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The first stygobiont species of Coleoptera from Portugal, with a molecular phylogeny of the Siettitia group of genera (Dytiscidae, Hydroporinae, Hydroporini, Siettitiina)

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Abstract

Iberoporus pluto sp. n., the first stygobiont beetle from Portugal (Dytiscidae, Hydroporinae), is described from a single female from the cave Soprador do Carvalho (Coimbra). The species is highly troglomorphic, depigmented, blind, and with elongated appendages not adapted for swimming. A molecular phylogeny based on a combination of three mitochondrial and two nuclear genes showed the new species to be sister to I. cermenius Castro & Delgado, 2001 from Córdoba (south of Spain), within the subtribe Siettitiina of the tribe Hydroporini. Both species are included in a clade with Siettitia avenionensis Guignot, 1925 (south of France) and Rhithrodytes agnus Foster, 1992 and R. argaensis Fery & Bilton, 1996 (north of Portugal), in turn sister to the rest of species of genus Rhithrodytes Bameul, 1989, in what is here considered the Siettitia group of genera. We resolve the paraphyly of Rhithrodytes by transferring the two Portuguese species to Iberoporus Castro & Delgado, 2001, I. agnus (Foster, 1992), comb. n. and I. argaensis (Fery & Bilton, 1996), comb. n.

Keywords

Diving beetles, groundwater, new species, stygofauna, troglomorphy
Introduction

The knowledge of the subterranean fauna from Portugal has significantly increased over the last decade, with the description of a high number of obligate subterranean species (tripling their number) and the establishment of new biogeographic patterns (Reboleira 2012). A high number of these species are stygobiont (i.e., confined to groundwater), mostly from wells in the north of the country, where evapotranspiration is higher (Reboleira et al. 2011, 2013). They include 62 species of crustaceans, mostly asellids, syncarids and amphipods, and one species of annelid (Reboleira et al. 2013).

In this work we describe the first stygobiont species of Coleoptera from Portugal, a diving beetle of the subtribe Siettitiina (Dytiscidae, Hydroporinae, Hydroporini; type genus: *Siettitia* Abeille de Perrin, 1904). Siettitiina includes the only known European genera of Dytiscidae which have stygobiont members: *Siettitia*, with two species in France, *Iberoporus* Castro & Delgado, 2001, with one species in south Spain, *Etruscodytes* Mazza et al., 2013, with one Italian species, and *Graptodytes* Seidlitz, 1887, with the Moroccan *G. eremitus* Ribera & Faille, 2010 among several epigean members (Ribera and Faille 2010, Nilsson and Hájek 2018a). The subtribe also includes some North American stygobiont species, with an uncertain phylogenetic position (Miller et al. 2013, Kanda et al. 2016, Miller and Bergsten 2016, Nilsson and Hájek 2018b). The new species is known from a single female found in a well-studied cave in central Portugal. Despite multiple visits to the same cave no additional specimens have been found, so we describe here the species on the basis of its morphological singularity and of the molecular data that places it unambiguously among the west Mediterranean species of Siettitiina.

Material and methods

Taxon sampling, DNA extraction and sequencing

For the phylogenetic placement of the new species we used the datasets of Ribera and Faille (2010) and Abellán et al. (2013), with the inclusion of additional sequences (mostly nuclear genes) and taxa (Table 1). Most notably is the inclusion of *Siettitia avenionensis* Guignot, 1925, the second oldest described stygobiont water beetle worldwide. Partial sequences of the genes COI and 18S were obtained from a larva preserved in 70% ethanol, collected in 1989 (Table 1). Other attempts to extract and sequence different larvae from the same locality collected in 1984 and 1992 (Ph. Richoux leg.) proved unsuccessful. Extractions of single specimens were non-destructive, using a standard phenol-chloroform method or the DNeasy Tissue Kit (Qiagen GmbH, Hilden, Germany). Vouchers and DNA samples are kept in the collections of the Museo Nacional de Ciencias Naturales, Madrid (MNCN), the Institute of Evolutionary Biology, Barcelona (IBE) and the Natural History Museum of Denmark (NHMD).
**Table 1.** Material used in the molecular phylogeny of the *Siettitia* group of genera, with locality, collector, and EMBL accession numbers. Newly obtained sequences are in bold typeface. Nomenclature follows Nilsson and Hájek (2018a).

