DFM- 0985 - 0986	Aporrectodea longa ripicola	France	Auvergne- Rhône-Alpes, Ain	45.966	5.174
DFM- 0987	Aporrectodea giardi voconca	France	Bourgogne- Franche- Comté, Côte- d'Or	47.062	4.760
DFM- 0988 - 0991	Aporrectodea longa ripicola	France	Bourgogne- Franche- Comté, Côte- d'Or	47.017	5.127
DFM- 0992 - 0996	Aporrectodea gogna	France	Bourgogne- Franche- Comté, Doubs	47.170	6.168
DFM- 0997	Aporrectodea longa	France	Bourgogne- Franche- Comté, Doubs	47.170	6.168
DFM- 0998 - 1002	Aporrectodea velox	France	Bourgogne- Franche- Comté, Haute- Saône	47.836	6.359
DFM- 1003 - 1007	Aporrectodea velox	France	Grand Est, Vosges	47.988	6.512
DFM- 1009 - 1010	Aporrectodea longa	France	Grand Est, Haut-Rhin	47.799	7.026
DFM- 1011	Aporrectodea longa ripicola	France	Grand Est, Haut-Rhin	47.799	7.026
DFM- 1012 - 1015	Aporrectodea rubra	France	Provence- Alpes-Côte d'Azur, Vaucluse	43.963	5.337
DFM- 1021 - 1025	Aporrectodea balisa	France	Provence- Alpes-Côte d'Azur, Vaucluse	43.965	5.323
DFM- 1026 - 1030	Aporrectodea longa ripicola	France	Provence- Alpes-Côte d'Azur, Vaucluse	43.903	4.968
DFM- 1046	Aporrectodea rubra	France	Provence- Alpes-Côte d'Azur, Var	43.181	6.345
DFM- 1067 - 1071	Aporrectodea nocturna	France	Provence- Alpes-Côte d'Azur, Var	43.587	6.538

with confidently identified specimens of the same species.

Putative species were delimited using ASAP [20] based on simple distances (p-distances), and the model with the lowest ASAP score was selected.

An ultrametric phylogenetic tree was obtained in BEAST 1.10.4 [21]. jModelTest v. 2.1.3 [22] was used to select the best-fit evolutionary model by applying the Akaike information criterion (AIC [23]) and the Bayesian information criterion (BIC [24]). GTR + I + G was selected as the best-fit evolutionary model. The analysis was conducted using a constant coalescent model and an uncorrelated lognormal relaxed clock. A relative calibration was specified for the root of the tree as a normal prior with mean = 1 and standard deviation = 0.05. A uniform distribution with an initial value of 0.002 and a range of 0.00005–0.02 was specified through the ucld. mean parameter, and a uniform distribution with an initial value of 0.10 ranging from 0 to 10 was specified for the ucld. stdev parameter. Fifty million generations were specified for the Monte-Carlo Markov chain, sampling every 5,000th generation. The log file was visualized in Log Tracer v. 1.7 [25] to check for convergence and effective sampling sizes (ESS) greater than 100. The final tree was generated by TreeAnnotator v.1.10.4 [21]. with a burn-in of 2000 trees.

Ancestral area reconstruction was performed in RASP 4 [26] using the previous ultrametric tree as input, and the most suitable model was chosen using BioGeoBEARS (DIVALIKE + j) and allowing a maximum of three areas for each node. The areas considered were coded as follows: A (Bretagne), B (Normandie), C (Ile de France), D (Grand Est), E (Bourgogne-Franche-Comté), F (Nouvelle Aquitaine), G (Occitanie), H (Auverne-Rhône-Alps), I (Provence-Alpes-Côte d'Azur), J (Spain, United Kingdom, Norway, Sweden, Denmark).

Dataset 2 included one representative of the different species-level taxa within *Aporrectodea* (including the information from the species delimitation analysis) with additional molecular markers.

Total genomic DNA was extracted using the DNeasy Blood & Tissue Kit (Qiagen), from ventral integument samples of approximate size 5×5 mm, from two representative specimens of each putative species. Regions of the nuclear 28S rRNA and mitochondrial 16S rRNA, NADH dehydrogenase (ND1) and 12S rRNA were amplified by polymerase chain reaction (PCR), with the primers and conditions described in Refs. [14,27] (Suppl. Table 2). PCR products were purified and sequenced by the C.A.C.T.I Genomics service (University of Vigo) and by Macrogen Spain. The DNA sequences are available in Genbank, under accession numbers (to be provided).

Previously existing sequences of *Aporrectodea* sensu stricto, as well as *Aporrectodea limicola* (Michaelsen, 1890), *Ei. handlirschi* and *Ap. pseudoantipai* and representatives of other crown Lumbricidae genera were downloaded from Genbank and included in the dataset as references.

Sequences were aligned using MAFFT v.7 [28] with default settings and concatenated (as no major incongruence existed between mitochoindrial and nuclear markers) using BioEdit [29] resulting in a matrix of 3199 bp. The best fitting evolutionary model for each partition was selected using jModelTest v. 2.1.3. GTR + I + G was selected as the best-fit evolutionary model for COI, 28S, 12S and ND1 and HKY + I + G was selected for 16S.

Bayesian inference of the phylogeny was estimated using MRBAYES v.3.1.2 [30] as implemented in CIPRES Science Gateway V. 3.3. Parameters were set to 50 million generations and sampled every $5,000^{\text{th}}$ generation (10,000 trees). The best fitting evolutionary models were specified for each partition. Two independent runs were performed, each with four chains, and 20% of the trees were discarded as burn-in. The remaining trees were combined and summarized on a 50% majority-rule consensus tree.

