

High diversity of fish ectoparasitic monogeneans (Dactylogyrus) in the Iberian Peninsula: a case of adaptive radiation?

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1 High diversity of fish ectoparasitic monogeneans (Dactylogyrus) in the

2 Iberian Peninsula: a case of adaptive radiation?

- 3 Michal Benovics^{1*}, Yves Desdevises², Radek Šanda³, Jasna Vukić⁴, Mathilde Scheifler², Ignacio
- 4 Doadrio⁵, Carla Sousa-Santos⁶, Andrea Šimková¹

5

- 6 Department of Botany and Zoology, Faculty of Science, Masaryk University, Kotlářská 2, 61137
- 7 Brno, Czech Republic.
- 8 ² Sorbonne Université, CNRS, Biologie Intégrative des Organismes Marins, BIOM, Observatoire
- 9 Océanologique, F-66650, Banyuls/Mer, France
- ³ National Museum, Václavské Náměstí 68, 115 79 Prague, Czech Republic
- ⁴ Department of Ecology, Faculty of Science, Charles University, Viničná 7, 128 44 Prague, Czech
- 12 Republic
- ⁵ Biodiversity and Evolutionary Group, Museo Nacional de Ciencias Naturales-CSIC, C/José
- Gutiérrez Abascal 2, 28006, Madrid, Spain
- ⁶ MARE Marine and Environmental Sciences Centre, ISPA Instituto Universitário, Rua Jardin do
- Tabaco 34, 1149–041, Lisbon, Portugal

- 18 Yves Desdevises desdevises@obs-banyuls.fr
- 19 Radek Šanda radek.sanda@nm.cz
- 20 Jasna Vukić jasna.vukicova@natur.cuni.cz
- 21 Mathilde Sheifler mathilde.scheifler@obs-banyuls.fr
- 22 Ignacio Doadrio doadrio@mncn.csic.es
- 23 Carla Sousa-Santos carla.santos@ispa.pt
- 24 Andrea Šimková simkova@sci.muni.cz
- 25 *corresponding author: benovics@mail.muni.cz
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Abstract

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The epicontinental fauna of the Iberian Peninsula is strongly influenced by its geographical history. As the possibilities for dispersion of organisms into and from this region were (and still are) limited, the local fauna consists almost exclusively of endemic species. Almost all Iberian freshwater fishes of the families Leuciscidae and Cyprinidae are endemic and ongoing research on these taxa continually uncovers new species. Nevertheless, information on their host-specific parasites remains scarce. In this study, we investigate the diversity and phylogenetic relationships in monogeneans of the genus Dactylogyrus (gill ectoparasites specific to cyprinoid fish) in the Iberian Peninsula. Twenty-two species were collected and identified from 19 host species belonging to Cyprinidae and Leuciscidae. A high degree of endemism was observed, with 21 Dactylogyrus species reported from Iberia only and a single species, D. borealis, also reported from other European regions. Phylogenetic analysis split the endemic Iberian Dactylogyrus into two well-supported clades, the first encompassing Dactylogyrus parasitising endemic Luciobarbus spp. only, and the second including all Dactylogyrus species of endemic leuciscids and four species of endemic cyprinid. Species delimitation analysis suggests a remarkable diversity and existence of a multitude of cryptic Dactylogyrus species parasitising endemic leuciscids (Squalius spp. and representatives of Chondrostoma s.l.). These results suggest a rapid adaptive radiation of Dactylogyrus in this geographically isolated region, closely associated with their cyprinoid hosts. Moreover, phylogenetic analysis supports that *Dactylogyrus* parasites colonised the Iberian Peninsula through multiple dispersion events.

Keywords

49 Monogenea; phylogeny; host-specificity; species delimitation; Iberian Peninsula; Cyprinoidea

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1 Introduction

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The Iberian Peninsula has a remarkable biological diversity, harbouring more than 50% of 64 European animal and plant species (Médail and Quézel, 1997, Martín et al., 2000, Williams et 65 al., 2000, Araújo et al., 2007, Cardoso, 2008, Rueda et al., 2010, López-López et al., 2011, 66 Penado et al., 2016) and approximately 31% of all European endemic vertebrate and plant 67 species (Williams et al., 2000). This high species diversity is linked with several climatic and 68 geological changes occurring over the region since the Cenozoic period (Hsü et al., 1973, 69 Rosenbaum et al., 2001), when putative migration routes periodically emerged and 70 disappeared. However, the main factor influencing the degree of endemism is most likely 71 72 geographical isolation resulting from the elevation of the Pyrenees in the north-east combined 73 with the generally mountainous topography of the peninsula, which provided a multitude of refuges during glacial periods (Gante et al., 2009, Hewitt, 2011). 74 While the species diversity of Iberian freshwater ichthyofauna is relatively low in comparison 75 to other European regions (Kottelat and Freyhof, 2007), the majority of species are endemic. 76 77 The Peninsula hosts representatives of just a few native freshwater fish groups, with most species belonging to the Cyprinidae and Leuciscidae families (order Cyprinoidea; following 78 the classification proposed by Schönhuth et al., 2018). The Leuciscidae (previously 79 considered as Leuciscinae within Cyprinidae; Ketmaier et al., 2004, Levy et al., 2009, Perea 80 et al., 2010, Imoto et al., 2013) are represented by the monotypic genus Anaecypris, the 81 82 genera Phoxinus, Iberocypris and Squalius, and by four recently erected genera belonging to Chondrostoma sensu lato: Achondrostoma, Iberochondrostoma, Parachondrostoma and 83 Pseudochondrostoma (Kottelat and Freyhof, 2007, Robalo et al., 2007, Schönhuth et al., 84 2018). In contrast to the leuciscids, cyprinids are represented by just two genera: Barbus and 85 Luciobarbus (Kottelat & Freyhof, 2007, Gante, 2011, Gante et al., 2015). The distribution of a 86 given cyprinoid species is usually confined to a specific ichthyogeographic province and the 87