| No | Species         | Voucher    | Locality, date, and collector                                      | COI-5'       | COI-3'       | 16S   | 18S     | H3       |
|----|-----------------|------------|-------------------------------------------------------------------|--------------|--------------|-------|---------|----------|
| 1  | *Graptodytes* aequaticus | NHM-IR206  | Morocco: Debdelou, Mecon forestiere; 6.4.1999, I Riber‡a, P Aguiller, C. Hernandez, A Mill‡án | LS999725     | HM588264     | AY250910 | AJ850509 | EF670184 |
| 2  | *G. atlantis*   | MNCN-AJ921 | Morocco: Lac Afnouritz, Azrou; 29.4.2000, I Riber‡a               | LS999726     | HM588265     | HM588602 | LS999692 | LS999771 |
| 3  | *G. bilineatus* | MNCN-AI608 | Sweden: Vasterbotten prov., A§mele, Vindel‡lv‡en; 18.9.2005, AN Nilsson | LS999727     | HM588267     | HM588603 | LS999693 | LS999772 |
| 4  | *G. castilleanus* | MNCN-AI1316 | Spain: Navarra, Pitiar, pond in crossroad; 21.7.2004, I Riber‡a, A Cieslak | LF947943     | HM588268     | HM588604 | LS999694 | LS999773 |
| 5  | *G. delectus*   | MNCN-AI1092 | Tenerife (Spain): Chamorga, Bco. Roque Bermejo; 20.7.2006, A Castro | LS999728     | HM588269     | HM588605 | LS999695 | LS999774 |
| 6  | *G. eremitus*   | IBE-AF33   | Morocco: Tiqqi, cave Doussouli; 28.7.2000, JM Bichain et al.    | LS999729     | HM588271     | HM588606 | LS999696 | LS999775 |
| 7  | *G. flavipes*   | NHM-IR40   | Spain: Huelva, Almonte, poblado forestal; 26.7.1998, P Aguilera | –            | HM588273     | AY250914 | AJ850510 | EF670185 |
| 8  | *G. fractus*    | MNCN-AI627 | Spain: Cordoba, Sa. de Cordoba, Arroyo de los Arenales; 16.4.2005, A Castro | LS451100     | HM588274     | HM588608 | LS453474 | LS453168 |
| 9  | *G. granularis* | MNCN-AI609 | Sweden: Vasterbotten prov., A§mele, Vindel‡lv‡en; 18.9.2005, AN Nilsson | LS999730     | HM588278     | HM588611 | LS999697 | LS999776 |
| 10 | *G. ignotus*    | NHM-IR531  | Spain: Girona, Estanys de Capmany, 3.2001, P Aguilera            | LS999731     | HM588287     | AY250915 | AJ850510 | EF670185 |
| 11 | *G. kuchtae*   | MNCN-AI177 | Mallorca (Spain): Ternelles, Torrent de Ternelles; 14.10.2004, I Riber‡a, A Cieslak | LS999732     | HM588288     | HM588614 | LS999698 | LS999777 |
| 12 | *G. laeticulus* | MNCN-AI16  | Algeria: Algeria, Ain Damous; 24.8.2000, S Bouzid                | –            | HM588300     | HM588621 | LS999699 | LS999778 |
| 13 | *G. pictus*     | MNCN-AI660 | Poland: Zachodniopomorsky, Dygowo; pond; 16.8.2004, I Riber‡a, A Cieslak | LS999733     | HM588290     | HM588615 | LS999700 | LS999779 |
| 14 | *G. pietrii*    | MNCN-DM37  | Tunisia: Rd. Beja-Tebournouk, NW Tebournouk; 23.10.2001, I Riber‡a, A Cieslak | LS999734     | HM588292     | HM588616 | LS999701 | LS999780 |
| 15 | *G. sedilloti*  | NHM-IR358  | Cyprus; 3.2001, K Miller                                        | LS451098     | HM588294     | HM588619 | LS453473 | LS453167 |
| 16 | *G. sedilloti*  | MNCN-AI111 | Chios (Greece): Ternelles, Torrent de Ternelles; 14.10.2004, I Riber‡a, A Cieslak | LS999735     | HM588293     | HM588618 | LS999702 | LS999781 |
| 17 | *G. siculus*    | MNCN-AH162 | Sicily (Italy): Parco dei Nebrodi, Stream Trail Lago Urio; 13.6.2007, P Peller‡a, F Pica‡o | LS999736     | HM588295     | HM588620 | LS999703 | LS999782 |
| 18 | *G. varius*     | MNCN-AH160 | Sicily (Italy): Parco dei Nebrodi, Stream Trail Lago Urio; 13.6.2007, P Peller‡a, F Pica‡o | LS999737     | HM588297     | HM588622 | LS999704 | LS999783 |
| 19 | *G. vetetator*  | MNCN-AH161 | Sicily (Italy): Parco dei Nebrodi, Stream Trail Lago Urio; 13.6.2007, P Peller‡a, F Pica‡o | LS451095     | HM588304     | HM588625 | LS453472 | LS453105 |
| 20 | *G. vetetator*  | MNCN-AI774 | Turkey: Düzce, Rd. to Kartalkaya from Caydure; 23.4.2000, I Riber‡a | LS999738     | HM588303     | HM588624 | LS999705 | LS999784 |
| 21 | *Iberoporus* cemenius | NHM-IR276  | Spain: Cordoba, Priego de Cordoba; 29.4.2000, A Castro           | LS451107     | AY250958     | AY250918 | AJ850511 | EF670186 |
| 22 | *I. pluto* sp. n. | IBE-AN151  | Portugal: Soprador do Carvalho; 24.10.2014, APS Reboleira        | LS999739     | LS999756     | LS999763 | LS999706 | LS999785 |
| 23 | *Metaporus* meridionalis | NHM-IR34 | Spain: Albacete, Robledo, Ojos de Villaverde; 7.9.1997, I Riber‡a | –            | HM588307     | AY250919 | AJ318730 | EF670187 |
| 24 | *Porithypus* geni | IBE-RA86   | Algeria: Garat Ain Necha, nr Ben-Azrouz (Skikla); 29.6.2000, S Bouzid | LS999740     | HP931320     | HP931543 | LS999707 | LS999786 |
| N | Species             | Voucher       | Locality, date, and collector                                                                 | COI-5'   | COI-3'   | 16Sr   | 18S    | H3    |
