1	Original article
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3	Running header: Biogeography of Mediterranean Deronectes diving beetles
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5	Reconstructing ancient Mediterranean crossroads in Deronectes diving
6	beetles
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## 25 ABSTRACT

26 Aim To reconstruct the evolutionary history of a genus of freshwater beetle with a pan-

27 Mediterranean distribution, to test classic hypotheses which proposed a Miocene origin

28 for groups with high biodiversity in the Iberian and Anatolian peninsulas.

29 **Location** Mediterranean basin.

30 Methods We sequenced four mitochondrial and one nuclear gene from 51 specimens of 31 30 of the ca. 60 extant species of *Deronectes* (Dytiscidae), all typical of mid-mountain 32 streams from North Africa and Iberia over most of Europe to the Middle East. We used 33 Maximum Likelihood, Bayesian probabilities with an a priori evolutionary rate and a 34 Dispersal-Extinction-Cladogenesis model to reconstruct their biogeographical history. 35 **Results** *Deronectes* has two major lineages which originated in the mid Miocene; one 36 including mostly eastern and another mainly western and central Mediterranean species. 37 From these two areas range expansions, mainly at the end of the Miocene and beginning 38 of the Pliocene, resulted in the many species groups and some of the extant species of the 39 genus. Most of the current diversity and distributions are, however, of Plio-Pleistocene 40 origin, particularly in widespread European species.

41 Main conclusions In line with traditional hypotheses, we found an ancient division 42 between eastern and western Mediterranean lineages of Deronectes, likely resulting from 43 the isolation of Europe west of the Alps from the Balkans and Anatolia during the early-44 middle Miocene. The history of the genus was strongly influenced by major geological 45 and climatic events, with successive cycles of fragmentation and subsequent eastward 46 and westward range expansions, resulting in a steady accumulation of species across the 47 basin. Most of these range movements took place through the north side of the 48 Mediterranean, with only local displacements in the south during the Messinian and a 49 recent (Pleistocene) colonization of the Italian peninsula, which remained largely 50 submerged through most of the genus' evolutionary history.

51

#### 52 Keywords

Biodiversity hotspot, dispersal, diversification, Dytiscidae, Mediterranean, Messinian
salinity crisis, phylogeny.

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#### 58 INTRODUCTION

59 The Mediterranean region, with its complex geological history, is an ideal system to 60 study the effects of palaeogeographical events on evolutionary diversification. The region 61 has had a 'reticulated' biogeographical history, in which the constituent landmasses have 62 repeatedly split, collided, and split again in different configurations over time 63 (Rosenbaum et al., 2002; Meulenkamp & Sissingh, 2003; Popov et al., 2004), resulting in 64 repeated episodes of vicariance and dispersal (Oosterbroek & Arntzen, 1992; Sanmartín et al., 2001). Whilst the geological evolution of the basin is relatively well understood, 65 66 the detailed geographical and temporal origins of most Mediterranean organisms remain 67 unknown, especially in diverse groups such as insects.

68 Traditional hypotheses proposed a Miocene origin for many terrestrial and 69 freshwater Mediterranean lineages, with close relationships between the fauna at the two 70 extreme ends of the basin – the so-called Kiermack disjunction (see e.g. Brehm, 1947 on 71 Iberian and Balkan plants, Bănărescu, 1991 on the Mediterranean freshwater fauna, or 72 Ribera & Blasco-Zumeta, 1998 on insects with disjunct distributions between the steppe 73 areas of NE Spain and those of the eastern Mediterranean and central Asia). Distribution 74 throughout the Mediterranean region became possible for terrestrial and freshwater taxa 75 during the Late Oligocene-Early Miocene, after the formation of a continuous landmass 76 connecting western Europe with the area roughly corresponding to the Balkans and 77 Turkey, separating the Tethys and Paratethys Oceans (Rögl & Steininger, 1983; 78 Oosterbroek & Arntzen, 1992). During the Miocene, the re-establishment of occasional 79 marine connections between the Tethys and Paratethys and successive landmass suture 80 events between the Eastern and Western Mediterranean Basins likely resulted in the 81 diversification of many Mediterranean groups (Oosterbroek & Arntzen, 1992; Montreuil, 82 2007). Alternatively, other biogeographical studies consider North Africa and the 83 Gibraltar Strait, which closed during the Messinian salinity crisis (end Miocene), to be an 84 alternative dispersal route by which lineages could have achieved circum-Mediterranean 85 distributions (e.g. Sanmartin, 2003).

Many of these early hypotheses were based on the presence of the same, or very closely related species, on both sides of the Mediterranean in similar ecological conditions (see Ribera & Blasco-Zumeta, 1998 for a review), but without a wider phylogenetic context. Similarly, and given the lack of fossils in most Mediterranean groups, the estimated temporal origin of these relationships were based on circumstantial evidence alone.

The widespread use of molecular data to obtain reliable, calibrated phylogenies
has resulted in a proliferation of studies on Mediterranean lineages (e.g. Levy *et al.*,
2009; Santos-Gally *et al.*, 2012 or Condamine *et al.*, 2013). There are, however, very few
data on freshwater invertebrates encompassing the entire Mediterranean area, most works
to date focussing on only parts of the basin (e.g. Trizzino *et al.*, 2011 for northern
Mediterranean freshwater beetles, or Solà *et al.*, 2013 for eastern Mediterranean
freshwater planarians).