ranges of different species rarely overlap (Doadrio, 1998, Gante et al., 2015), suggesting that speciation is closely linked with the formation of river basins (Zardoya and Doadrio, 1998, Machordom and Doadrio, 2001, Doadrio et al., 2002, Mesquita et al., 2007, Casal-López et al., 2017, Sousa-Santos et al., 2019). In contrast to the thorough previous and on-going research on Iberian cyprinoids, data on their helminth parasites are scarce (da Costa Eiras, 2016). In previous studies focussed on freshwater fishes in different regions of the northern hemisphere (e.g. Mexico and the Balkans), it has been suggested that the biogeography of fish helminth parasites reflects the historical dispersion and current distribution of their hosts (e.g. Choudhury and Dick, 2001, Pérez-Ponce de León et al., 2005, Benovics et al., 2018). However, very few studies have been carried out on cyprinoid monogeneans in the Iberian Peninsula, by far the most thorough being those of el Gharbi et al. (1992) and Šimková et al. (2017). The former study, describing seven species of *Dactylogyrus* from six cyprinid species (relying on morphological data only) suggested that the pattern of geographical distribution of Dactylogyrus spp. follows the distribution of their cyprinid hosts, for which they are highly host-specific. The study by Šimková et al. (2017) focussed on phylogenetic relationships between endemic *Dactylogyrus* from cyprinids in Iberia and *Dactylogyrus* from Central Europe and north-west Africa. The authors suggested multiple origins of endemic *Dactylogyrus* in the Iberian Peninsula as the presence of Dactylogyrus lineages in different Luciobarbus lineages was associated with specific dispersion events. Gill monogeneans belonging to *Dactylogyrus* are currently the most species-diversified group within the Platyhelminthes (more than 900 nominal Dactylogyrus species, mostly described from morphology, are presently known according to the latest review by Gibson et al., 1996). Dactylogyrus species are strictly specific to cyprinoids and many Dactylogyrus species are specific to a single host species (Šimková et al., 2006b). However, the degree of host

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specificity across *Dactylogyrus* species differs and, in some cases, host specificity is likely to reflect the ecology and recent distribution of their hosts (Benovics et al., 2018). Dactylogyrus species with a narrow host range are most common in regions with a high number of endemic host species. In Europe, such regions include the Balkan Peninsula, where a multitude of strictly host-specific endemic Dactylogyrus species has been documented (Dupont and Lambert, 1986, Benovics et al., 2017, 2018), and the Iberian Peninsula, where many Dactylogyrus endemic species have been documented for Luciobarbus (el Gharbi et al., 1992). It has been suggested that such a high degree of endemism in *Dactylogyrus* is the result of co-speciation with their hosts over long evolutionary periods in geographically isolated regions (Dupont, 1989). Over time, the *Dactylogyrus* parasites have developed an attachment organ (haptor) that is highly specialised toward their host (Šimková et al., 2000, Jarkovský et al., 2004, Šimková and Morand, 2008). As such, the shape and size of monogenean haptoral sclerites are considered to be species specific and represent suitable morphological characters for species determination. Nevertheless, some species exhibit haptoral sclerites that are very similar in shape and size (see Pugachev et al., 2009); thus, species identification is often difficult from the observation of haptoral sclerotised structures only. It has been suggested, therefore, that the shape of the sclerotised parts of copulatory organs are more suitable for the identification of monogeneans to species level due to their putative faster evolutionary rate (Pouvaud et al., 2006, Šimková et al., 2006b, Vignon et al., 2011, Mendlová et al., 2012, Mandeng et al., 2015, Benovics et al., 2017). Rapid morphological diversification in the monogenean copulatory organs is hypothesised to be a mechanism to avoid hybridisation (Rohde 1989), which is especially likely for *Dactylogyrus* species living on the same hosts in overlapping microhabitats (Šimková et al., 2002, Šimková and Morand, 2008). Compared with Central Europe, Dactylogyrus communities in the southern European Peninsulas generally appear to be species poor. Cyprinoids with a wide European distribution

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range, such as Rutilus rutilus and Squalius cephalus, harbour up to nine Dactylogyrus species 138 (e.g. Šimková et al., 2000, Seifertová et al., 2008). In contrast, a maximum of five 139 Dactylogyrus species per cyprinoid species have been reported from the southern European 140 Peninsulas (Dupont and Lambert, 1986, el Gharbi et al., 1992, Galli et al., 2002, 2007, 141 Benovics et al., 2018). 142 In comparison to other European regions, cyprinoid monogenean communities have been 143 underexplored in the Iberian Peninsula. Thus, the main objective of the present study was to 144 investigate the diversity of *Dactylogyrus* spp. parasitising endemic cyprinoids in this 145 geographical region. A species delimitation method was applied to assess the species status of 146 147 Dactylogyrus identified in this study based on genetic variability within and among each 148 species, and to compare these results to species defined from morphology only. Moreover, the present study investigates the evolutionary history and phylogenetic relationships between 149 endemic Iberian Dactylogyrus and Dactylogyrus from other Peri-Mediterranean regions, 150 including cyprinoid species with a wide European distribution range, in order to (1) shed new 151 light on cyprinoid phylogeography, (2) infer potential historical contacts between cyprinoids 152 from different regions, and (3) evaluate the evolution of *Dactylogyrus* species diversity (using 153 both morphology and species delimitation methods). 154

2 Material and Methods

2.1 Parasite collection

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Fish were collected over years 2016 and 2017 from 17 localities in Portugal and Spain (Figure 1). In total, 257 specimens representing 19 fish species were examined for presence of *Dactylogyrus* parasites (Table 1). Fish were dissected following the standard protocol described by Ergens and Lom (1970). *Dactylogyrus* specimens were collected from the gills, mounted on slides and fixed in a mixture of glycerine and ammonium picrate (GAP; Malmberg, 1957) for further identification. Determination to species level was performed on

the basis of the size and shape of the sclerotised parts of the attachment apparatus (anchor hooks, marginal hooks and connective bars of the haptor) and the reproductive organs (male copulatory organ and vaginal armament) following Pugachev et al. (2009). At least five specimens of each *Dactylogyrus* species from each host species examined were bisected using fine needles. One-half of the body (either the anterior part containing the reproductive organs or the posterior part with the attachment organ) was mounted on a slide and used for morphological identification. The other half was individually preserved in pure ethanol for subsequent DNA extraction.

2.2 DNA extraction, PCR and sequencing

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DNA extraction was performed using the DNeasy Blood & Tissue Kit (Quiagen, Hilden, Germany) based on the standard protocol provided by the manufacturer. Two DNA regions were amplified. The partial gene coding 18S rRNA and complete ITS1 region was amplified using the primers S1 (forward, 5'-ATTCCGATAACGAACGAGACT-3') and Lig5.8R (reverse, 5'-GATACTCGAGCCGAGTGATCC -3') (Šimková et al., 2003, Blasco-Costa et al., 2012). Each amplification reaction was performed in a final volume of 20 µl, the reaction mixture comprising 1.5 U Taq polymerase (Fermentas), 1x buffer, 1.5 mM MgCl2, 0.2 mM of dNTPs, 0.1 mg/ml BSA, 0.5 µM of each primer and 2 µl of pure DNA (20 ng/µl). PCR was carried out using the following steps: 3 min initial denaturation at 95 °C, followed by 40 cycles of 40 seconds at 94 °C, 30 seconds at 52 °C and 45 seconds at 72 °C, and 4 minutes of final elongation at 72 °C. The second marker, a part of the gene coding 28S rRNA, was amplified using the primers C1 (forward, 5'-ACCCGCTGAATTTAAGCA-3') and D2 (reverse, 5'-TGGTCCGTGTTTCAAGAC-3') (Hassouna et al., 1984), following the PCR protocol described in Šimková et al. (2006a). The PCR products were purified prior to sequencing using the ExoSAP-IT kit (Ecoli, Bratislava, Slovakia), following the standard protocol, and directly sequenced using the PCR primers and the BigDye Terminator Cycle

Sequencing kit (Applied Biosystems, Foster City, CA). Sequencing was carried out on an ABI 3130 Genetic Analyzer (Applied Biosystems). The newly generated sequences were deposited in GenBank (see Table 1 for accession numbers).