|---|--------------------|---------------|-----------------------------------------------------------------------------------------------|----------|----------|--------|--------|-------|
| 25 | *P. lineatus*      | NHM-IR24      | England (UK): Somerset Levels, Chilton Treasury; 4.7.1998, I Ribera                           | LS999741 | AY250973 | AY250933 | AJ318743 | EF670188 |
| 26 | *P. obliqueognatus* | IBE-RA147     | Italy: Piano Grande, Piano di Castelluccio; 20.7.2009, M Toledo                              | LS999742 | HF931305 | LS999764 | LS999708 | LS999787 |
| 27 | *P. vicinus*       | MNCN-AH113    | Portugal: Cereal, ephemeral pond btw. Cereal and Vilanova; 24.1.2008, I Ribera             | LS999743 | HF931132 | HF931350 | LS999709 | LS999788 |
| 28 | *R. lineatus*      | MNCN-AH1007   | Portugal: Viana do Castelo, N Ponte de Lima, W Labruja; 28.5.2006, H Fery                   | LS999744 | HF931143 | HF931362 | LS999710 | LS999789 |
| 29 | *R. argensis*      | MNCN-AI179    | Portugal: Serra de Arga, Pools on summit; 9.5.2005, D'I Bilton                            | HF948005 | HF931183 | HF931405 | LS999711 | LS999790 |
| 30 | *R. bimaculatus*   | IBE-RA727     | Spain: Huasca, Aragues del Puerto; 23.7.2011, I Esteban                                     | LS999745 | LS999757 | LS999765 | LS999712 | LS999791 |
| 31 | *R. crux*          | MNCN-AI302    | Italy: Alessandria, stream; 2.5 km S Praglia; 18.10.2002, I Ribera, A Cieslak              | LS451084 | HF931187 | HF931410 | LS453475 | LS453108 |
| 32 | *R. numidicus*     | MNCN-DM34     | Tunisia: Rd. Tabarka-Ain-Draham, stream Ain-Draham; 23.10.2001, I Ribera, A Cieslak         | –        | LS999758 | LS999766 | LS999713 | LS999792 |
| 33 | *R. sagittatus*    | NHM-IR183     | Corsica (France): Porto-Veccio: l'Ospedale; 18.9.1999, I Ribera, A Cieslak                  | –        | AY250975 | AY250936 | AJ850513 | EF670190 |
| 34 | *S. avenionensis*  | MNCN-AI897    | France: Barbentane; 22.2.1992, J Dalmon                                                    | –        | LS999759 | –        | –       | –      |
| 35 | *S. abellani*      | IBE-PA312     | Spain: Ciudad Real, PN Cabañeros; 7.7.2008, A Millán and col.                              | LS451083 | HS991328 | HS931530 | LS453469 | LS453169 |
| 36 | *S. azernensis*    | NHM-IR661     | Morocco: Moyen Atlas, nr. Azrou, Col du Zad; 16.4.2001, Pellecchia, Pizzetti              | LS999746 | AY250979 | AY250940 | LS999715 | LS999793 |
| 37 | *S. canariensis*   | IBE-AF114     | Gran Canaria (Spain): Barranco Guigüi grande; 1.4.2008, J Hájek, K Kalikóvi                  | LS999747 | HF931113 | HF931330 | LS999716 | LS999794 |
| 38 | *S. epleuricus*    | MNCN-AH73     | Portugal: Serra de São Mamede, Portalegre: r. Caia; 25.7.1998, I Ribera                    | LS999748 | LS999760 | LS999767 | LS999717 | –      |
| 39 | *S. escheri*       | MNCN-AH107    | Morocco: Aïlilah, rd. N1, stream ca.; 4 km S Aïlilah; 27.3.2008, I Ribera, P Aguilera, C Hernando | LS999749 | HF931130 | HF931349 | LS999718 | LS999795 |
| 40 | *S. fornasii*      | MNCN-AH108    | Morocco: Aïlilah, rd. N1, stream ca.; 4 km S Aïlilah; 27.3.2008, I Ribera, P Aguilera, C Hernando | LS999750 | HF931131 | LS999768 | LS999719 | LS999796 |
| 41 | *S. lepidus*       | MNCN-AI632    | Spain: Córdoba, Sierra Morena, cta. Villavicosa; 16.4.2005, A Castro                       | LS999751 | LS999761 | LS999769 | LS999720 | LS999797 |
| 42 | *S. occidentalis*  | NHM-IR529     | Portugal: Algarve; 2001, P Aguilera                                                       | –        | AY250980 | AY250942 | –       | LS999798 |
| 43 | *S. optatus*       | MNCN-AI1089   | Spain: Jair, Sierra de Cazorla, cta. Del Tranco; 3.8.2006, A Castro                      | LS999752 | LS999762 | LS999770 | LS999721 | LS999799 |
| 44 | *S. optatus*       | NHM-MsC       | Corsica (France): Porto-Veccio: l'Ospedale; 18.9.1999, I Ribera, A Cieslak                 | –        | AY250981 | AY250943 | AJ850514 | EF670192 |
| 45 | *S. rebeccae*      | MNCN-AH72     | Portugal: Serra Estrela, Sabugueiro, r. above village; 12.5.2005, I Ribera                | LS999753 | FR851207 | FR851208 | LS999722 | LS999800 |
| 46 | *S. rufulus*       | MNCN-AI259    | Sardinia (Italy): Road from Oschiri to Mount Limbara; 17.10.2006, GN Foster                | LS999754 | HF931179 | HF931400 | LS999723 | LS999801 |
| 47 | *S. samai*         | IBE-AF142     | Algeria: Oued Bagrar; 24.3.2006, S Bouzid                                                  | LS999755 | HF931119 | HF931336 | LS999724 | LS999802 |