99 Whilst there remains a dearth of detailed analyses of lineages with wide 100 Mediterranean distributions, general hypotheses on the origin and composition of the 101 Mediterranean biota as a whole can only be tested by investigating such taxa. In this work 102 we study one of these lineages, the diving beetle genus Deronectes Sharp (family 103 Dytiscidae). With ca. 60 described species, *Deronectes* has a predominantly 104 Mediterranean distribution, ranging from North Africa and the Iberian Peninsula over 105 most parts of Europe and the Middle East, with some species reaching central Asia. 106 *Deronectes* are poor dispersers, with species usually restricted to relatively small 107 geographical ranges particularly in mountain regions, making them eminently suitable for 108 biogeographical reconstructions. There are in addition some widespread species with 109 continental-scale distributions (Abellán & Ribera, 2011), demonstrating their potential for 110 range expansion. *Deronectes* usually live amongst gravel, stones or submerged tree roots 111 in small streams with sparse vegetation (Fery & Brancucci, 1997).

112 Previous work based on mitochondrial genes and with incomplete sampling 113 identified two main lineages within the genus, mostly corresponding to species with a 114 western (Iberian Peninsula) or eastern (Anatolia and Middle East) distribution (Ribera et 115 al., 2001; Ribera, 2003; Abellán & Ribera, 2011), but the precise relationships of the 116 species, their geographical origin and the temporal framework of their diversification 117 remained obscure. In this work we use a comprehensive molecular phylogeny of the 118 genus to reconstruct its biogeographical history, the geographical origin of major lineages 119 and the events that led to their current distributions through the Mediterranean basin.

120

# 121 MATERIAL AND METHODS

122

#### 123 **Taxon sampling**

124 Deronectes contains 58 species and four subspecies (Nilsson & Hájek, 2015), most of

125 them revised by Fery & Brancucci (1997) and Fery & Hosseinie (1998). These authors

126 divided the genus into 10 groups based on morphology, to which Hájek *et al.* (2011)

- 127 added an 11<sup>th</sup> for a single species from Turkey (*D. ermani* Hájek *et al.*) (See Appendixes
- 128 S1a,b in Supporting Information for a checklist of the species and subspecies with
- 129 distributions). We studied 51 specimens of 30 species of *Deronectes*, with an emphasis
- 130 on the western clade (20 out of 24 known species) but including representatives of all
- recognised species groups with the exception of the recently described *D. ermani*. We
- 132 also included three of the four described subspecies (Appendix S1c). For some species
- 133 more than one specimen was included to detect possible unrecognised variation. We used
- 134 37 species of closely related genera of Hydroporini as outgroups, following the
- 135 phylogenies of Dytiscidae in Ribera *et al.* (2008) and Miller & Bergsten (2014).
- 136

# 137 **DNA extraction and sequencing**

138 Specimens were collected and preserved in absolute ethanol directly in the field. We 139 obtained the DNA non-destructively, either with standard phenol-chloroform extraction 140 or commercial kits (mostly DNeasy Tissue Kit, Qiagen GmbH, Hilden, Germany and 141 Charge Switch gDNA Tissue Mini Kit, Invitrogen, Carlsbad, CA, USA), following 142 manufacturer's instructions. Voucher specimens and DNA extractions are kept in the 143 collections of the Institut de Biología Evolutiva, Barcelona (IBE), Museo Nacional de 144 Ciencias Naturales, Madrid (MNCN) and Natural History Museum, London (NHM). 145 Six gene fragments from five different genes (four mitochondrial and one nuclear)

146 were obtained in four different amplification reactions: 1) 5'end of the Cytochrome c

147 Oxidase Subunit 1 gene (the barcode fragment, Hebert et al., 2003, COI-5'); 2) 3'end of

- 148 Cytochrome c Oxidase Subunit 1 (COI-3'); 3) 3'end of 16S rDNA plus tRNA transfer of
- Leucine plus 5'end of NADH dehydrogenase subunit 1 (nad1) (16S); 4) an internal
- 150 fragment of the nuclear gene Histone 3 (H3) (see Appendix S2a for details on primers
- 151 used and Appendix S2b for PCR conditions).
- 152

#### 153 **Phylogenetic analyses**

Edited sequences were aligned using MAFFT v.6 with the G-INS algorithm and defaults for other parameters (Katoh & Toh, 2008). We used jModeltest 0.1.1 (Posada, 2008) to

- estimate the evolutionary model that best fitted the data for each gene separately, using
- 157 AIC (Akaike Information Criterion) scores as selection criteria and default values for
- 158 other parameters (Katoh & Toh, 2008). To infer the phylogeny of *Deronectes* we used
- 159 Bayesian probabilities (Bp) and Maximum Likelihood (ML).

160 For Bayesian analyses we used MrBayes 3.2 (Ronguist *et al.*, 2012) implementing 161 the most similar evolutionary models to those selected by jModeltest. For the analyses of 162 combined mitochondrial and nuclear data we used five partitions corresponding to COI-163 5', COI-3', 16S rRNA+tRNA-Leu, nad1 and H3. We also analysed the mitochondrial 164 (with 4 partitions as above) and nuclear data separately. The program was left running 165 until we obtained a sufficient number of trees after the two independent runs converged, 166 according to the ESS (Effective Sample Size) and PSRF (Potential Scale Reduction 167 Factor) criteria as estimated in Tracer v1.5. (Drummond & Rambaut, 2007) and MrBayes 168 respectively. Convergence and burn-in values were estimated visually after examining a 169 plot of the standard deviation of split frequencies between the two simultaneous runs.