2.3 Phylogenetic and species delimitation analysis

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Partial sequences coding 18S rRNA and 28S rRNA, and complete sequences of the ITS1 region were concatenated and aligned using the fast Fourier transform algorithm implemented in MAFFT (Katoh et al., 2002) using the G-INS-i refinement method. Out of 71 DNA sequences used in the alignment, 35 were newly sequenced in this study. Sequences from 35 other *Dactylogyrus* species, used as representative species from different European regions, and sequences of Ancyrocephalus percae, used as an outgroup (phylogenetically closely related to Dactylogyrus according to Mendoza-Palmero et al., 2015), were obtained from GenBank (see Table S1 in supplementary material for accession numbers). Gaps, hypervariable regions and ambiguously aligned regions were removed from the alignment using GBlocks v. 0.91 (Talavera and Castresana, 2007). The optimal DNA evolutionary model was selected separately for each part of the alignment corresponding to one of the three markers analysed (18S, ITS1, 28S) using the Bayesian information criterion in jModelTest v. 2.1.10 (Guindon and Gascuel, 2003, Darriba et al., 2012). Maximum likelihood (ML) analysis was conducted in RAxML v. 8.2.11 (Stamatakis, 2006, 2014), applying the general time-reversible model (GTR; Lanave et al., 1984) of nucleotide substitution. Internal node support was assessed by running 1000 bootstrap pseudoreplicates. Bayesian inference (BI) analysis was performed in MrBayes v. 3.2.6 (Ronguist et al., 2012) using two parallel runs, each with four Markov chains (one cold and three heated) of 10⁷ generations with trees sampled every 10² generations. The first 30% of trees were discarded as initial burn-in. Convergence was indicated by an average standard deviation of split frequencies per parallel run of < 0.01, subsequently checked using Tracer v. 1.7.1 (Rambaut et al., 2018). Posterior probabilities were calculated as the frequency of samples recovering particular clades. To investigate genetic diversity in the commonly used genetic markers between well-defined endemic *Dactylogyrus* species, uncorrected pair-wise genetic distances (p-distances) were computed for 12 selected taxa in MEGA X (Kumar et al., 2018). Three sequence alignments were used: the partial gene coding 18S rRNA, the complete ITS1 region and the partial gene coding 28S rRNA. All positions containing gaps and missing data were removed from final computations. The Bayesian implemented Poisson Tree Processes model (bPTP; Zhang et al., 2013) was applied to the phylogram resulting from BI in order to infer putative species of Iberian Dactylogyrus. The bPTP method only requires a phylogenetic tree as its input and uses branch lengths to estimate the mean expected number of substitutions per site between two branching events. Within species, branching events will be frequent whereas they will be rarer between species. The model implements two independent classes of Poisson process (one describing speciation and the other describing coalescent processes) and searches for transition points between interspecific and intraspecific branching events. Potential species clusters are then determined by identifying the clades or single lineages that originate after these transition points. The computation was run for 5 x 10⁵ generations with the first 30% of trees discarded as initial burn-in. The distant outgroup taxon was removed from the final analysis to improve

3 Results

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Twenty-two *Dactylogyrus* species (identified using morphological characters, i.e. sclerotised parts of the haptor and reproductive organs) were collected from endemic Iberian cyprinoid species (Table 1). From one to five *Dactylogyrus* species were recorded per host species, with highest species richness found on *Luciobarbus* spp. (5 species on *L. guiraonis*, 4 species on *L.*

graellsii, and 4 species on L. sclateri). Both Parachondrostoma species, Barbus haasi, 238 Iberochondrostoma almacai and Phoxinus bigerri were parasitised by a single Dactylogyrus 239 species. Overall, Dactylogyrus bocageii exhibited the widest host range across the Iberian 240 241 Peninsula, parasitising four Luciobarbus species. Minor genetic variation was observed between D. bocageii collected from different hosts (p-distance ≤ 0.002 in the partial gene for 242 28S rRNA, p-distance \leq 0.020 in the ITS1 region; Tables 2 and 3). 243 The final concatenated alignment of partial genes for 18S rRNA, 28S rRNA and the ITS1 244 region included 71 sequences contained 1533 unambiguous nucleotide positions. The most 245 suitable evolutionary models were TrNef+I+G, TPM2uf+G and GTR+I+G for the partial 246 247 genes coding 18S rRNA, the ITS1 region and part of the gene for 28S rRNA, respectively. 248 Both ML and BI analyses produced trees with congruent topologies varying only in some 249 support values for individual nodes (Figure 2). Phylogenetic analysis divided all taxa into three strongly supported clades. 250 The first group (Clade A; Figure 2) included the majority of *Dactylogyrus* species from 251 252 Europe, and especially the species parasitising Leuciscidae. In addition, several *Dactylogyrus* species from Barbus and Luciobarbus (Cyprinidae) were also placed in this clade (i.e. 253 254 Dactylogyrus of Barbus spp. and Luciobarbus spp. from Central Europe and the Balkans, and D. balistae, D. legionensis, D. linstowoïdes and D. andalousiensis of Iberian Luciobarbus 255 spp.). Dactylogyrus from Iberian cyprinoids were divided into seven lineages within Clade A. 256 257 Dactylogyrus polylepidis of Achondrostoma arcasii was in a well-supported sister position to the morphologically similar D. vistulae. Dactylogyrus from European cyprinids formed three 258 well-supported groups within Clade A. Dactylogyrus legionensis, D. balistae and D. 259 linstowoïdes were grouped in a sister position to common Dactylogyrus species from Central 260 European Barbus spp. (D. malleus, D. prespensis and D. petenyi). The second group 261 contained D. andalousiensis from two Iberian Luciobarbus species, and D. omenti from 262