Examples of most species of Palaearctic *Siettitiina* were included, including all stygobiont or interstitial species with the exception of *Graptodytes aurasius* Jeannel, 1907 (Algeria), *Siettitia balsetensis* Abeille de Perrin, 1904 (France) and *Etruscodytes netbuns*
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Mazza et al., 2013 (Italy). Trees were rooted in the split between *Graptodytes*+*Metaporus* Guignot, 1945 and the rest of Siettitiina, based on previous phylogenetic results (Ribera et al. 2008, Abellán et al. 2013).

Fragments of five genes in five sequencing reactions were sequenced, three mitochondrial (1) 5′ end of cytochrome c oxidase subunit 1 (COI-5, “barcode” fragment of Hebert et al. 2003); (2) 3′ end of cytochrome c oxidase subunit 1 (COI-3); (3) 5′ end of 16S RNA plus the Leucine tRNA plus 5′ end of NADH dehydrogenase subunit I (16S); and two nuclear fragments (4) an internal fragment of the small ribosomal unit, 18S RNA (18S) and (5) an internal fragment of Histone 3 (H3). Details on primers used are provided in Table 2. Sequences were assembled and edited with Geneious v6.0.6 (Kearse et al. 2012); new sequences (111) have been submitted to the EMBL database with accession numbers LS999692–LS999802 (Table 1).