For ML analysis we used a fast approximate algorithm as implemented in RAXML 7.1 (Stamatakis *et al.*, 2008) using GTR+G as an evolutionary model and the same partitions as in MrBayes. The optimum topology was that of the best likelihood amongst 100 replicates, and node support was estimated with 1000 bootstrap replicates using the CAT approximation (Stamatakis *et al.*, 2008).

175

#### 176 Estimation of ages of divergence

177 We obtained an estimate of divergence dates amongst species with BEAST v1.7 178 (Drummond & Rambaut, 2007). There are no fossils or unambiguous biogeographical 179 events that could be used to calibrate the phylogeny of *Deronectes*, so we used an *a priori* 180 substitution rate for the combined mitochondrial sequence of 0.01 substitutions/site per MY (million years) (standard deviation 0.002), similar to that obtained in related beetle 181 182 groups for the same combination of mitochondrial protein coding and ribosomal genes 183 (Papadopoulou et al., 2010; Ribera et al., 2010; Andujar et al., 2012). We excluded the 184 nuclear sequence (H3), and to ensure that the topology obtained with mitochondrial 185 sequences was the same as that obtained with the combined matrix we constrained all 186 well supported nodes (with a posterior probability in MrBayes > 0.95 and a bootstrap 187 support in RAxML > 70%) after deleting the outgroups. We used a GTR+I+G 188 evolutionary model, an uncorrelated lognormal relaxed clock and a Yule speciation 189 model. We executed two independent runs with the same settings that were allowed to 190 run until they had converged and the number of trees was sufficient according to ESS 191 values, as measured in Tracer v1.5. The consensus tree of the two runs was compiled 192 with Tree Anotator v1.7 (Drummond & Rambaut, 2007).

#### 194 Diversification

- 195 To have an estimation of possible changes in diversification rates through the evolution 196 of the group we used a log-lineage through time approach (LTT) (Barraclough & Nee, 197 2001). Only the western clade could be studied, with an almost complete taxon sampling 198 (20 out of 22 species). We used the R library 'ape' (Paradis et al., 2004) to compile the 199 LTT plot using the ultrametric tree obtained in BEAST after deleting duplicated 200 specimens of monophyletic taxa. We used the  $\gamma$ -statistic (Pybus & Harvey, 2000) for 201 measuring the relative timing of diversification, i.e. whether there is a constant 202 diversification rate through the tree, or the interior nodes are closer to the tips, or to the 203 root, than expected under a pure birth process. The  $\gamma$  -values of complete reconstructed 204 phylogenies follow a standard normal distribution. If  $\gamma < 0$ , the internal nodes can be said 205 to be closer to the root than expected under a pure birth process, and vice versa (Pybus & 206 Harvey, 2000).
- 207

#### 208 Ancestral area reconstruction

- 209 To estimate ancestral areas of distribution we used a dispersal-extinction-cladogenesis 210 model implemented in the package Lagrange c++0.1, a ML inference model in which 211 parameters are estimated for rates of migratory events between areas (range expansions) 212 and local extinctions within areas (range contraction) (Almeida et al., 2012). Lagrange 213 considers branch lengths and allows the definition of a number of areas with an 214 associated probability matrix of dispersal between them (Ree & Smith, 2008). We 215 considered eight geographical areas, based on the current distribution of *Deronectes* 216 species: A) southeastern Iberian Peninsula including Mallorca; B) centre and north of the 217 Iberian Peninsula; C) Italy (including Sicily) and southeastern France; D) Corsica and 218 Sardinia; E) Balkan peninsula; F) Turkey; G) northern and central Europe; and H) 219 Maghreb (See Fig. 1 for the distribution of the main lineages and Appendix S1a and 220 Appendix S1b for the distribution of the species of *Deronectes*). We used the tree 221 obtained in BEAST after pruning duplicated specimens. In all analyses a maximum of 222 four possible ancestral areas was allowed.
- 223

For the reconstruction of ancestral areas we used three different time slices 224 corresponding to the Pleistocene, Pliocene and Miocene, and a different 225 palaeogeographical scenario for each: present, Piacenzian/Gelasian (1.8-3.4 Ma) and Late 226 Tortonian (7-8 Ma) respectively, adapted from Meulenkamp & Sissingh (2003). For each 227 scenario we identified the geographical barriers between our eight pre-defined areas, and

- assigned probabilities of dispersal to the land or sea barriers in different combinations,
- including a null model with all probabilities equal to 1 (Table 1). We used the likelihood
- 230 of the reconstruction to select the model best fitting the current distribution of the species.
- A difference equal to or greater than 2 log-likelihood units was considered significant
- 232 (Ree *et al.*, 2005; Ree & Smith, 2008).