Aulopyge huegelii (Balkan endemic species). The third group contained D. carpathicus and D. crivellius (two common species of Barbus spp.) and two yet undescribed endemic Dactylogyrus species of endemic Balkan Luciobarbus species (L. albanicus and L. graecus). The phylogenetic position of Dactylogyrus sp. 1 from S. aradensis and S. torgalensis (morphologically identical but genetically slightly different; p-distance = 0.010) was not fully resolved and its sister position to D. folkmanovae was only supported by BI. The majority of Iberian Dactylogyrus species (Dactylogyrus sp. 2 to Dactylogyrus sp. 10) formed a welldefined phylogenetic lineage that also included D. caucasicus, D. ergensi and D. tissensis. The three latter species and the Iberian *Dactylogyrus* in this lineage all have the same or very similarly shaped male copulatory organs commonly classified as 'ergensi' of the 'chondrostomi' type (see Pugachev et al., 2009). Generalist Dactylogyrus species within Clade A (i.e. D. legionensis, D. polylepidis, Dactylogyrus sp. 1, Dactylogyrus sp. 7 and Dactylogyrus sp. 8) exhibited intraspecific genetic variability. The second major group (Clade B) comprised five *Dactylogyrus* species specific to Iberian *Luciobarbus*. Where intraspecific genetic variability was documented, all genetic variants formed well supported clades (i.e. D. bocageii, D. guadianensis and D. lenkoranoïdes). The last strongly supported group (Clade C) encompassed Dactylogyrus species host specific to Carassius spp. and/or Cyprinus carpio distributed across the Europe and Asia. Genetic distances were computed between morphologically similar species from Clade B (Figure 2). Three alignments of 12 sequences representing five *Dactylogyrus* species of group B were analysed to compare intra- and interspecific genetic variability calculated using genetic markers commonly used in monogeneans. The alignments comprised 486 nucleotide positions for the partial gene coding 18S rRNA combined, 716 nucleotide positions for the ITS1 segment and 807 nucleotide positions for the partial gene coding 28S rRNA. The lowest genetic variability was observed for the partial gene coding 18S rRNA. No intraspecific/inter-

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population genetic variability was observed (p-distance = 0.000) and interspecific pair-wise nucleotide diversity varied from 0.002 to 0.010 (Table 4). Low pair-wise interspecific diversity was also observed for the partial gene coding 28S rRNA (0.006 to 0.020); however, minor intraspecific genetic variability was observed in this gene (p-distance < 0.002). Slight genetic distance in part of the gene for 28S rRNA was observed between different populations of D. bocageii (0.001 to 0.002) and between individuals from different populations of D. guadianensis (p-distance = 0.001). The highest genetic diversity was observed in the ITS1 region, in which intraspecific distances varied from 0.000 (D. lenkoranoïdes) to 0.020 (D. bocageii). The pair-wise interspecific diversity in the ITS1 region varied from 0.031 between D. doadrioi and D. guadianensis to 0.135 between D. doadrioi and D. mascomai. The species status of Dactylogyrus collected from endemic Iberian cyprinoids was investigated using the bPTP method, with the addition of *Dactylogyrus* species parasitising cyprinoids in other parts of Europe used as a reference of previously delimited species (Benovics et al., 2018). The results of the bPTP analysis were largely consistent with the species previously described on the basis of morphology (Figure 3), though the ML solution suggested a higher species diversity. Based on ML results, D. legionensis encompasses two species, each being host-specific (one to L. graellsii and the other to L. guiraonis), as well as Dactylogyrus sp. 1 (S. aradensis and S. torgalensis). Both BI and ML supported solutions, obtained from bPTP analysis, suggested a generalist status for D. andalousiensis, D. bocageii, D. lenkoranoïdes and D. guadianensis (i.e. there were no host-specific parasites within these delimited species). A potentially new species, *Dactylogyrus* sp. 7, was also supported by the species delimitation analysis as a generalist, parasitising both S. carolitertii and S. pyrenaicus. This analysis also suggested that D. borealis, determined using morphological characters, is a common parasite of *Phoxinus* spp. in other parts of Europe and is also found on *P. bigerri* in the Iberian Peninsula. bPTP analysis also suggested that Parachondrostoma miegi and P.

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turiense are both parasitised by a single *Dactylogyrus* species (*Dactylogyrus* sp. 8) that is morphologically similar and phylogenetically close to *Dactylogyrus* sp. 9, parasitising *P. duriense*. Finally, species delimitation analysis supported the discovery of at least 11 unknown *Dactylogyrus* species in the Iberian Peninsula, as all other Iberian genetic variants were identified as individual host-specific species.

The Iberian Peninsula harbours a high diversity of cyprinoids that have been the subject of

4 Discussion

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4.1 Parasite diversity and distribution

321 extensive research; nevertheless, the species diversity of their host-specific parasites is still underexplored, especially in areas with a high diversity of endemic cyprinoids. Following 322 previous research on the *Dactylogyrus* (or Monogenea in general) of Iberian cyprinids (el 323 Gharbi et al., 1992, Lacasa-Millán and Gutiérrez-Galindo, 1995, Gutiérrez-Galindo and 324 Lacasa-Millán, 2001), this study is the first to investigate the overall diversity of Iberian 325 Dactylogyrus, including molecular data for both cyprinoid fish and their host-specific 326 Dactylogyrus. 327 328 The present study revealed the presence of several potentially new *Dactylogyrus* species to science, all of which were well supported by the bPTP species delimitation method. This 329 strongly suggests that endemic Iberian cyprinoid species harbour an endemic Dactylogyrus 330 fauna, as previously suggested for Iberian Luciobarbus species by el Gharbi et al. (1992). In 331 contrast to the Balkan and Apennine Peninsulas (Dupont and Lambert, 1986, Dupont and 332 Crivelli, 1988, Dupont, 1989, Galli et al., 2002, 2007, Benovics et al., 2018), Iberian 333 334 Dactylogyrus spp. appear to exhibit a higher degree of host specificity as the majority of Dactylogyrus species from Leuciscidae were restricted to a single host species. Benovics et al. 335 (2018) proposed that southern European endemic cyprinoids harbour species-poor 336 Dactylogyrus communities compared with European cyprinoids with a wide distribution 337