**Phylogenetic analyses**

Edited sequences were aligned using the online version of MAFFT 7 with the G-INS-I algorithm (Katoh and Toh 2008).

BEAST 1.8 (Drummond and Rambaut 2007) was used for Bayesian phylogenetic analyses, using a molecular-clock approach for estimating divergence times. We applied a partition by genes with uncorrelated lognormal relaxed clocks to estimate substitution rates and a Yule speciation process as the tree prior, using GTR+I+G and HKY+I+G evolutionary models. We calibrated the tree using rates estimated in Andújar et al. (2012) for a genus of Carabidae (*Carabus* Linnaeus, 1758), in the same suborder Adephaga (rate of 0.0113 [95% confidence interval 0.0081 – 0.0147] substitutions per site per million years (subst/s/Ma) for COI-5; 0.0145 [0.01 – 0.0198] subst/s/Ma for COI-3 and 0.0016 [0.001 – 0.0022] subst/s/Ma for 16S+tRNA). Analyses were run for 100 million generations, assessing that convergence was correct and estimating

| Gene         | Primer   | Sequence                                                                 | Reference           |
|--------------|----------|--------------------------------------------------------------------------|---------------------|
| COI-3'       | Jerry (5') | CAACATTATAATTTGATTTTTTTTGG                                                | Simon et al. (1994) |
|              | Pat (3')  | TCTATGCACTAATCTGACCATATATA                                               | Ribera et al. (2010) |
|              | Chy (5')  | T(A/T)TGACCTCAA(T/C)TTTCAAAA(T/G)GT                                      |                     |
|              | Tom (3')  | AC(A/G)TAATGAA(A/G)TGCTGAC(T/A)                                          |                     |
| COI-5'       | Uni LepF1b| TAAATATGACGACTACTATAGGGAGTTCCAAATCATAAAGATATTGGAAAC                     | Hebert et al. (2004) |
|              | Uni LepR1 | ATTAACCTCTCACATGAACCTCCTGGAATGCTCAGCTTCAAAAATCA                          |                     |
| 16S+trnL+nad1| 16SαR (5')| CGCTGTGTTTACAGAAAAACAT                                                   | Simon et al. (1994) |
|              | ND1 (3')  | GGCCTCCTAGTTGATTCCTGAATATATCC                                           |                     |
|              | 16Sb      | CCGGTCTGAACATCACGTATG                                                   |                     |
| 18S          | 18S 5'    | GAACACCTGTTGATTCCTGCAAG(T/CAGT)                                          | Shull et al. (2001) |
|              | 18S 18Sb  | TAAACGCACCAACCTTAT                                                  |                     |
| H3           | H3αF (5') | ATGGCCTGTGTTACAGAAAAGCACRCG                                             | Colgan et al. (1998) |
|              | H3αR (3') | ATATCCGGTCGCTGCAATGATG                                                  |                     |
the burn-in fraction with Tracer v1.6 (Drummond and Rambaut 2007). We also used a fast Maximum Likelihood (ML) heuristic algorithm in RAxML-HPC2 (Stamatakis 2006) in the CIPRES Science Gateway (Miller et al. 2010), using the same partition scheme as in BEAST with a GTR+G evolutionary model independently estimated for each partition and assessing node support with 100 pseudoreplicates with a rapid bootstrapping algorithm (Stamatakis et al. 2008).

**Results**

The two BEAST analyses (GTR and HKY evolutionary models) resulted in identical topologies and very similar branch lengths, although convergence for GTR evolutionary models was poor for some genes (nad1, 18S), so we present here only the results of the HKY models (Fig. 1). The topology was also almost identical to that obtained with RAxML (Fig. 1).

We obtained a well-supported, well-resolved phylogeny of Siettiina (Fig. 1). In agreement with previous results we recovered three clades, Graptodytes+Metaporus, Stictonectes Brinck, 1943 + Porhydrus Guignot, 1945, and the Siettitia group of genera as here defined, including Siettitia, Rhithrodytes, and Iberoporus (plus most likely Etruscodytes, see Discussion). The new species was placed as sister to Iberopus cermenius Castro & Delgado, 2001 with strong bootstrap support (BS = 73%), although in the Bayesian analyses the support was lower (posterior probability, pp = 0.73). Both species were in turn sister to Rhithrodytes argaensis Bilton & Fery, 1996 plus R. agnus Foster, 1992 in a very well supported clade (BS = 94; pp = 0.97), and then to Siettitia (Fig. 1). All other sampled species of Rhithrodytes were placed as sister to this clade, rendering the genus paraphyletic. In order to preserve the monophyly of Rhithrodytes we thus transfer the two species to the genus Iberoporus, Iberoporus agnus (Foster, 1992) comb. n. and Iberoporus argaensis (Bilton & Fery, 1996), comb. n.