To account for topological uncertainty we used the best settings as selected above in a Bayes-Lagrange analysis with a selection of 1000 trees from amongst the last 50,000 trees of the stationary period (post burn-in) of the BEAST analysis. Using a custom script in R and a spreadsheet we parsed the output and estimated the frequency of each

- combined area reconstruction for the nodes present in the consensus tree, as well as the
- 238 individual frequency of each of the eight areas.
- 239

#### 240 **RESULTS**

## 241 **Phylogenetic analyses**

There were no length differences in protein coding genes, and the length of ribosomal genes ranged between 685-693 bp in the ingroup. The MrBayes runs of combined and nuclear H3 analyses reached a standard deviation of split frequencies below 0.01 at 15 and 4 million generations (MG) respectively, and below 0.005 at 18.5 MG in the analysis of the mitochondrial matrix. These were considered the burn-in fractions, after which analyses were left to run until they reached convergence (at 23, 10 and 30 MG respectively).

Differences between topologies obtained with Bp and ML were minimal, and affected only the degree of resolution and support of some nodes (Fig. 2; Appendix S3a and S3b). The monophyly of *Deronectes* was strongly supported, as well as its separation into two major clades, one including species predominantly distributed in the eastern Mediterranean ("eastern clade"); and a clade of species predominantly distributed in the western and central Mediterranean ( "western clade") (Figs 1 and 2).

255 The eastern clade was further subdivided into two species groups as follows:

1) *D. parvicollis* group, including species from large parts of Asia, Turkey and the
Caucasus to southern Siberia and Central Asia. Only one species, *D. parvicollis*(Schaum), extends into Europe (Balkans).

2) *D. latus* group, four species from eastern Turkey to the Iberian Peninsula and
throughout central and northern Europe, including the British Isles and Scandinavia. This

group included the most widespread species of the genus, *D. latus* (Stephens), ranging
over most of Europe north of the Pyrenees and the Apennines.

Within the western clade we recovered four well-supported species groups plus two isolated species (*D. sahlbergi* Zimmermann and *D. doriae* Sharp), but the relationships amongst them were not well resolved. (Fig. 2). These four clades were:

266 1) *D. opatrinus* group, including mostly species endemic to the Iberian Peninsula,
267 with only one (*D. hispanicus* (Rosenhauer)) reaching northern Morocco and two (*D. hispanicus* and *D. opatrinus* (Germar)) southern France.

269 2) *D. aubei* group, with three species and one subspecies distributed from the
270 Cantabrian mountains in northwestern Spain to Sicily, including the Alps and southern
271 Germany.

3) *D. moestus* group, including species with a predominantly western
Mediterranean distribution, from the Maghreb and the Iberian Peninsula to the Balkans
through southern France, Italy and Sicily. The two missing north African species from the
western clade (*D. perrinae* Fery & Brancucci and *D. peyerimhoffi* (Régimbart)) most
likely belong here, as they are morphologically very similar to *D. moestus* and *D.
<i>fairmairei* (Leprieur) respectively (Fery & Brancucci, 1997).

4) *D. platynotus* group, including two species and two subspecies from the
Balkans, Central Europe and northwest Iberia.

280 Most species with more than one sequenced specimen were monophyletic, with 281 some exceptions. There were three paraphyletic complexes of closely related species 282 (Fig. 2): 1) the widespread D. latus, with the Iberian endemic D. angusi Fery & 283 Brancucci nested within it; 2) the D. aubei group, with one clade west of the Rhone river, 284 from the French Massif Central to the Cantabrian Mountains including D. a. sanfilippoi 285 Fery & Brancucci and D. delarouzei (Jacquelin du Val), and another east of the Rhone, 286 from the Alps to Sicily with D. a. aubei (Mulsant) and D. semirufus (Germar) (Appendix 287 S1b); and 3) D. ferrugineus Fery & Brancucci and D. wewalkai Fery & Fresneda, both 288 Iberian endemics. There was also one case (D. moestus) with a deep intraspecific 289 divergence, with the specimen from Morocco (MNCN-AI937) sister to D. brannani 290 (Schauffus) (a Mallorcan endemic), and both sister to the other specimens of D. moestus, 291 from northern Spain to Bulgaria, including specimens of the two recognised subspecies 292 (Fig. 2).

The analysis of the nuclear sequence (H3) showed lower resolution and an absence of support at some nodes, with polytomies in some groups (e.g. *D. moestus* or *D*.

- *latus*) but with a topology compatible with that obtained from the mitochondrial
- sequence, with a single exception (Appendixes S3c and S3d). Whilst with the
- 297 mitochondrial sequence the two subspecies of *D. aubei* were recovered as paraphyletic,
- and respectively sisters to the geographically closest species of the group, the nuclear
- sequence recovered a monophyletic *D. aubei* as sister to the other two species of the
- 300 group (D. delarouzei and D. semirufus).
- 301

#### 302 Estimation of ages of divergence and mode of diversification

- The origin of extant species of *Deronectes* and the separation of the Eastern and Western clades was estimated to have occurred in the Middle Miocene (ca. 14 Ma, with a 10.0-17.5 95% confidence interval) (Fig. 3). The origin of the well-supported species groups was estimated to have occurred over a relatively short time period at the end of the Miocene and beginning of the Pliocene.
- Some extant species originated during the Pleistocene, particularly within the *D*. *latus*, *D. aubei* and *D. platynotus* groups, but most species of the Iberian clade (*D. opatrinus* group) and the *D. moestus* group were estimated to be of Pliocene or even late
  Miocene origin. Observed intraspecific variation was also limited to a Pleistocene origin,
  except in the case of *D. moestus* (Fig. 3).
- 313 The LTT plot (Fig. 4) reflecting the temporal pattern of diversification of species 314 of the western clade showed a steady increase in lineages over time. The  $\gamma$ -statistic was 315 negative (-0.96) but not significantly different from zero (*P*= 0.33).
- 316