3. Results

3.1. Species delimitation and phylogeny

The clustering hypothesis obtained by ASAP with the lowest ASAP score (5.5) consisted of 13 putative species (Supplementary Fig. 2). *Aporrectodea nocturna, Ap. arverna, Ap. balisa, Ap. gogna* and *Ap. velox* were each recovered as a single species. *Aporrectodea longa* was split in two putative species, and *Ap. longa ripicola* was recovered as two separate putative species (one of them corresponding to *Ap. longa ripicola viridis*). *Aporrectodea giardi* was recovered as a single species. Finally, *Ap. giardi voconca* was recovered as a separate putative species. Finally, *Ap. rubra* and *Ap. rubra acidicola* were shown as different putative species.

The Bayesian (Fig. 1) phylogenetic tree recovered a monophyletic and well supported *Aporrectodea* sensu stricto comprising all of the studied species of *Aporrectodea* except *Ap. limicola* (closely related to *Eisenia, Eisenoides, Bimastos* and *Scherotheca*), *Ap. rosea* and *Ap. pseudoantipai*. The latter clustered with *Eiseniona handlirschi* in a strongly supported clade, more closely related to *Lumbricus* than to *Aporrectodea*.



Fig. 1. Phylogenetic relationships between *Aporrectodea* sensu stricto species and their closest relatives obtained by Bayesian inference based on the concatenated sequence of five molecular markers (COI, ND1, 16 S, 12S and 28S). Support (posterior probability) values are shown above the corresponding branches. The taxa added in this study are shown in bold.

Aporrectodea rubra and Ap. arverna were recovered as the earliestbranching clades within Aporrectodea, followed by Ap. gogna. The next clade comprised Ap. caliginosa, the closely related Ap. tuberculata and Aporrectodea ripicola stat. nov., the latter clearly separated from Ap. longa. One of the last two clades included Aporrectodea voconca stat nov. (neatly separated from Ap. giardi), Ap. velox, Ap. balisa and one of the lineages of Ap. trapezoides (plus a second one according to Maximum Likelihood analysis); the other included Ap. giardi and Ap. longa, and an Ap. nocturna clade with strong internal divergence.

3.2. Ancestral area reconstruction

The topology of the ultrametric tree on which the Ancestral Area Reconstruction was based (Fig. 2) differed slightly from the Bayesian and ML tree in regard to the sister relationship of *Ap. rubra* and *Ap. gogna* (instead of *Ap. arverna*) and *Ap. velox* and *Ap. balisa* (instead of *Ap. voconca stat. nov.*). This was probably due to exclusion of the cosmopolitan species *Aporrectodea caliginosa/Aporrectodea tuberculata* and *Aporrectodea trapezoides*).

The oldest divergence was between *Ap. rubra* and *Ap. gogna*, and between both *Ap. voconca stat. nov.* and *Ap. nocturna* and their closest relatives. The divergence between *Ap. balisa-Ap. velox* and *Ap. giardi-Ap. longa* was comparatively recent.

Aporrectodea rubra, Ap. ripicola stat. nov., Ap. nocturna and Ap. longa displayed deeper internal branches than the other species of which several populations were sampled.

Ancestral area reconstruction (Fig. 2) estimated the most likely area of origin for the common ancestor of the studied *Aporrectodea* to be Auvergne-Rhône-Alps (Southeastern France). The ancestral area for most of the clades and species corresponded to Southeastern-Eastern France (Bourgogne-Franche-Comté, Auvergne-Rhône-Alps, ProvenceAlpes-Côte d'Azur), suggesting most of the diversification of the genus occurred in this region. The exception was the *Ap. giardi-Ap. longa* clade, whose ancestral area was estimated to be in north-central France (Ile de France). For *Ap. longa*, the ancestral area was estimated to be the surrounding countries (Spain, United Kingdom, Norway, Sweden, Denmark), but this is probably due to an overrepresentation of populations from outside of France.

4. Discussion

4.1. Systematic implications

The wealth of subspecies and varieties described by Bouché [6] for *Nicodrilus* (here *Aporrectodea*) already suggested a greater diversity of the genus than is usually recognized. Phylogenetic inference and species delimitation analyses supported that the subspecies *giardi voconca* and *longa ripicola* are not closely related to their parent taxa *Ap. giardi* and *Aporrectodea longa* thus should be elevated to species status.

4.1.1. Proposed taxonomic changes
Phylum Annelida Lamarck, 1802.
Subphylum Clitellata Michaelsen, 1919.
Class Oligochaeta Grube, 1850.
Superorder Megadrili Benham, 1890.
Order Crassiclitellata Jamieson, 1988.
Family Lumbricidae Rafinesque-Schmaltz, 1815.
Tribe Lumbricini Qiu and Bouché 1998.
Genus Aporrectodea Orley, 1885.

4.1.1.1. Aporrectodea voconca stat. nov. (Bouché, 1972)



Fig. 2. Ancestral area reconstruction for *Aporrectodea* sensu stricto populations from France and surrounding countries (excluding the cosmopolitan *Aporrectodea caliginosa/Aporrectodea tuberculata* and *Aporrectodea trapezoides*). Only the most likely reconstruction for each node is shown for key nodes. The areas considered were coded as follows: A (Bretagne), B (Normandie), C (Ile de France), D (Grand Est), E (Bourgogne-Franche-Comté), F (Nouvelle Aquitaine), G (Occitanie), H (Auvergne-Rhône-Alps), I (Provence-Alpes-Côte d'Azur), J (Spain, United Kingdom, Norway, Sweden, Denmark).

Nicodrilus terrestris voconcus Bouché, 1972: 321.