range (e.g. Rutilus rutilus, Squalius cephalus). The same pattern was also observed in the Iberian Peninsula, where one to five Dactylogyrus species were found on a single cyprinoid host species. It should be noted, however, that parasite community composition may be strongly influenced by seasonal abiotic factors (e.g. González-Lanza and Alvarez-Pellitero, 1982, Lux, 1990, Appleby and Mo, 1997, Šimková et al., 2001b, Poulin and Morand, 2004, Zhang et al., 2015, Sinaré et al., 2016). Until now, knowledge of *Dactylogyrus* diversity in southern European Mediterranean Peninsulas has been based on studies taking place in summer only (Benovics et al, 2018, this study) as the *Dactylogyrus* diversity is expected to be highest during this period (Šimková et al., 2001b). In this study, a higher number of *Dactylogyrus* species was observed on *Luciobarbus* species. While the overall species richness on these fish was in accordance with the observations of el Gharbi et al. (1992), the species composition in the present study differed slightly from their data. In line with the study of el Gharbi et al. (1992), D. bocageii was the most common species (occurring on five Luciobarbus species), though its distribution range was wider, as proposed by Lambert and el Gharbi (1995), stretching via Zujar and Torgal rivers to the south-western part of the peninsula (south-west Iberian province, Filipe et al., 2009). Interestingly, unlike other European regions, the only endemic representative of the genus Barbus in Iberia, Barbus haasi, harbours Dactylogyrus species typical of Luciobarbus spp.. In the Balkans, endemic Barbus spp. are parasitised by common Dactylogyrus species for this fish genus (e.g. D. dyki and D. crivellius), whilst Luciobarbus spp. are parasitised by different, strictly host-specific species (Benovics et al., 2017, 2018). In accordance with our own findings, El Gharbi et al. (1992) showed that B. haasi is a common host of D. bocageii, D. mascomai and D. lenkoranoïdes, whilst D. dyki and D. carpathicus (commonly distributed on European Barbus spp.) were only found in previous studies on B. haasi x B. meridionalis hybrids in the north-eastern part of the Peninsula. Nevertheless, Gutiérrez-Galindo and

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Lacasa-Millán (1999) also reported the latter two Dactylogyrus species from B. haasi in the River Llobregat (north-east Spain). However, the fish hosts from this study could potentially also be hybrids, as presence of the B. haasi x B. meridionalis hybrids was previously documented in Llobregat basin (Machordom et al., 1990). In contrast to the aforementioned studies, only D. lenkoranoïdes was collected from B. haasi in this study (Uldemo River; Ebro basin). This low parasite diversity may be linked with the seasonal fluctuation in parasite communities previously documented among Iberian Dactylogyrus (e.g. D. legionensis (González-Lanza and Alvarez-Pellitero, 1982) or D. balistae (Simón-Vicente, 1981)). In addition to the common parasitisation of Iberian Barbus by Dactylogyrus parasites typically recognised as specific to Luciobarbus, several cases of infection by Dactylogyrus species common for Barbus were also reported in Iberian Luciobarbus species. Gutiérrez-Galindo and Lacasa-Millán (2001) also reported that L. graellsii was parasitised by D. dyki and D. extensus (host specific parasites of Barbus spp. and Cyprinus carpio, respectively). However, the presence of D. dyki on Luciobarbus spp. may result from non-detected instances of hybridisation, as hybrids of cyprinoid species are usually parasitised by *Dactylogyrus* specific for each of the parental species (Šimková et al., 2013, Krasnovyd et al. 2017). Hybridisation between Iberian Luciobarbus spp. (potentially also between Luciobarbus and Barbus; Gante et al., 2015) appears to be quite common, especially between congeners living in sympatry (e.g. Luciobarbus spp.; Almodóvar et al., 2008, Sousa-Santos et al., 2018). Thus, hostswitching is possible, most likely occurring between species from phylogenetically close genera (i.e. Barbus and Luciobarbus; Yang et al., 2015) in north-eastern Iberian drainages where the distribution ranges of Central European barbels (e.g. B. meridionalis; see Kottelat and Freyhof, 2007 for its distribution range) and Iberian barbels overlap. Despite the presence of high numbers of endemic *Dactylogyrus* species in Iberia, *P. bigerri* was parasitised by D. borealis, a common species on European Phoxinus spp. (Moravec,

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2001, Šimková et al., 2004, Benovics et al., 2018). The presence of this common European Dactylogyrus species is in contrast to the expected high degree of endemism in south European peninsulas (Williams, 2000, Hewitt, 2011). Other common European Dactylogyrus species are absent from Iberia; for example, D. vistulae, which parasitises the highest number of cyprinoid species across Europe, is absent from Iberia, and only the closely related D. polylepidis is found on Iberian cyprinoids. These findings suggest that either (1) D. borealis was only recently introduced into the Iberian Peninsula with another *Phoxinus* species coming from different European areas (see Corral-Lou et al., 2019), or (2) D. borealis represent an extremely slowly evolving species, meaning that the Iberian lineage would be morphologically and genetically similar to D. borealis from other European areas. In the present study, D. polylepidis, originally described from Pseudochondrostoma polylepis (Alvarez-Pellitero et al., 1981), was found for the first time on three host species (all members of the Leuciscidae). The wider host range recorded for D. polylepidis indicates that this species represents a true generalist parasite, probably endemic to this region. In contrast to D. polylepidis, the morphologically similar and phylogenetically closely related D. vistulae is a typical generalist in Europe (except Iberia) and Asia, parasitising a multitude of cyprinoid species and genera (Moravec, 2001, Benovics et al., 2018). Dactylogyrus polylepidis and D. vistulae share remarkably similar morphological traits, including an enlarged seventh pair of marginal hooks, large anchor hooks and a similar size and shape of the copulatory organs (see Pugachev et al., 2009). It has previously been hypothesised that large attachment structures (or structures with variable size and shape) in monogeneans increases the probability of switching to fish species of different body sizes, which is in accordance with the low degree of host specificity observed in D. vistulae (e.g. Šimková et al., 2001a, Benovics et al., 2018) and D. polylepidis (this study). Compared to endemic cyprinids, endemic leuciscids harbour species poor Dactylogyrus communities, though leuciscid Dactylogyrus species exhibit a

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higher degree of host specificity, with most species harbouring at least one specific Dactylogyrus species. The majority of new species recorded are morphologically similar, with Dactylogyrus sp. 2 and Dactylogyrus sp. 10, for example, sharing the 'ergensi' type of male copulatory organ but differing in the shape and size of the haptoral hard parts. Phylogenetic analyses and species delimitation analyses supported their species identities, i.e. nine new species were recognized. Species delimitation has received much attention recently, and numerous methods have now been developed that help identify species by using molecular data in a rigorous framework alongside morphological examination (Carstens et al., 2013, Zhang et al., 2013, Grummer et al., 2014). DNA-based delimitation methods have also been used to confirm or invalidate morphologically determined species, to identify cryptic species or highlight significant intraspecific genetic variability. The aforementioned diversity in haptoral part shape and size appears to be common in *Dactylogyrus* spp. and was previously hypothesised to be the result of adaptations to specific microhabitats (i.e. specific positions on fish gills; Šimková et al., 2001a, Jarkovský et al., 2004). Thus, minor morphological variabilities in the attachment organs may be observed in species with ongoing speciation parasitising phylogenetically distant hosts, as is the case in the Iberian Peninsula.