## 317 Ancestral area reconstruction

318 Amongst models tested, the best likelihood in Lagrange was found for the geography of 319 the Pleistocene, assigning the same penalty value for dispersal through one of the pre-

- defined land areas or a sea barrier shorter than 100 Km, and a zero probability of
- 321 dispersal over marine barriers longer than 100 Km or through two or more land areas
- 322 (Table 1). These settings were used to reconstruct ancestral areas using the 1000 post-
- 323 burnin trees in BEAST. The use of Pliocene or Miocene palaeogeographical scenarios,
- 324 either alone or in combination, resulted in significantly worse likelihoods (Table 1).
- 325 Within the same geographical scenario, results were less sensitive to small changes in the
- 326 values of the cost of dispersal, with differences of less than two units logL, but always
- 327 significantly better than the null model of all probabilities equal and equal to one (Table
- 328 1). In any case, results were very similar for all paleaogeographical scenarios, cost

matrixes or topologies, with differences only in the relative proportion of some of the
 reconstructed areas of the deeper nodes, including a large number of species with wide
 geographical distributions.

Most of the nodes present in the consensus tree had a well supported reconstructed ancestral area, with only 5 out of 34 lacking at least one area present in more than 90% of reconstructions, and only 2 (10 and 11 in Fig. 3) where the most likely area was present in less than 80% of the 1000 trees (Table 2). Most of the nodes were also well resolved, with 22 (65%) with only two areas with a frequency higher than 90%, and only three with four areas (the maximum number allowed in the settings) with a frequency higher than 90% (Table 2).

The eastern clade of *Deronectes* was unequivocally reconstructed as having an origin in Turkey (region F), with an expansion to Italy and the Balkan peninsulas (areas C and E) at the origin of the *D. latus* group (node 26 in Fig. 3). There was a subsequent expansion to central and northern Europe and the Iberian Peninsula (areas B and G) during the Pleistocene, at the origin of *D. latus* and *D. angusi* respectively (Fig. 3; Table 2).

345 The reconstructed origin of the western clade was more ambiguous. Although 346 centred in the southwestern Mediterranean region, only central and northern Iberia and 347 Corsica and Sardinia (areas B and D) occurred at a frequency higher than 90% in the 348 1000 trees, but the Maghreb (area H) also had a high frequency (89%, Fig. 3; Table 2). 349 Within the western clade, the *D. opatrinus* group (node 4) had a well supported central 350 and north Iberian origin (area B), with an expansion to the southeastern Iberian Peninsula 351 (area A) at the end of the Messinian (nodes 7 and 8), at the origin of the endemic species 352 D. algibensis Fery & Fresneda and D. depressicollis (Rosenhauer) (Fig. 3). This lineage 353 experienced further expansions during the Pliocene to north Africa and within the Iberian 354 Peninsula.

The *D. moestus* group (node 20) was reconstructed as most likely having a Maghrebian or Corso-Sardinian origin (areas H and D), with subsequent expansions to SE Iberia during the Messinian and the rest of the Iberian Peninsula and Mallorca during the Pliocene (Fig. 3; Table 2).

The reconstructed origins of the *D. aubei* (node 12) and *D. platynotus* groups (node 18) were more ambiguous. For the first, three areas had a 100% frequency in the set of 1000 trees: central and north Iberia, central and north Europe and the Italian peninsula (areas B, C and G; Table 2). The range expansion of this group apparently took

- 363 place between the late Miocene and middle Pleistocene. Similarly, two areas were
- 364 reconstructed with a frequency of more than 90% at the origin of the *D. platynotus* group:
- 365 central and northern Iberia and the Balkan peninsula (areas B and E) (Fig. 3; Table 2).
- 366 Two more expansions to the east (Balkan peninsula and Turkey) were unambiguously
- 367 reconstructed at the origin of *D. doriae* and *D. sahlbergi*, during the Messinian (Fig. 3;
- 368 Table 2).
- 369

### 370 **DISCUSSION**

371 A Miocene basal split in Deronectes

372 According to our results it seems highly likely that the ancestor of extant *Deronectes* was 373 found on the northern shores of the Mediterranean during the early Miocene. Our 374 estimation of the age of the basal split between eastern and western lineages is in good 375 agreement with the increased isolation of Europe west of the Alps from the Balkans and 376 Anatolia during the middle Miocene. During this time, climate change and tectonic 377 movements associated with Carpathian uplift resulted in a succession of sea level 378 fluctuations in the central and eastern Paratethys basins (Ter Borgh et al., 2014). The 379 extension of the Carpathian Foreland in a narrow deep-sea basin towards the west 20-15 380 Ma (Dercourt et al., 1985; Meulenkamp & Sissingh, 2003) could have contributed to the 381 isolation of strictly freshwater species in the two areas. The basal split in Deronectes 382 agrees with the estimated age of similar western and eastern lineages within the 383 freshwater beetle genus Hydrochus (Hidalgo-Galiana & Ribera, 2011), and with many 384 other comparable splits within Mediterranean lineages, although in most cases no age 385 estimates are available (see e.g. examples in Oosterbroek & Arntzen, 1992).