Aporrectodea giardi voconca Blakemore, 2008: 21

4.1.2. Redescription

4.1.2.1. Diagnosis. Specimens of Aporrectodea voconca can be distinguished from other known species of Aporrectodea by the position of the clitellum in segments XXVI-XXXVI, spectacle-shaped tubercula pubertatis in segments XXXII-XXXIV, strongly developed genital papillae in segments X, XII (sometimes in XI,XII) and a partially pennate typhlosole which changes to trifid around segment XXVII (Table 2).

4.1.2.2. Etymology. The name *voconca* refers to the Gallic tribe of the Vocontii, which inhabited the lower Rhône valley.

4.1.3. Material examined

Holotype. France • Adult; Auvergne-Rhône-Alpes, Ain, Saint-Sorlinen-Bugey 26-Feb-1968; Marcel Bouché leg. ; voucher 61-401-212;

deposited in MNHN.

Additional material. France • 3 adult specimens; Auvergne-Rhône-Alpes, Ain, Jasseron; latitude/longitude: 46.216/5.340; elevation: 373 m asl; 30-Oct-2021; T. Decaëns, D. Fernández Marchán leg. (2 specimens) BOLD Sample ID: DFM-0980, DFM-0981; deposited in ECO&SOLS. Bourgogne-Franche-Comté, Côte-d'Or, Bouze-les-Beaune; latitude/longitude, 47.062, 4.76; elevation, 490 m asl; 30-Oct-2021; T. Decaëns, D. Fernández Marchán leg. (1 specimen); BOLD Sample ID: DFM-0987; deposited in ECO&SOLS.

4.1.4. Morphological description

4.1.4.1. External morphology. Body pigmentation brown with anteroposterior and dorso-ventral gradients (Fig. 3).

Mean length (fixed specimens) 13 cm (12.1–13.7 cm, n = 2 adults); body cylindrical in cross-section, slightly flattened tail; mean number of segments 207 (n = 2 adults; 192 segments in the holotype). Mean weight (fixed specimens): 2.65 g (2.27–3.03 g, n = 2 adults). Prostomium epilobous, closed. Transverse furrows from segment VII or VII. First dorsal

Table 2

Morphological characters of *Aporrectodea voconca* stat. nov, *Aporrectodea ripicola* stat. nov. and the other species of *Aporrectodea* sensu stricto. Average length in centimeters, average weight in grams. 1st D.P: First dorsal pore. G.P: Genital papillae. T.P: Tubercula pubertatis. * Obtained from the minimum and maximum values shown in Bouché [6].

	Av. length	# segments	Av. weight	1st D.P	G.P	Clitelum	T.P.	Typholosole
Aporrectodea voconca stat. nov	13	192–222	2.65	11/12	10,12 or 11,12	26–36	32-34 spectacle-shaped	Pennate (up to 28), trifid
Aporrectodea giardi	20*	108–203	2.4	(10/11) 11/12	(9)10,11,12	(1/n 26)27–35 (2/3 36)	32-34 spectacle-shaped	Pennate (up to 32), trifid
Aporrectodea ripicola stat. nov.	10.5	155–173	1.26	12/13, 13/14	(9)10,11	1/2 28 (28)-1/2 35 (35)	32-1/2 34 oval shaped	Pennate (up to 40+), simple
Aporrectodea longa	15*	160-200	1.65	12/13	9–11	(1/2 27)28–35 (1/2 36)	32-34 gutter-shaped	Pennate (up to 32), trifid
Aporrectodea nocturna	13.5*	180–250	1.68	(8/9)9/ 10	(9)10-12	(1/y 27)28–34 (1/2 35)	31-33 spectacle-shaped	Trifid-pentafid
Aporrectodea velox	23.6	483	7	(9/10) 10/11	(9)10-12	(1/n 24)25–1/2 36 (36)	1/n 30-34 gutter-shaped	Pennate (up to 32), bifid T
Aporrectodea gogna	17.3	215-240	6.45	11/12	11,13,14	24–37	1/2 31–35 gutter-shaped	Pennate
Aporrectodea balisa	21.3	280-315	5.4	11/12	9–11	(1/n 26)27–35 (1/n 36)	(1/n 29)1/2 30–1/2 34 (34) band- shaped, ventral protuberances in 31, 33	Pennate
Aporrectodea rubra	5.5	95–115	0.5	(4/5,5/ 6)6/7	(9,10,11,12)	(25)1/3 25–1/2 32 (1/a 33)	(1/y 26)1/3 27–30 (1/n 31) gutter- shaped	Pennate
Aporrectodea arverna	5.9	115–126	0.8	(9/10) 10/11	10,11 (12)	(25)1/n 26–34 (1/y 35)	(1/2 31)2/3 31–33 band-shaped, ventral protuberances in 1/4 32, 1/4 33	Pennate
Aporrectodea caliginosa	5.5	130–165	0.4	(9/10) 10/11	9–11	(26)27–34 (1/a 35*)	31-33 spectacle-shaped	Pentafid
Aporrectodea trapezoides	5.9	130–170	0.6	(8/9)9/ 10	9–11	27–34	31-33 arc-shaped band	Pennate (up to 32), trifid



Fig. 3. External morphology of *Aporrectodea voconca* stat. nov. (a,b) and *Aporrectodea ripicola* stat. nov. (c,d) in lateral (a,c) and ventral (b,d) views; typhlosole of e) *Aporrectodea voconca* stat. nov. f) *Aporrectodea ripicola* stat. nov. and g) *Aporrectodea longa*. Scale bar: 2 cm.

pore at intersegmental furrow 11/12. Nephridial pores inconspicuous aligned in *C* close to *b*. Spermathecal pores at intersegmental furrows 9/10 and 10/11 in *c*. Male pores in segment XV, surrounded by a well-developed porophore. Female pores on the posterior part of segment XIV, inconspicuous. Clitellum saddle-shaped in segments XXVI-XXXVI. Spectacle-shaped tubercula pubertatis in segments XXXII-XXXIV. Chaetae closely paired. Strongly developed chaetophores/genital papillae in segments X, XII (sometimes in XI,XII).