4.2 Phylogeny of endemic *Dactylogyrus*

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Phylogenetic reconstruction of *Dactylogyrus* parasitising Iberian cyprinoids suggests that Iberian *Dactylogyrus* belong to two well-supported phylogenetic lineages (Clade A and Clade B; Figure 2). One of these clades contains *Dactylogyrus* from endemic Cyprinidae only (representatives of five *Luciobarbus* species and *Barbus haasi*), while the second includes *Dactylogyrus* endemic to Iberian cyprinoids (both Cyprinidae and Leuciscidae) and *Dactylogyrus* parasitising cyprinoids from other parts of Europe. This was previously reported by Šimková et al. (2017) following analysis of phylogenetic relationships between *Dactylogyrus* from north-west Africa and those from the Iberian Peninsula, the authors

suggesting multiple origins for *Dactylogyrus* from both Mediterranean areas in association with the historical biogeography of their cyprinid hosts. Clade B comprises Dactylogyrus species described by el Gharbi et al. (1992), using morphological characteristics of the haptor and reproductive organs. According to their study (also supported by our own morphometric data), all these species achieve a small body size and display remarkably similar morphological features (i.e. sclerotised parts of attachment and copulatory organs), in accordance with their phylogenetic proximity. Previously, their description was based on small differences in the shape and size of sclerotised parts only (e.g. spiralisation of the male copulatory organ and size of haptoral sclerites). However, as has been previously documented, such variability may be present within single species and is common in the different monogenean taxa (e.g. Rohde and Watson, 1985, Boeger and Kritsky, 1988, Vignon and Sasal, 2010), including *Dactylogyrus* (Rahmouni et al., 2017). Nonetheless, the species status of each taxon in Clade B was supported by phylogenetic and species delimitation analyses, which was in concordance with their morphological determination. According to Šimková et al. (2017), Iberian *Dactylogyrus* species of this lineage are phylogenetically close to Dactylogyrus from north-west African Carasobarbus fritschii, suggesting different historical origins of Dactylogyrus in Clade B and Clade A. According to previous reports and the data presented here, each *Dactylogyrus* species within Clade B parasitises several endemic Luciobarbus species. Considering the monophyletic origin of Iberian Luciobarbus (Yang et al., 2015), its probable historical dispersion via northern Africa (Bianco, 1990, Doadrio, 1990, Zardova and Doadrio 1998), and the phylogenetic relatedness of *Dactylogyrus* from Clade B with north-west African Dactylogyrus (Šimková et al., 2017), we may postulate that these species originated on the *Luciobarbus* ancestor, and may have host-switched in the past to endemic north-west African Carasobarbus, subsequently dispersing to the Iberian Peninsula during its historical connection with North Africa. The high number of morphologically

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similar species exhibiting a low molecular divergence (e.g. D. bocageii, D. mascomai, D. guadianensis, D. lenkoranoïdes and D. doadrioi) suggests subsequent rapid speciation, most likely linked with the radiation of Luciobarbus across individual river basins within the Iberian Peninsula (Doadrio, 1998, Zardoya and Doadrio, 1998, Doadrio et al., 2002, Mesquita et al., 2007, Gante et al., 2015, Casal-López et al., 2017). Addition of *Dactylogyrus* species from Asian Capoeta (phylogenetically sister group to Iberian Luciobarbus; Yang et al., 2015) to phylogenetic reconstruction and assessing coevolutionary scenarios involving these parasites and their hosts may shed more light into origin of the Dactylogyrus of Iberian Luciobarbus and finally resolve the phylogenetic relationships within this group of Dactylogyrus. In contrast to *Dactylogyrus* from Clade B, the phylogenetic proximity of Iberian *Dactylogyrus* within Clade A to Central European and Balkan *Dactylogyrus* species supports their European origin. In accordance with the phylogeny proposed by Šimková et al. (2017), Dactylogyrus species from Iberian Luciobarbus form two well-supported lineages within Clade A, and cluster with Dactylogyrus from European Barbus. Two species within Clade A, D. balistae and D. legionensis, have large body-size, large haptoral sclerites and are missing the haptoral connective ventral bar (see el Gharbi et al., 1992). These species form a well-supported clade in sister position with another Iberian species, D. linstowoïdes. This clade is closely related to D. malleus, D. prespensis and D. petenvi, all host-specific parasites to European Barbus. In contrast to D. legionensis and D. balistae, these three species have small body size, similarly shaped small haptoral elements and a ventricular ventral bar (see Pugachev et al., 2009). Based on morphology, D. linstowoïdes represents the transient form between these two lineages, with the haptoral sclerites resembling Dactylogyrus of European Barbus and copulatory organs morphologically similar to Iberian species. Our results support a common origin for these species, with D. balistae, D. legionensis and D. linstowoïdes possibly

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evolving in Iberia from a common ancestor and thereafter switching to Luciobarbus, 488 following which D. balistae and D. legionensis secondarily lost their haptoral connective 489 ventral bar. 490 In this study, Leuciscids generally harboured poorer *Dactylogyrus* species communities than 491 cyprinids. However, due to the higher species richness of this fish family in the Iberian 492 Peninsula, a remarkably high species diversity was observed among their *Dactylogyrus* 493 parasites, and specifically among *Dactylogyrus* parasitising *Squalius* spp. and the genera 494 erected from *Chondrostoma* s.l.. Almost each genetic variant was supported as a species by 495 the species delimitation analysis. Dactylogyrus from Iberian leuciscids formed three major 496 497 phylogenetic lineages. The first comprised Dactylogyrus sp. 1 only, collected from two 498 endemic Squalius species, S. torgalensis and S. aradensis. Previous molecular phylogenetic studies suggested that these sister species have a basal position to other representatives of 499 500 Squalius in Iberia (Sanjur et al., 2003, Waap et al., 2011, Perea et al., 2016, Sousa-Santos et al., 2019). The distribution of S. torgalensis and S. aradensis is limited to the south-western 501 extremity of the Iberian Peninsula, and the same distribution range was found for 502 Dactylogyrus sp. 1. Extrapolating from the phylogenetic reconstruction, Dactylogyrus sp. 1 is 503 phylogenetically close to common *Dactylogyrus* species from European *Squalius* spp., i.e. *D*. 504 505 folkmanovae and D. nanoides (hypothesised to be genus specific according to Šimková et al., 2004 and Benovics et al., 2018), and probably represents an ancestral *Dactylogyrus* lineage 506 that has coevolved in Iberia with its endemic Squalius hosts. 507 The majority of endemic leuciscid *Dactylogyrus* formed a well-supported clade, with D. 508 caucasicus from Alburnoides spp. and D. ergensi from Chondrostoma spp. in sister position. 509 Benovics et al. (2018) has previously suggested that D. caucasicus originated from the 510 ancestor of D. ergensi by host-switching to Alburnoides. The species delimitation analysis 511 suggested the existence of nine potentially new species (Dactylogyrus sp. 2 to Dactylogyrus 512