386 Of the two main lineages of *Deronectes*, the eastern clade was unambiguously 387 reconstructed as having an origin in Anatolia, but the precise origin of the western clade 388 was more uncertain due to the wide geographical ranges of some species within it and the 389 lack of statistical support for the nodes connecting the main groups. During most of the 390 Miocene, the Italian peninsula was mostly submerged or partly merged with the future 391 Balkan and Anatolian peninsulas (Dercourt et al., 1985; Rosenbaum et al., 2002; 392 Meulenkamp & Sissingh, 2003; Popov et al., 2004), something which could explain the 393 absence of ancient lineages in this area, apparently colonised by *Deronectes* only during 394 the Pleistocene.

395

#### **Tortonian disaggregation**

397 We traced the origin of the main species groups within *Deronectes* to the late Tortonian 398 and the transition to the Messinian, in most cases with relatively poor topological 399 resolution, suggesting a rapid succession of isolation events. The diversification of the 400 western clade involved successive splits between the Iberian Peninsula, north Africa and 401 Corsica and Sardinia, resulting in the main species groups recovered in our phylogeny. 402 These species groups were mostly in agreement with those obtained with previous 403 molecular (Ribera et al., 2001; Ribera, 2003; Abellán & Ribera, 2011) and morphological 404 analyses (Fery & Brancucci, 1997; Fery & Hosseinie, 1998). The main differences in our 405 study were the recognition of an Iberian clade, divided into several groups of species not 406 previously thought to be closely related (Fery & Brancucci, 1997), and the composition 407 of the D. moestus group (Appendix S1a).

408 The Tortonian was characterised by strong tectonic activity and changes in sea 409 level in the area between southeastern Iberia and the Maghreb (Alvinerie et al., 1992; 410 Martín *et al.*, 2009), favouring vicariance events that led to allopatric speciation in a 411 number of groups (e.g. Jolivet et al., 2006; Hidalgo-Galiana et al., 2011; Faille et al., 412 2014). The development of more continental climates, with changes in precipitation 413 seasonality, have been associated with the appearance of more open vegetation in some 414 areas in southern Europe after the Middle Miocene Optimum, particularly in the 415 southwestern Mediterranean (e.g. Barrón et al., 2010; Casas-Gallego et al., 2015). The 416 causal relationship between climate and vegetation changes is, however, not well 417 established (Bruch et al., 2011).

418 The origin of species of the western group in the Balkans, east of the Paratethys 419 basin, is more uncertain. In the Lagrange analyses they were reconstructed as having a 420 western origin, requiring subsequent range expansion towards the east. However, the 421 relative lack of support allowed an alternative scenario (as seen in the topology of Fig. 2), 422 with the eastern-most species within the western clade (D. platynotus group plus the 423 isolated D. doriae and D. sahlbergi) sister to the remaining western lineages, something 424 which would not require a range expansion from the west, but instead a western 425 migration of the Iberian member of the group (D. costipennis) in the Plio-Pleistocene.

426

## 427 Messinian crossroads

428 The onset of the Messinian salinity crisis 5.96-5.33 Ma ago and the establishment of new

- 429 land corridors seem to have facilitated the expansion of some species of *Deronectes*,
- 430 although these movements were relatively local and mostly centred in the southwest of

the Mediterranean basin. After the closing of the Tortonian sea corridors between

432 mainland Iberia, the Betic-Rifian area and mainland North Africa (Martín *et al.*, 2009)

433 there were expansions of the Iberian lineages towards the southeast (D. opatrinus group)

434 and of the *D. moestus* group towards the northwest and the Balearic islands, both likely

435 crisscrossing the Gibraltar area. Both expansions continued during the Pliocene, some

436 species of the Iberian clade towards north Africa, and species of the *D. moestus* group437 towards southern Europe.

In the east, range movements associated with the Messinian likely include the
crossing of the Bosphorus strait by *D. doriae*, currently known only from Turkey,
Armenia and Iran (Fery & Brancucci, 1997; Nilsson & Hájek, 2015), and possibly *D. sahlbergi*, known from Turkey but also from Greece (Nilsson & Hájek, 2015), meaning
that Asian populations may have a relatively recent origin.

443 We did not find any evidence of large scale range movements during the Messinian in the south or central Mediterranean basin, or along the coast of the Sarmatic 444 445 Sea (the Paratethys) which could correspond to the "lago mare" dispersal routes proposed 446 by e.g. Bianco (1990). During the Messinian, changes in climate or vegetation on the 447 northern side of the Mediterranean were not very pronounced (Favre et al., 2007), but it 448 is possible that the newly formed land corridors did not have the ecological conditions to 449 allow the dispersal of species restricted to fast flowing freshwater streams (Roveri et al., 450 2014).