4.1.4.2. Internal anatomy. Septa 6/7–8/9 strongly thickened, 9/10–11/ 12 slightly thickened. Lateral hearts in segments VI-XI. Calciferous glands in segments X-XIV, with paired diverticula in X. Crop in segments XV-XVI, gizzard in segments XVII-XVIII. Typhlosole pennate (composed of transverse folds resembling ribs) from the start to segments XXVII-XXVIII, trifid (composed of three longitudinal lamellae) from that point to the end (Fig. 3). Male sexual system holandric, testes and funnels (not enclosed in testes sacs, but with sperm present) located ventrally in segments 10 and 11. Four pairs of reniform seminal vesicles in segments IX, X, XI and XII, with the latter two pairs being larger. Ovaries and female funnels in segment XIII, ovarian receptacles (ovisacs) in segment XIV. Two pairs of globular spermathecae in segments IX and X. Nephridial bladders inverted J shaped, proclinate in anterior and posterior segments.

4.1.4.3. Aporrectodea ripicola stat. nov. (Bouché, 1972)

Nicodrilus longus ripicola Bouché, 1972: 325.

Aporrectodea longa ripicola Blakemore, 2008: 22

4.1.5. Redescription

4.1.5.1. Diagnosis. Specimens of Aporrectodea ripicola can be

distinguished from other known species of *Aporrectodea* by the position of the clitellum in segments 1/2 XXVIII (XXVIII)-1/2 XXXV (XXXV), oval-shaped tubercula pubertatis in segments XXXII-1/2 XXXIV, partially pennate typhlosole which changes to simple beyond segment XL and frequent green pigmentation (Table 2).

4.1.5.2. *Etymology*. The name *ripicola* refers to the preference of this earthworm for moist soils.

4.1.5.3. Material examined. Holotype. France • Adult; Centre-Val de Loire, Indre et Loire, Loches 07-Nov-1968; Marcel Bouché leg. Voucher 64-1021-0231; deposited in MNHN.

Additional material. France • 12 adult specimens; Provence-Alpes-Côte d'Azur, Vaucluse, Caumont-sur-Durance; latitude/longitude: 43.903/4.968; elevation: 49 m asl; 25-Nov-2021; T. Decaëns, D. Fernández Marchán, Y. Capowiez leg. (5 specimens); BOLD Sample ID: DFM-1026, DFM-1027, DFM-1028, DFM-1029, DFM-1030; deposited in ECO&SOLS. Auvergne-Rhône-Alpes, Ain, Crans; latitude/longitude, 45.966/5.174; elevation, 299 m asl; 29-Oct-2021; T. Decaëns, D. Fernández Marchán leg. (2 specimens); BOLD Sample ID: DFM-0985, DFM-0986; deposited in ECO&SOLS. Bourgogne-Franche-Comté, Coted'Or, Pouilly-sur-Saone; latitude/longitude, 47.017/5.127; elevation, 177 m asl; 30-Oct-2021; T. Decaëns, D. Fernández Marchán leg. (4 specimens); BOLD Sample ID: DFM-0988, DFM-0989, DFM-0990, DFM-0991; deposited in ECO&SOLS. Grand Est, Haut-Rhin, Bourbach-le-Haut; latitude/longitude, 47.799/7.026; elevation, 572 m asl; 03-Nov-2021; T. Decaëns, D. Fernández Marchán leg. (1 specimen); BOLD Sample ID: DFM-1011; deposited in ECO&SOLS.

4.1.6. Morphological description

4.1.6.1. *External morphology*. Body pigmentation faint brown to brown with antero-posterior and dorso-ventral gradients, sometimes with a green hue which is more intense in the tail region (Fig. 3).

Mean length (fixed specimens) 10.5 cm (9.2–11.9 cm, n = 3 adults); head and clitellum cylindrical in cross-section, tail strongly flattened dorso-ventrally; mean number of segments 167 (n = 3 adults; 173 segments in the holotype). Mean weight (fixed specimens): 1.26 g (0.99–1.60 g, n = 3 adults). Prostomium epilobous, closed. Transverse furrows starting at segment V or VI. First dorsal pore at intersegmental furrow 12/13 or 13/14. Nephridial pores inconspicuous aligned in *C* close to *b*. Spermathecal pores at intersegmental furrows 9/10 and 10/ 11 in *c*. Male pores in segment XV, surrounded by a well-developed porophore. Female pores on the posterior part of segment XIV, inconspicuous. Clitellum saddle-shaped in segments 1/2 XXVIII (XXVIII)-1/2 XXXV (XXXV), oval-shaped tubercula pubertatis in segments XXXII-1/2 XXXIV. Chaetae closely paired. Chaetophores/genital papillae or glandular areas in segments (IX) X, XI.

4.1.6.2. Internal anatomy. Septa 6/7–11/12 slightly thickened. Lateral hearts in segments VI-XI. Calciferous glands in segments X-XIV, with paired diverticula in X. Crop in segments XV-XVI, gizzard in segments XVII-XVIII. Typhlosole pennate (composed of transversal folds resembling ribs) from the start to segments XL or beyond, simple from that point (Fig. 3). Male sexual system holandric, testes and funnels (not enclosed in testes sacs, but with sperm present) located ventrally in segments 10 and 11. Four pairs of reniform seminal vesicles in segments IX, X, XI and XII, with the latter two pairs being larger. Ovaries and female funnels in segment XIII, ovarian receptacles (ovisacs) in segment XIV. Two pairs of globular spermathecae in segments IX and X, intraseptal. Small nephridial bladders inverted J shaped, proclinate in anterior and posterior segments.