According to our calibration, the separation between the new species and Iberoporus cermenius was dated at ca. 10 Ma (95% HPD 13.4-6.9 Ma), with a similar age for the split from I. agnus + I. argaensis (11.4 Ma [15.0-8.3]), during the Tortonian (Fig. 1).

**Taxonomy**

*Iberoporus pluto* sp. n.

http://zoobank.org/3F0A115A-F9F0-4AE5-95BC-E4E918FA04BB

Figures 2–4, 6

**Type locality.** Portugal, Penela, Gruta Soprador do Carvalho (39°59’N, 8°23’W) (Fig. 6).
Figure 1. Phylogeny of the Siettitia group of genera, obtained with Bayesian methods. Numbers in nodes, Bayesian posterior probabilities/maximum likelihood bootstrap support (obtained in RAxML); c.n., constrained node in the Bayesian analysis. See Table 1 for details on the specimens.

Type material. Holotype female (NHMD) Portugal, Penela, Gruta Soprador do Carvalho, ASPS Reboleira leg., 24.X.2014, with red holotype label and DNA voucher label “IBE-AN151”.
Diagnosis. A blind and depigmented species of *Iberoporus*, larger and wider than the other subterranean species of the genus, with a cordiform pronotum without lateral stria, less prominent constriction between pronotum and elytra and with a more transverse pronotum. Appendages longer and more slender, especially antennae and pro- and mesotibiae. Male unknown.

Description. Body length 2.8 mm, maximum width 1.1 mm. Habitus: Body elongate, strongly parallel-sided (including pronotum and head) (Fig. 2), flattened in lateral view (Fig. 3a); in dorsal view lateral outline with a slight discontinuity between posterior angles of pronotum and base of elytra. Body and appendages uniformly pale orange (cuticle appears translucent after DNA extraction due to digestion of soft tissue).

Head (Fig. 2): Wide, anterior margin almost perfectly semicircular, deeply encased in pronotum, with two lateral dark scars in place of eyes; surface smooth, with very sparse small shallow punctures, surface weakly micro-reticulated, stronger on margins, glabrous. Antennae with ovoid pedicel, distal antennomeres conical, more elongate.

Pronotum (Figs 2, 3): Cordiform, margins sinuated, anterior part slightly wider than head, posterior part narrower than head and base of elytra; anterior margin more or less straight (except angles), angles strongly acute; posterior margin sinuated, angles acute; sides without rim, anterior margin with transverse depression with irregular row of large punctures; posterior margin with some sparse large punctures very loosely forming a row. Pronotum without sublateral stria on each side, with only a slight depression and very irregular row of larger punctures. Surface smooth, with fine shallow punctures denser on disk, with very fine microreticulation, stronger near margins, cells not contiguous; centre of disc with small longitudinal rectangular mark. Pronotum with long lateral sensorial setae (Fig. 3b).

Elytra (Figs 2, 3): almost parallel-sided on basal 2/3, apical third regularly acuminate. Sides of elytra with weak rim, not visible from above. In lateral view margin of elytra almost straight, only very weakly ascending to humeral angle in anterior quarter; epipleuron not visible until shoulders. Surface with same structure as on pronotum, with very sparse larger punctures; larger punctures forming very loose and irregular lines on elytra; more distinct near to suture and on disk. With long sensorial setae on margins (Fig. 3b). Without traces of hind wings.

Ventral surface (Fig. 4): Uniformly pale, colour similar to dorsal surface. Prosternal process lanceolate, apex acuminate; not reaching anteromedial metaventral process. Epipleuron becoming narrower short before mid-length, without oblique carina near shoulder. Metepisternum more or less triangular in shape. Metacoxal lines obsolete; joint hind margin of metacoxal processes incised; lobes of processes rounded.

Legs (Figs 2–4): long and slender, especially posterior legs. Metafemora very thin, not enlarged, regularly curved; without natatorial setae.

Etymology. From “Πλούτων” (Ploutōn), the ruler of the underworld in the Greek mythology. Name in apposition.