451

## 452 The establishment of current distributions in the Plio-Pleistocene

453 The best model for the ancestral reconstruction in Lagrange was that reflecting present 454 geography, suggesting that current distribution patterns within the genus are largely 455 dominated by range movements since the Messinian. This is in contrast with the results 456 obtained with other groups of very poorly dispersing species, such as Trechus fulvus 457 group ground beetles (Carabidae), which include many subterranean taxa and have a 458 distribution still dominated by their late Miocene biogeography (Faille et al., 2014). Also 459 supporting the importance of the Plio-Pleistocene in the evolutionary history of 460 Deronectes is the high number of species estimated to have originated during this period, 461 reflected by the constancy of the diversification rate estimated from the LTT plot. 462 Exceptions are an island endemic (D. lareynii from Corsica), the Moroccan D. theryi and 463 the isolated eastern species *D. doriae* and *D. sahlbergi*, all apparently of late Miocene 464 origin. Most of the geographically restricted species in the Iberian clade also have a

- relatively ancient (Pliocene) origin, most likely driven by vicariance between the main
  mountain systems (Ribera, 2003). There are other known examples of freshwater
  Coleoptera with similar biogeographical patterns for example, in the Hydraenidae
- 468 (subgenus *Enicocerus*) and Hydrochidae (genus *Hydrochus*) Iberian endemics are mostly
- 469 of late Miocene origin (Ribera *et al.*, 2010; Hidalgo-Galiana & Ribera, 2011).

470 Such an ancient origin is, however, not a generalised pattern, as in other groups of freshwater Coleoptera most Iberian endemics, many of them restricted to the same 471 472 mountain systems as endemic Deronectes, are apparently of Pleistocene origin. This is 473 the case for most species of the "Haenydra" lineage (Ribera et al., 2010; Trizzino et al., 474 2011), some species groups of *Limnebius* (Abellán & Ribera, 2011) (both Hydraenidae), 475 and several Iberian endemic diving beetles from different genera (Dytiscidae, Ribera, 476 2003; Ribera & Vogler, 2004). Similarly, all speciation events within some groups of 477 Deronectes are of Pleistocene origin (D. aubei, D. platynotus and D. latus groups), which 478 are also the groups including most non-monophyletic species in our analyses. Most of 479 these can be explained either by incomplete lineage sorting due to their recent divergence 480 (D. ferrugineus-wewalkai, D. latus-angusi) or the presence of previously unrecognised 481 species-level diversity (D. moestus complex), except for the discordance between 482 mitochondrial and nuclear data within the D. aubei group. Incomplete lineage sorting is 483 not expected to leave any predictable biogeographical pattern (Funk & Omland, 2003), so 484 is unlikely to be the reason for the grouping of the mitochondrial haplotypes in two 485 clusters, west and east of the Rhone river (the later including the Pleistocene expansion of 486 D. semirufus to peninsular Italy and Sicily). This clear geographic pattern is more 487 consistent with introgressive hybridization between closely related taxa sharing the same 488 geographical range, a pattern seen commonly in areas hypothesized to be glacial refugia 489 (e.g. Berthier et al., 2006; Schmidt & Sperling, 2008 or Nicholls et al., 2012). An 490 alternative possibility could be Wolbachia infection, known to alter patterns of mtDNA 491 variability (Jiggins, 2003). Our data do not allow further interpretations of this pattern, 492 which may require a more comprehensive taxon sampling in potential refugial areas (e.g. 493 Massif Central or Black Forest) and the sequencing of additional molecular markers.

Another western lineage which apparently diversified in the Pleistocene is the *D*. *platynotus* group, which was reconstructed to have expanded westwards from the
Balkans, giving rise to the Iberian endemic *D. costipennis*. In Fery & Brancucci (1997)
another species (*D. hakkariensis* Wewalka) known from a single specimen from
southeastern Turkey was tentatively included in the *D. platynotus* group, although

- 499 because of its deviating morphology and geographical distribution this relationship was
- 500 considered doubtful. Unfortunately we could not obtain material of this rare species for
- 501 our analyses, but if shown to genuinely belong to the *D. platynotus* group, *D.*
- 502 hakkariensis would represent a further expansion to the east, most likely during the
- 503 Pliocene, constrained by the stem (Messinian) and crown (lower Pleistocene) ages of the504 group.
- In the eastern clade, the diversification and expansion of the *D. parvicollis* group, to occupy large areas of the Middle East and central Asia, with one species (*D. parvicollis*) expanding westward into the Balkans, most likely took place during the Plio-Pleistocene. The other main lineage within the eastern clade, the *D. latus* group, also expanded during the Pleistocene, but in this case towards the west, first to give rise to the only Italian endemic of the genus (*D. angelinii* Fery & Brancucci) and then to reach
- 511 Iberia and most of Europe north to Scotland and Scandinavia, as testified by Holocene
- 512 remains of *D. latus* in Britain and Sweden (Abellán *et al.*, 2011).
- 513

## 514 CONCLUDING REMARKS

515 Our reconstruction of the evolutionary and biogeographical history of Deronectes shows 516 that its diversification has been shaped by geological and climatic changes around the 517 Mediterranean since the Miocene. These have produced successive rounds of 518 fragmentation, subsequent range expansion leading again to further fragmentation - the 519 overall result of which has been a steady accumulation of species. This pattern of range 520 expansions under favourable conditions followed by fragmentation when conditions 521 change has been described for other groups of lotic Coleoptera (Ribera et al., 2010), and 522 may be a more general pattern contributing substantially to the overall richness of the 523 Mediterranean biodiversity hotspot.