4.1.6.3. General remarks. Representative specimens of Ap. rubra acidicola and A. ripicola stat nov. var. viridis were separated from their parent species by species delimitation analysis. However, in the absence of clear morphological differences (*Ap. rubra acidicola* mainly differed in its soil preferences and *Ap. ripicola viridis* in its stronger green pigmentation), it is more reasonable not to elevate them to species. In addition, *viridis* was originally a variety described after 1960, and according to the International Code of Zoological Nomenclature the name is considered unavailable.

A remarkably high intraspecific genetic diversity was observed within *Ap. nocturna*, which had already been hinted by the local scale comparative phylogeography of Marchán et al. [31]. However, species delimitation did not separate the intraspecific lineages into putative species, which suggests that they do not constitute cryptic species.

The opposite case was found for *Ap. longa*, with two deeply divergent lineages which were delimited as different putative species. Those lineages were already described by Martinsson et al. [32], who found the existence of a barcoding gap but no evidence of cryptic speciation from nuclear markers (ITS2 and Histone 3).

The different diversification patterns of the two aforementioned species may simply be due to a more incomplete sampling of the range of *Ap. longa*, resulting in a genetic gap between the observed lineages. However, other evolutionary causes (such as different dispersal-extinction histories or different isolation patterns) may have been involved and appear worthy of further exploration.

Aporrectodea longa and Ap. nocturna, which have previously been suggested to form a single species according to molecular analyses [14], appeared as well-separated taxa in our analyses. This may be due to the increased geographical sampling and more complete coverage of their intraspecific diversity. However, a few individuals identified as Ap. longa appeared nested within Ap. nocturna. Interestingly, Bouché [6] described Ap. longa var. ambiguus with a very similar morphology to Ap. nocturna. Our results suggest that those earthworms were actually morphs of Ap. nocturna which resembled Ap. longa.

The typhlosole had previously been described as pennate ("penné") for most of the species of Aporrectodea sensu stricto in Bouché [6], except for Ap. giardi and Ap. caliginosa (bifid). For Aporrectodea longa, it was described as "pennate, sometimes organized in 2 or three lamellae", while for Ap. nocturna it was described to be "pennate or formed by 2, 3 or 5 longitudinal lamellae". Detailed observation of the typhlosole of representatives of those species has revealed a more complex scenario: Ap. giardi, Ap. longa, Ap. voconca stat. nov., Ap. ripicola stat. nov., Ap. velox and Ap. trapezoides (Eurosiberian lineage) possess pennate typhlosoles (composed of transverse folds resembling ribs) which change into three to one longitudinal lamellae at somewhat specific positions. For Ap. voconca stat. nov. and Ap. ripicola stat. nov, in which this transition is displaced forwards or backwards relative to the other species, this character can be considered of taxonomic utility. Within a genus with a limited amount of taxonomically informative characters, this discovery might help to resolve the troublesome systematics of the different lineages of Ap. caliginosa and Ap. trapezoides. Furthermore, the division of the typhlosole in two types of organization suggests that there may be some regionalization and functional specialization of the intestine in the genus Aporrectodea, a possibility which has not been previously explored and that could be related to the ecological flexibility of these species.

Aporrectodea is one of the most conspicuous genera within human managed habitats in its native range (Europe) and also in temperate latitudes al around the globe. Its complicated, unclear taxonomy has hindered soil ecology and applied research for decades, with results which are not really cross-comparable due to the different treatment of the taxa. The taxonomic advances in this work should pave the way for more rigorous inclusion of those species into the knowledge of soil functioning, with only the status of a few species remaining to be solved (see 4.3.).

4.2. Biogeographic and evolutionary implications

Ancestral area reconstruction based on the endemic *Aporrectodea* species restricted to France enabled location of the origin of this successful and globally distributed genus in the Auvergne-Rhône-Alpes, in Southeastern France. The peri-Alpine region appears to have been an important area in the diversification and evolution of Lumbricini (*Lumbricus* [33], *Allolobophora* [13]), which is supported by this result. The closely related genus *Scherotheca* has been suggested to have originated between Provence and Corsica [34], although more complete sampling of the Provençal species will be necessary to confirm this possibility. These non-overlapping geographical origins could explain the surprisingly parallel evolution of both genera and their distinct biogeographical patterns.

It is worth mentioning that Ancestral Area Reconstruction methods heavily depend on sampling extent, and that the possibility of the actual ancestral area not having been sampled (and thus not being taken into account in the analysis) is never 0. This is unlikely for the whole genus *Aporrectodea*, as the earliest branching taxa (which contain the most information about the ancestral area) possess very restricted ranges; however, this could be the case for other more widely distributed species (such as Ap. *giardi* or *Ap. longa*).

Biogeographic reconstruction also showed that most of the diversification of the genus Aporrectodea occurred in the area of origin, before the genus expanded across the rest of France and neighbouring countries. However, a second centre of diversification appears to be located in the North of France (Ile de France), where the common ancestor of Ap. giardi and Ap. longa was inferred to have lived. Interestingly, this region is further north than the limit usually reconstructed for permafrost during the Last Glacial Maximum (LGM, ca. 33,000-15,000 years ago [35]): it is usually believed that earthworm species and populations occurring further north than this limit became extinct, and those areas were repopulated at a later date [6]. The very high genetic diversity detected in this northern area (including two distinct lineages of Ap. longa) suggests two hypotheses: either the species diversified in-situ and survived in glacial refuges, or northern France was recolonized independently from the south by separate lineages. This surprising result may be explained by the limited sampling of Ap. giardi (and to a lesser extent Ap. longa) ranges. It is also possible that the inclusion of several populations of these species from a region with very high human activity (Ile de France) could have distorted the results due to the likely repeated transfer of specimens from distant regions. A wider, denser coverage of the ranges of Ap. giardi and Ap. longa could offer further insight into these hypotheses.