sp. 10) phylogenetically related to D. ergensi (the species with the widest distribution range across Europe), which may indicate that endemic Dactylogyrus sp. 2 to Dactylogyrus sp. 10 also share a common ancestor with D. ergensi. As suggested by Robalo et al. (2007), the ancestor of *Chondrostoma* s.l. could have dispersed into Iberia prior to the Messinian period. when the host-specific ancestral *Dactylogyrus* species associated with these hosts most likely colonised Iberia. Our data suggest that the rapid radiation of *Chondrostoma*-related species promoted the speciation of their host-specific *Dactylogyrus*. Even if parasite phylogeny is not fully congruent with that of their hosts, all Iberian Dactylogyrus species, excluding Dactylogyrus sp. 8 (collected from Parachondrostoma species only distributed in rivers of the Mediterranean slope (Kottelat and Freyhof, 2007)), parasitise leuciscids in river basins of the Atlantic slope (distribution according to Kottelat & Freyhof, 2007, Robalo et al., 2007, Sousa-Santos et al., 2019). Considering that the distribution of cyprinoid species in Iberia is almost non-overlapping, the incongruence between host and parasite phylogenies could be the result of secondary contacts between fish host species, as recently documented in some Iberian rivers (e.g. Doadrio, 2001, Sousa-Santos et al., 2019). Dactylogyrus sp. 7, for example, was collected from two separate species, S. pyrenaicus and S. carolitertii. Sousa-Santos et al. (2019) and Waap et al. (2011) suggested that S. pyrenaicus consists of two different species, each associated with different river basins. Previous multilocus phylogenetic analyses (Sousa-Santos et al., 2019) have supported that S. pyrenaicus is paraphyletic, as genetic variants of this species from the Tagus and Colares basins were both grouped with S. carolitertii. Exactly the same pattern was observed among genetic variants of *Dactylogyrus* sp. 7, with individuals collected from S. pyrenaicus being in paraphyly and individuals from the River Colares grouped with individuals from S. carolitertii. A similar situation has also been observed in Dactylogyrus spp. from the Balkans, where the phylogenetic positions of two populations of D. vistulae within the D. vistulae clade (i.e. paraphyly) and molecular dissimilarity between

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the two populations (Benovics et al., 2018) supported the existence of two different *Alburnoides* species, as previously proposed by Stierandová et al. (2016).

In general, *Dactylogyrus* species diversity within the Iberian Peninsula appears to be associated with the historical dispersion of their cyprinoid hosts, with subsequent adaptive radiation following the peninsula's geographical isolation due to elevation of the Pyrenees (Munoz et al., 1986, Puigdefàbregas et al., 1992; Stange et al., 2016). At least two historical origins can be inferred for Iberian *Dactylogyrus*, each associated with the different dispersion routes proposed for cyprinoids (Banarescu, 1989, 1992, Doadrio, 1990, Doadrio & Carmona, 2003, Perea et al., 2010). Despite well-supported delineation between a multitude of endemic

Dactylogyrus species, the phylogenetic relationships between Dactylogyrus species do not

fully correspond to the phylogeny of their hosts, suggesting secondary contacts and host-

switching between endemic Iberian cyprinoids.

Figure 1. Map of collection localities in the Iberian Peninsula. The same codes are used in Table 1 as locality IDs. Figure 2. Phylogenetic tree of 70 Dactylogyrus haplotypes reconstructed by Bayesian inference (BI). The tree is based on combined parts of genes coding 18S and 28S rRNA, and the complete ITS1 region. Values between branches indicate posterior probabilities from BI and bootstrap values from ML analysis. Values below 0.80 (BI) and 50 (ML) are shown as dashes (-). The letters A-C represent specific well-supported lineages, as described in the Results section.

Figure 3. Results of species bPTP delimitation analysis applied to clades comprising endemic Dactylogyrus. Brackets at the terminal branches indicate different species, as suggested by BI and ML analyses.

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Table 1. List of cyprinoid species including localities of their collection and list of collected Dactylogyrus species from respective hosts.

Host species	N	ID	Locality	Dactylogyrus species	185	28S
Achondrostoma arcasii	15	S1	Chico River, flow of Palancia	D. polylepidis	MN365664	MN338198
	10	S2	Tera River	D. polylepidis	MN365665	MN338199
Achondrostoma occidentale	13	P1	Alcabrichel	Dactylogyrus sp. 2	MN365666	MN338200
				Dactylogyrus sp. 10	MN365667	MN338201
Barbus haasi	4	S 3	Beceite, Uldemo River	D. lenkoranoïdes	MN365668	MN338202
Iberochondrostoma almacai	19	P2	Torgal River, Mira basin	Dactylogyrus sp. 3	MN365669	MN338203
Iberocypris alburnoides	12	S4	Near Llera, Retin River	Dactylogyrus sp. 5	MN365670	MN338204
Luciobarbus bocagei	6	Р3	Colares (Portugal)	D. balistae	KY629344	MN338205
9			, 5,	D. bocageii	MN365671	KY629347
	10	S 5	Ucero River (Spain)	D. mascomai	no seq	MN338206
Luciobarbus comizo	5	S6	Peraleda de Zancejo, Zujar River	D. andalousiensis	MN365672	MN338207
				D. bocageii	MN365673	MN338208
				D. guadianensis	MN365674	MN338209
Luciobarbus graellsii	1	S 3	Beceite, Uldemo River	D. legionensis	MN365678	MN338210
			,	D. lenkoranoïdes	MN365676	MN338211
	5	S7	upstream Mella, tributary of	D. bocageii	MN365675	MN338212
			Materraña			
				D. lenkoranoïdes	MN365677	MN338213
				D. legionensis	MN365679	MN338214
				D. mascomai	MN365680	MN338215
Luciobarbus guiraonis	6	S8	Magro River	D. bocageii	MN365681	MN338216
				D. legionensis	KY629330	KY629350
				D. doadrioi	MN365682	KY629346
	4	S 9	Turia River	D. linstowoïdes	KY629329	KY629349
				D. mascomai	-	KY629348*
Luciobarbus sclateri	5	P2	Torgal River, Mira basin	D. andalousiensis	KY629331	KY629351
				D. bocageii	MN365684	MN338218
	10	S10	Benehavis, Guadalmina River	D. andalousiensis	MN365683	MN338217
				D. guadianensis	MN365685	MN338219
Parachondrostoma miegii	12	S 3	Beceite, Uldemo River	Dactylogyrus sp. 8	MN365686	MN338220
Parachondrostoma turiense	18	S 9	Turia River	Dactylogyrus sp. 8	MN365687	MN338221
Phoxinus bigerri	12	S5	Ucero River	D. borealis	MN365688	MN338222
Pseudochondrostoma duriense	9	S 5	Ucero River	Dactylogyrus sp. 9	MN365689	MN338223
				D. polylepidis	no seq	no seq
Pseudochondrostoma polylepis	10	P4	Alcoa, Fervenca	Dactylogyrus sp. 6	MN365690	MN338224
	15	Р3	Colares	-	-	-
Squalius aradensis	5	P5	Seixe	Dactylogyrus sp. 1	MN365691	MN338225
	6	P6	tributary of Seixe	-	-	-
Squalius carolitertii	15	P7	Arunca, Mondego basin (Vermoil)	Dactylogyrus sp. 7	MN365692	MN338226
			-	Dactylogyrus sp. 11	MN365693	MN338227
				D. polylepidis	-	_
Squalius pyrenaicus	5	Р3	Colares	Dactylogyrus sp. 7	MN365694	MN338228
•	5	S 6	Peraleda de Zancejo, Zujar River	Dactylogyrus sp. 7	MN365695	MN338229
Squalius torgalensis	10	P2	Torgal River, Mira basin	Dactylogyrus sp. 1	MN365696	MN338230
,			<u> </u>	Dactylogyrus sp. 4	MN365697	MN338231