Notes on the habitat. Soprador do Carvalho is a cave with approximately 4 km of horizontal development (Fig. 7). It is the largest cave of the so-called Dueça Speleologi-
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Figure 2. Habitus of *Iberoporus pluto* sp. n., dorsal view (holotype, after DNA extraction). Scale bar: 1 mm.

cal System, located in the north-eastern part of the Sicó karst area in central Portugal (Neves et al. 2005). The subterranean stream feeds the spring of the Dueça River, a contributor to the Mondego River. The substrate of the river is mostly composed of
Figure 3. *Iberoporus pluto* sp. n., holotype. **a** Lateral view (scale bar, 1 mm) **b** Detail of the sensory setae of pronotum and elytra (both previous to DNA extraction).
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**Figure 4.** *Iberoporus pluto* sp. n., holotype, ventral view (previous to DNA extraction).

**Figure 5.** Habitus of the species of *Iberoporus*. a *I. cermenius* (modified from Millán et al. 2014) b *I. agnus* comb. n. c *I. argaensis* comb. n. (both modified from Fery 2016).
clasts and gravel, with large clay deposits on the margins. The specimen was found in the bottom of a clay pond connected to the margin of the subterranean stream. Other invertebrate stygobionts are found in this stream, such as a new species of the asellid genus *Proasellus* and of the amphipod genus *Pseudoniphargus*, and unidentified copepods (Reboleira 2012). In the terrestrial compartment of the cave, several cave-adapted species are known: the pseudoscorpion *Occidenchthonius duecensis* Zaragoza & Reboleira, 2018; the millipede *Scutogona minor* Enghoff & Reboleira, 2013; the woodlice *Trichoniscoïdes sicoensis* Reboleira & Taiti, 2015 (which has an amphibian behaviour and can be collected inside the stream totally submerged) and *Porcellio cavernicolus* Vandel, 1946; and the dipluran *Podocampa cf. fragiloides* Silvestri, 1932 (Enghoff and Reboleira 2013, Reboleira et al. 2015, Zaragoza and Reboleira 2018). Over recent years, the cave is being explored for tourism. This may represent a major threat, as tourists constantly trample the bottom of the subterranean stream where the new species was found.

**Remarks.** *Iberoporus pluto* sp. n. is most similar in its external morphology to *I. cermenius*. Both share a similar shape of the head, a cordiform pronotum without

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**Figure 6.** Distribution map of the Iberian species of *Rhithrodytes* and *Iberoporus*. Key: red star, *I. pluto* sp. n.; blue diamond, *I. cermenius*; filled purple circle, *I. argaensis* comb. n.; empty purple circle, *I. agnus* comb. n.; black circles, *R. bimaculatus* (data from Millán et al. 2014).
lateral stria, and similar general appearance (Figs 2, 5a). In the absence of males of *I. pluto* sp. n. (and in addition to the genetic differences), both species can be easily separated by the body shape, larger and wider in *I. pluto* sp. n., and with a less prominent constriction between pronotum and elytra (clearly visible in *I. cermenius*) and with a more transverse pronotum. The appendages of *I. pluto* sp. n. are also longer and more slender, especially the antennae and the pro- and mesotibiae (Figs 2, 5a). *Iberoporus cermenius* has also well-defined parasutural rows on the elytra formed by large punctures, which are absent in *I. pluto* sp. n.

**Discussion**

We obtained for the first time a phylogeny of Siettitina including a species of its type genus, *Siettitia*. Despite the incomplete data, there is strong support for the existence of a clade including *Siettitia, Iberoporus, and Rhithrodytes*, what we call the *Siettitia* group of genera. Our results also clearly demonstrate the paracyphyly of *Rhithrodytes*, and the need to transfer two of the species to maintain its monophyly. The relationships between *Rhithrodytes* and the other three European stygobiont genera of Siettitina (*Siettitia, Iberoporus, and Etruscodytes*), although widely recognised, had not been clearly established. Originally, the genus *Rhithrodytes* was erected for a group of species...
of *Graptodytes* (the group IV of Zimmermann 1919, or the group “crux” of Guignot 1947) with a curved apex of the median lobe of the aedeagus, a lateral stria running the whole length of the pronotum (Bameul 1989) and (as recognised later), a transverse carina in the epipleura (Fery 2013). With the exception of the epipleural carina, the rest of the characters are shared with the subterranean genus *Siettitia*, which has been for long recognised to be closely related to some of the species included in *Rhithrodytes* (e.g., *R. bimaculatus* (Dufour, 1852); Régimbart 1905, Zimmermann 1932) (Table 3).