In relation to this same conundrum, Ap. gogna and Ap. velox are narrow endemics whose range is restricted to an area located in the region putatively affected by permafrost during the LGM. Martínez Navarro et al. [13] suggested that the Val de Saône-Vosges area (which they inhabit) could have acted as a glacial refuge for endemic Lumbricidae, as distinct vegetation types (such as ombrotrophic bogs) were present in the area during the LGM [36]. The relatively deep branches between the two sampled populations of Ap. gogna are more consistent with pre-glacial diversification and survival in situ, while the extremely low genetic divergence between the two populations of Ap. velox suggests post-glacial colonization or speciation associated with glacial isolation. Very little is known about the distribution of these two species: Ap. gogna was previously known from a single locality, but in this work a second population was found, located 120 km further to the north. Likewise, a second population of Ap. velox was found during our sampling. In this case, Bouché [6] already indicated that the range of the later may be much larger, having observed unusually large casts in localities separated by 250 km. However, the deep-burrowing and fast-retreating behaviour of these species have greatly restricted sampling.

The identification of the putative ancestral area of *Aporrectodea* and several of its species provides information about their native range and

their ability to adapt to different environmental conditions and biotic interactions. Studies based on this new knowledge could improve our understanding of the effect of globalization and climate change on the role of those important ecosystem engineers in soils.

Reconstruction of the whole phylogenetic tree of Aporrectodea offers insight into the evolution of functional traits in this ecologically successful genus. The earliest branching taxa, Ap. rubra and Ap. arverna, are small, reddish, epigeic/epianecic species [37]. This may lead to the conclusion that the ancestral life form of the genus corresponded to those character states, surprisingly similar to several Lumbricus species, Eisenia, Bimastos and Satchellius, which belong to the same clade [38]. The scenario is complicated by the other early branching species, Ap. gogna, which has an unusually large body (the second largest within the present definition of the genus), very dark pigmentation and a fully anecic lifestyle. The other species that shares such specialized adaptation, Ap. velox, appears in the crown group and is not closely related. Hence, even though there appears to be a general trend towards larger body size and a more anecic lifestyle within the genus (as shown by the nocturna-giardi-longa clade), extreme phenotypes were reached convergently in two unrelated lineages. The only endogeic, (mostly) unpigmented species (Ap. caliginosa and Ap. tuberculata) appear to be derived taxa closely related to the epianecic Ap. ripicola, so that this lifestyle and trait combination can be considered to be derived within Aporrectodea. Marchán et al. [33] found that most of the early-branching species of Scherotheca are unpigmented endogeics or faintly pigmented endoanecics, suggesting the ancestral state of this genus may be different from that of Aporrectodea. However, more comprehensive sampling of the diversity of Scherotheca must be included in phylogenetic analyses to enable construction of a robust hypothesis. The acquisition of trait syndroms explaining life forms (ecological categories, sensu [6,37]) within closely related lumbricid genera is an intriguing topic, and the progressively more complete molecular dataset on these taxa is bringing us closer to understanding the situation. The relationship between earthworm functional traits, ecological categories and their effect on soil functioning is a growing focus of interest for soil biologists and could lead to a greatly improved leverage of their ecosystem services.

4.3. Remaining questions

As this work focused on endemic species of *Aporrectodea*, *Ap. caliginosa* and *Ap. trapezoides* were purposefully left aside. A large amount of species delimitation, neotyping and description remains to be performed with those peregrine, taxonomically troublesome species. The location of a putative diversification centre for the genus in Southeastern France suggests that intensive sampling and genotyping of *Ap. caliginosa* and *Ap. trapezoides* in this region could help to disentangle their systematics.

Several Central European species assigned to Aporrectodea or Eiseniona, depending on the authors (sineporis, smaragdina, pannoniella, bohiniana, kozjekensis, predalpina, etc.), remain to be included in a molecular phylogenetics frame. With inclusion of the above-mentioned Central European species, we could attempt to resolve this question as their clustering with Ap. trapezoides or with Eiseniona handlirschi would place them in their appropriate genus.

Two species which have been recurrently assigned to *Aporrectodea* but which do not appear to be closely related (thus remaining within *Aporrectodea* sensu lato) are *Ap. rosea* and *Ap. limicola. Aporrectodea* rosea has appeared as a rogue taxon (i.e. with uncertain position in a phylogenetic tree) in several molecular phylogenetic analyses [12,14] but the addition of rare species (*Panoniona satchelli* (Bouché, 1972)) suggests that it may be related to *Panoniona* [13] (but not according to the results of our work). In any case, the *Ap. rosea* species complex remains to be properly delimited and described. It appears to constitute its own genus and should receive a new name in the near future. *Aporrectodea limicola* has consistently been recovered as a sister taxon to *Scherotheca*, while not being nested within it. The inclusion of this

species in a phylogenomic (such as Anchored Hybrid Enrichment [38]) dataset could confirm this relationship and justify the need for a new name for *Ap. limicola*.