N = number of processed fish individuals from respective locality, ID = code corresponding with localities marked in Figure 1 and codes in following tables, numbers in columns 18S and 28S correspond to sequence accession numbers for the respective genetic markers in GenBank; 18S = sequences of partial gene coding 18S rRNA combined with complete ITS1 region, 28S = sequences or partial gene coding 28S rRNA. Sequence not used in the present study is marked by asterisk (*). Dashes represent localities where no *Dactylogyrus* parasites were collected and/or missing sequences.

Table 2. Uncorrected pair-wise genetic distances between individuals from clade B (Figure 2) collected from different Barbus and Luciobarbus species in Iberian Peninsula.

No.	Dactylogyrus species	Host species	ID	1	2	3	4	5	6	7	8	9	10	11
1	D. bocageii	L. bocageii	Р3	Х										
2	D. bocageii	L. comizo	S6	-	Χ									
3	D. bocageii	L. graellsii	S7	0.001	0.001	Χ								
4	D. bocageii	L. guiraonis	S8	0.001	0.001	0.002	Χ							
5	D. bocageii	L. sclateri	P2	0.001	0.001	0.002	0.002	X						
6	D. lenkoranoïdes	B. haasi	S 3	0.016	0.016	0.017	0.015	0.017	Χ					
7	D. lenkoranoïdes	L. graellsii	S 3	0.016	0.016	0.017	0.015	0.017	-	Χ				
8	D. lenkoranoïdes	L. graellsii	S7	0.016	0.016	0.017	0.015	0.017	-	-	Χ			
9	D. guadianensis	L. comizo	S6	0.019	0.019	0.020	0.017	0.020	0.015	0.015	0.015	Χ		
10	D. guadianensis	L. sclateri	S10	0.017	0.017	0.019	0.016	0.019	0.014	0.014	0.014	0.001	Χ	
11	D. doadrioi	L. guiraonis	S8	0.017	0.017	0.019	0.016	0.019	0.006	0.006	0.006	0.016	0.015	Χ
12	D. mascomai	L. graellsii	S7	0.010	0.010	0.011	0.009	0.011	0.009	0.009	0.009	0.011	0.010	0.010

Distances are computed from the alignment of partial genes coding 28S rRNA. Identical sequences are marked by dashes (-). ID = code corresponding with localities marked in Figure 1 and specified in Table 1.

Table 3. Uncorrected pair-wise genetic distances between individuals from clade B (Figure 2) collected from different Barbus and Luciobarbus species in Iberian Peninsula.

No.	Dactylogyrus species	Host species	ID	1	2	3	4	5	6	7	8	9	10	11
1	D. bocageii	L. bocageii	Р3	Χ										
2	D. bocageii	L. comizo	S 6	0.004	X									
3	D. bocageii	L. graellsii	S7	0.015	0.020	X								
4	D. bocageii	L. guiraonis	S8	0.018	0.020	0.003	X							
5	D. bocageii	L. sclateri	P2	0.001	0.003	0.017	0.020	X						
6	D. lenkoranoïdes	B. haasi	S 3	0.095	0.096	0.085	0.088	0.094	Χ					
7	D. lenkoranoïdes	L. graellsii	S 3	0.095	0.096	0.085	0.088	0.094	-	X				
8	D. lenkoranoïdes	L. graellsii	S7	0.095	0.096	0.085	0.088	0.094	-	-	X			
9	D. guadianensis	L. comizo	S 6	0.078	0.080	0.070	0.073	0.077	0.094	0.094	0.094	Χ		
10	D. guadianensis	L. sclateri	S10	0.078	0.082	0.071	0.074	0.080	0.096	0.096	0.096	0.008	X	
11	D. doadrioi	L. guiraonis	S8	0.109	0.110	0.096	0.099	0.108	0.031	0.031	0.031	0.101	0.105	Х
12	D. mascomai	L. graellsii	S7	0.109	0.110	0.099	0.101	0.108	0.131	0.131	0.131	0.119	0.123	0.135

Distances are computed from complete sequences of ITS1 region. Identical sequences are marked by dashes (-).

ID = code corresponding with localities marked in Figure 1 and specified in Table 1.

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Table 4. Uncorrected pair-wise genetic distances between individuals from clade B (Figure 2) collected from different Barbus and Luciobarbus species in Iberian Peninsula.

No.	Dactylogyrus species	Host species	ID	1	2	3	4	5	6	7	8	9	10	11
1	D. bocageii	L. bocageii	Р3	Х										
2	D. bocageii	L. comizo	S6	-	Х									
3	D. bocageii	L. graellsii	S7	-	-	X								
4	D. bocageii	L. guiraonis	S8	-	-	-	Χ							
5	D. bocageii	L. sclateri	P2	-	-	-	-	X						
6	D. lenkoranoïdes	B. haasi	S 3	0.008	0.008	0.008	0.008	0.008	Χ					
7	D. lenkoranoïdes	L. graellsii	S 3	0.008	0.008	0.008	0.008	0.008	-	Х				
8	D. lenkoranoïdes	L. graellsii	S7	0.008	0.008	0.008	0.008	0.008	-	-	Х			
9	D. guadianensis	L. comizo	S6	0.002	0.002	0.002	0.002	0.002	0.010	0.010	0.010	Х		
10	D. guadianensis	L. sclateri	S10	0.002	0.002	0.002	0.002	0.002	0.010	0.010	0.010	-	X	
11	D. doadrioi	L. guiraonis	S8	0.008	0.008	0.008	0.008	0.008	0.004	0.004	0.004	0.010	0.010	X
12	D. mascomai	L. graellsii	S7	_	_	-	-	-	0.008	0.008	0.008	0.002	0.002	0.008

Distances are computed from the alignment of partial genes coding 18S rRNA. Identical sequences are marked by dashes (-). ID = code corresponding with localities marked in Figure 1 and specified in Table 1.