Subsequent to the description of *Rhithrodytes* two genera were described each for a single European stygobiont species: *Iberoporus* and *Etruscodytes*. *Iberoporus cermenius* shares the structure of the male genitalia with *Rhithrodytes* and *Siettitia*, but it is in particular very similar to that of *I. agnus* and *I. argaensis*. These two species (formerly in *Rhithrodytes*) have a more straight median lobe and a different shape of the apex of the parameres (Bilton and Fery 1996, Fery 2016).

The body shape of *I. agnus* and *I. argaensis* has also some similarities to the species of *Iberoporus*, parallel-sided and elongated (Figs 5b, c; see figs 12–19 in Fery 2016). *Iberoporus cermenius* shares with *Siettitia* the structure of the metacoxal processes, something that could be related to the subterranean habitat and a poor swimming ability (Castro and Delgado 2001).

*Etruscodytes*, described from a male and a female, also shares with *Rhithrodytes* and *Siettitia* the general structure of the aedeagus (note that the tip of the aedeagus in the figure of Mazza et al. 2013 seems to be broken) and the long lateral striae of the pronotum (Table 3), but nevertheless was described in a separate genus due to some morphological peculiarities (Mazza et al. 2013). Thus, according to the description by Mazza et al. (2013), the species would have (1) head wide and “subsquare” (regularly rounded in *Siettitia* and *Rhithrodytes*; although more similar to that of *Iberoporus*); (2) presence of short and flattened setae on pronotum and elytra; (3) prosternal process contacting anteromedial process of metaventrite (also in *Rhithrodytes*, not in *Siettitia* and *Iberoporus*, Table 3); (4) anteromedial process of metaventrite rounded (pointed in *Siettitia* according to Mazza et al. 2013); (5) ventrites II and III not fused (fused in *Siettitia* and *I. cermenius*, not in *I. pluto* sp. n. or *Rhithrodytes*); (6) elytra not completely fused (fused in *Siettitia*, not in *Iberoporus* and *Rhithrodytes*). Some of these characters seem to be clear autapomorphies related to the subterranean life (fusion of elytra or ventrites, particularly shaped setae, lack of lateral striae on the pronotum, lack of carina on the epipleuron), and others are of uncertain interpretation. Thus, the structure of the prosternal process is sometimes difficult to appreciate, but there do not seem to be fundamental differences between the species (note that in fig. 7 in Mazza et al. 2013 the prosternal process seems to fit below the anteromedial process of the metaventrite, which is likely an artefact), being the differences consequence of the different position of the mesocoxa (contiguous or not) and ultimately the width of the body, which in turn may depend on the habitat and ecology of the species. More data, especially molecular sequences of *Etruscodytes* and *Siettitia*, and the likely discovery of other subterranean taxa would contribute to the understanding of the evolution of this western Mediterranean lineage.
**Table 3.** Summary comparison of some character states among the taxa of the *Siettitia* group of genera (character states of *Etruscodytes* obtained from Mazza et al. 2013).

| Character and character state                  | *Siettitia* | *Etruscodytes* | *Iberoporus cermenius, I. pluto sp. n.* | *Iberoporus agnus, I. argaensis* | *Rhithrodytes sensu novo* |
|-----------------------------------------------|-------------|----------------|----------------------------------------|----------------------------------|--------------------------|
| sublateral pronotal stria                    | long        | long           | absent                                 | long                             | long                     |
| subhumeral epipleural carina                 | absent      | absent?        | absent                                 | present                          | present                  |
| pigmentation of elytra                       | weak        | weak           | weak                                   | strong                           | generally strong         |
| eyes                                          | absent      | absent         | absent                                 | present                          | present                  |
| body shape, general                          | parallel    | parallel       | parallel                               | oval-parallel                    | generally oval           |
| constriction at bases of pronotum and elytra | absent      | absent         | absent                                 | absent                           | absent                   |
| contact between prosternal process and       | absent      | present        | absent                                 | present                          | present                  |
| anteromedial metasternal process             | fused       | not fused      | fused in *I. cermenius*                | not fused                        | not fused                |
| ventrites II and III                         | fused       | partly fused?  | not fused                              | not fused                        | not fused                |
| elytra                                       |             |                |                                        |                                  |                          |

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