5. Conclusions

Multilocus phylogenetic analyses supported the status of two subspecies of *Ap. longa* and *Ap. giardi* as *Ap. ripicola* stat. nov. and *Ap. voconca* stat. nov., while *Ap. longa*, *Ap. nocturna* and *Ap. giardi* were confirmed as separate species. The typhlosole of *Aporrectodea* species appears to be more complex than initially thought and adopts different character states between species, which could allow further advances to be made in their taxonomy. The ancestral area for the genus *Aporrectodea* was inferred to be the southeastern perialpine region of France, with most of the diversification of the genus occurring in the same area. The earliest branching taxa (*Ap. arverna*, *Ap. rubra* and *Ap. gogna*) suggest a complex evolution of functional traits within *Aporrectodea*, with small epianecic and large anecic forms in its early evolution.

This work has resulted in a homogeneous *Aporrectodea*, with just a few Central European species (likely belonging to *Eiseniona*), *Ap. rosea* and *Ap. limicola* remaining systematically unresolved in *Aporrectodea* sensu lato, and the species complexes *Ap. caliginosa* and *Ap. trapezoides* to be delimited and formally redescribed.

Declaration of competing interest

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Data availability

Data will be made available on request.

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Appendix A. Supplementary data

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References

- J.P. McDaniel, G. Butters, K.A. Barbarick, M.E. Stromberger, Effects of Aporrectodea Caliginosa on soil hydraulic properties and solute dispersivity, Soil Sci. Soc. Am. J. 79 (3) (2015) 838–847.
- [2] G.H. Baker, M. Amato, J. Ladd, Influences of Aporrectodea trapezoides and A. rosea (Lumbricidae) on the uptake of nitrogen and yield of oats (Avena fatua) and lupins (Lupinus angustifolius): the 7th international symposium on earthworm ecology. Cardiff. Wales- 2002, Pedobiologia 47 (5–6) (2003) 857–862.

- [3] K.Y. Chan, G.H. Baker, M.K. Conyers, B. Scott, K. Munro, Complementary ability of three European earthworms (Lumbricidae) to bury lime and increase pasture production in acidic soils of south-eastern Australia, Appl. Soil Ecol. 26 (3) (2004) 257–271.
- [4] L. Örley, A palaearktikus övben élő Terrikoláknak revíziója és elterjedése.
- Értekezések a Természettudományok Köréből, Magyar Akad 15 (18) (1885) 1–34. [5] P. Omodeo, Contributo alla revisione dei Lumbricidae, Arch. Zool. Ital. 41 (1956) 129–212.
- [6] M.B. Bouché, Lombriciens de France. Écologie et systématique, 671, Institut national de la recherche scientifique, 1972.
- [7] J.P. Qiu, M.B. Bouché, Révision des taxons supraspécifiques de Lumbricoidea, Documents pédozoologiques et intégrologiques 3 (1998) 179–216.
- [8] G.E. Gates, Contributions to a revision of the Lumbricidae. XIV. What is *Enterion terrestris* Savigny, 1826 and what are its relationships? Megadrilogica 2 (4) (1975) 10–12.
- [9] C. Csuzdi Cs, A. Zicsi, *Earthworms Of Hungary (Annelida: Oligochaeta, Lumbricidae)*. Pedozoologica Hungarica No. 1, Budapest: Hungarian Natural History Museum, Budapest, 2003, p. 271, 273.
- [10] J.W. Reynolds, The Earthworms (Lumbricidae, Megascolecidae and Sparganophilidae) in Canada, Canada Food Inspection Agency, Ottawa, ii, 2022, p. 185.
- [11] R.J. Blakemore, An updated list of valid, invalid and synonymous names of Criodriloidea and Lumbricoidea (Annelida: Oligochaeta: Criodrilidae, Sparganophilidae, Ailoscolecidae, Hormogastridae, Lumbricidae, Lutodrilidae), in: M.T. Ito, N. Kaneko (Eds.), A Series of Searchable Texts on Earthworm Biodiversity, Ecology and Systematics from Various Regions of the World, Yokohama University, Yokohama pp. 1–80.
- [12] J. Domínguez, M. Aira, J.W. Breinholt, M. Stojanovic, S.W. James, M. Pérez-Losada, Underground evolution: new roots for the old tree of lumbricid earthworms, Mol. Phylogenet. Evol. 83 (2015) 7–19.
- [13] A. Martínez Navarro, S. Jiménez, T. Decaëns, M. Hëdde, M. Novo, D. Trigo, D. F. Marchán, Catch-all no more: integrative systematic revision of the genus *Allolobophora* (Crassiclitellata, Lumbricidae) with the description of two new relict earthworm genera, Org. Divers. Evol. (2023) 1–14.
- [14] M. Pérez-Losada, M. Ricoy, J. Domínguez, J. Marshall, Phylogenetic assessment of the earthworm *Aporrectodea caliginosa* species complex (Oligochaeta, Lumbricidae) based on mitochondrial and nuclear DNA sequences, Mol. Phylogenet. Evol. 52 (2009) 293–302.
- [15] R. Fernández, A. Almodóvar, M. Novo, B. Simancas, D.J.D. Cosín, Adding complexity to the complex: new insights into the phylogeny, diversification and origin of parthenogenesis in the Aporrectodea caliginosa species complex (Oligochaeta, Lumbricidae), Mol. Phylogenet. Evol. 64 (2) (2012) 368–379.
- [16] S.V. Shekhovtsov, E.V. Golovanova, S.E. Peltek, Different dispersal histories of lineages of the earthworm *Aporrectodea caliginosa* (Lumbricidae, Annelida) in the Palearctic, Biol. Invasions 18 (3) (2016) 751–761.
- [17] R. Fernández, M. Novo, D.F. Marchán, D.J. Díaz Cosín, Diversification patterns in cosmopolitan earthworms: similar mode but different tempo, Mol. Phylogenet. Evol. 94 (2016) 701–708.
- [18] P.D.N. Hebert, A. Cywinska, S.L. Ball, J.R. deWaard, biological identifications through DNA barcodes, Proc. Biol. Sci. 270 (2003) 313–321.
- [19] L.M. Hernández-Triana, S.W. Prosser, M.A. Rodríguez-Perez, L.G. Chaverri, P.D. N. Hebert, T.R. Gregory, Recovery of DNA barcodes from blackfly museum specimens (Diptera: Simuliidae) using primer sets that target a variety of sequence lengths, Molecular Ecology Ressources 14 (2014) 508–518.
- [20] N. Puillandre, S. Brouillet, G. Achaz, ASAP: assemble species by automatic partitioning, Molecular Ecology Resources 21 (2) (2021) 609–620.
- [21] M.A. Suchard, P. Lemey, G. Baele, D.L. Ayres, A.J. Drummond, A. Rambaut, Bayesian phylogenetic and phylodynamic data integration using BEAST 1.10, Virus Evolution 4 (2018) vey016, https://doi.org/10.1093/ve/vey016.
- [22] D. Darriba, G.L. Taboada, R. Doallo, D. Posada, jModelTest 2: more models, new heuristics and parallel computing, Nat. Methods 9 (8) (2012) 772.
- [23] H. Akaike, Information theory and an extension of the maximum likelihood principle, in: Proceedings of the 2nd International Symposium on Information, bn petrow, f. Czaki, Akademiai Kiado, Budapest, 1973.
- [24] G. Schwarz, Estimating the dimension of a model, Ann. Stat. 6 (2) (1978) 461–464.
 [25] A. Rambaut, A.J. Drummond, D. Xie, G. Baele, M.A. Suchard, Posterior summarisation in Bayesian phylogenetics using Tracer 1, Syst. Biol. 7 (2018)
- syy032, https://doi.org/10.1093/sysbio/syy032.[26] Y. Yu, C. Blair, X.J. He, Rasp 4: ancestral state reconstruction tool for multiple genes and characters, Mol. Biol. Evol. 37 (2) (2020) 604–606.
- [27] M. Pérez-Losada, J.W. Breinholt, M. Aira, J. Domínguez, An updated multilocus phylogeny of the Lumbricidae (Annelida: Clitellata: Oligochaeta) earthworms, Journal of Phylogenetics & Evolutionary Biology 2015 (2015).
- [28] K. Katoh, D.M. Standley, MAFFT multiple sequence alignment software version 7: improvements in performance and usability, Mol. Biol. Evol. 30 (2013) 772–780.
- [29] T.A. Hall, BioEdit: a user-friendly biological sequence alignment editor and analysis program for Windows 95/98/NT, 41, in: Nucleic Acids Symposium Series, vol. 41Information Retrieval Ltd., [London, 1999, pp. 95–98. c1979-c2000.
- [30] F. Ronquist, M. Teslenko, P. Van Der Mark, D.L. Ayres, A. Darling, S. Höhna, B. Larget, L. Liu, M. Suchard, J.P. Huelsenbeck, MrBayes 3.2: efficient Bayesian phylogenetic inference and model choice across a large model space, Syst. Biol. 61 (3) (2012) 539–542.
- [31] D.F. Marchán, T. Decaens, D.J.D. Cosin, M. Hedde, E. Lapied, J. Domínguez, French Mediterranean islands as a refuge of relic earthworm species: cataladrilus porquerollensis sp. nov. and Scherotheca portcrosana sp. nov.(Crassiclitellata, Lumbricidae), European Journal of Taxonomy 701 (2020).

- [32] S. Martinsson, C. Rhodén, C. Erséus, Barcoding gap, but no support for cryptic speciation in the earthworm Aporrectodea longa (Clitellata: Lumbricidae), Mitochondrial DNA Part A 28 (2) (2017) 147–155.
- [33] T. Milutinović, R. Tsekova, J. Milanović, M. Stojanović, Distribution, biogeographical significance and status of *Lumbricus meliboeus* Rosa, 1884 (Oligochaeta, Lumbricidae) at the European scale: first findings in Serbia and in Bulgaria, N. West. J. Zool. 9 (1) (2013).
- [34] Marchán, et al., The Cradle of Giants: Insights into the Origin of Scherotheca (Lumbricidae, Crassiclitellata) with Description of Eight New Species from Corsica, 2023. France. Marchán, D. F., Domínguez, J., Hedde, M., Decaëns, T. Zoosystema.
- [35] P.U. Clark, A.S. Dyke, J.D. Shakun, A.E. Carlson, J. Clark, B. Wohlfarth, A. M. McCabe, The last glacial maximum, Sci. Technol. Humanit. 325 (5941) (2009) 710–714.
- [36] V. Janská, B. Jimenez-Alfaro, M. Chytrý, J. Divíšek, O. Anenkhonov, A. Korolyuk, M. Culek, Palaeodistribution modelling of European vegetation types at the Last Glacial Maximum using modern analogues from Siberia: prospects and limitations, Quat. Sci. Rev. 159 (2017) 103–115.
- [37] N. Bottinelli, M. Hedde, P. Jouquet, Y. Capowiez, An explicit definition of earthworm ecological categories–Marcel Bouché's triangle revisited, Geoderma 372 (2020), 114361, https://doi.org/10.1016/j.geoderma.2020.114361.
- [38] D.F. Marchán, S.W. James, A.R. Lemmon, E.M. Lemmon, M. Novo, J. Domínguez, D. Trigo, A strong backbone for an invertebrate group: anchored phylogenomics improves the resolution of genus-level relationships within the Lumbricidae (Annelida, Crassiclitellata), Org. Divers. Evol. (2022) 1–10.