524 Within Deronectes, most of these eastward and westward range expansions 525 involved overland dispersal through the north side of the Mediterranean basin, with a 526 limited influence of Messinian land corridors and the total absence of these beetles in 527 north Africa from Libya to Egypt. This could be expected given their ecological 528 requirements, but what is more surprising is the irrelevance of the Italian peninsula 529 during most of the evolutionary history of the group. Most of the Italian peninsula 530 remained submerged until the Pliocene (Rosenbaum et al., 2002; Meulenkamp & 531 Sissingh, 2003; Popov et al., 2004), and all species currently found in mainland Italy 532 south of the Alps are of Pleistocene origin. This absence of ancient Italian species is

- 533 paralleled in freshwater Coleoptera which have Iberian endemics of Miocene or Pliocene
- 534 origin (*Enicocerus*, *Hydrochus*, Ribera *et al.*, 2010; Hidalgo-Galiana *et al.*, 2011), but not
- 535 by groups with an abundance of Pleistocene species, which have also Italian endemics
- 536 ("Haenydra" and Limnebius; Trizzino et al., 2011; Abellán & Ribera, 2011).
- 537 Our results clearly show that the timing of key diversification events may differ between
- taxa even when sharing the same habitat and geographic distribution, differences that
- 539 have shaped the current distribution of diversity in the Mediterranean hotspot.
- 540

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- 741

742	SUPPORTING INFORMATION
743	Additional Supporting Information may be found in the online version of this article:
744	
745	Appendix S1 Additional materials.
746	(a) Checklist of species of <i>Deronectes</i> , including geographical distribution, species group
747	according to Fery & Brancucci (1997) and clade in which the species was included
748	according to our results. Nomenclature follows Nilsson & Hájek (2015). In grey, species
749	studied in this work.
750	(b) Distribution maps of the species of <i>Deronectes</i> .
751	(c) List of the specimens included in the phylogeny, with specimen voucher, locality,
752	collector and Genbank accession numbers (in bold, new sequences). In grey,
753	Hydroporinae outgroups, with genus-groups according to Ribera et al. (2008).
754	
755	Appendix S2 Additional methods.
756	(a) Primers used for the amplification and sequencing. In brackets, length of the
757	amplified fragment.
758	(b) Standard PCR conditions for the amplification of the studied fragments.
759	
760	Appendix S3 Additional results.
761	(a) Phylogenetic tree obtained with MrBayes with the combined nuclear and
762	mitochondrial sequences and a partition by gene, including all outgroups. Numbers in
763	nodes: Bayesian posterior probabilities.
764	(b) Phylogenetic tree obtained with RAxML with the combined nuclear and
765	mitochondrial sequences and a partition by gene, including all outgroups. Numbers in
766	nodes: Bootstrap support values.
767	(c) Phylogenetic tree obtained with MrBayes using only the mitochondrial sequence data.
768	Numbers in nodes: Bayesian posterior probabilities.
769	(d) Phylogenetic tree obtained with MrBayes using only the nuclear sequence data (H3).
770	Numbers in nodes: Bayesian posterior probabilities.
771	
772	

773	BIOSKETCH
774	David García-Vázquez is a PhD student in the Institute of Evolutionary Biology in
775	Barcelona. This paper is part of his thesis dissertation, focussed on the origin of
776	widespread European species of lotic water beetles. This work is also part of a long-term
777	collaboration between the authors on the evolutionary history of Mediterranean
778	water beetles.
779	
780 781	Editor: Luiz Rocha
782	Author contributions: D.GV. and I.R. conceived the work; D.T.B., L.F.V. and I.R. led
783	the specimen collection; D.GV., R.A. and I.R. obtained the molecular data; D.GV. and
784	I.R. analysed the data and led the writing; all authors contributed to the discussion of
785	results and the writing.
786	
787	
788	

## 789 **TABLES**

- 790 **Table 1** Dispersal cost schemes used in Lagrange, and likelihood of the different
- 791 Lagrange models. (a) Dispersal probabilities across sea or land barriers in the six
- 792 combinations used (in bold, combination with the best likelihood score). "land barrier"
- refers to one of the pre-defined areas (see Fig. 3). (b) Likelihood of the six dispersal
- schemes in (a) for the three tested palaeogeographical scenarios. The matrix with the best
- 795 likelihood score for each scenario is shown in bold, with a star for the best overall
- scheme. (c) Matrix of dispersal probabilities between our pre-defined geographical areas
- for each palaeogeographical scenario according to the costs of scheme #3 in (a) (see Fig.

798 3 for the maps used for the reconstruction).

799

## 800 (a)

barrier	#1	#2	#3	#4	#5	#6
contiguous land areas	1	1	1	1	1	1
land barrier	0.2	0.2	0.4	0.2	0.1	0.2
sea barrier < 100 km	0.4	0.4	0.4	0.2	0.2	0.4
two land barriers	0	0.1	0	0	0	0.2
sea barrier >100 km	0	0.1	0	0	0	0
>2 land or 2 land+sea	0	0.1	0	0	0	0

801

802 (b)

palaeogeographical scenario	#1	#2	#3	#4	#5	#6
Pleistocene (present)	103.2	104.68	101.7*	103.2	103.9	103.7
Pliocene	107.5	108.4	105.1	107.5	108.4	107.8
Miocene (late Tortonian)	107.6	108.5	105.5	107.5	108.5	107.9

<sup>804 (</sup>c)

Pleistocene		Α	В	С	D	Е	F	G
SE Iberian Peninsula and Mallorca	Α	-						
Centre and N Iberian Peninsula	В	1	-					
SE France, Italy and Sicily	С	0.6	1	-				
Corsica and Sardinia	D	0	0	0.6	-			
Balkans	Е	0	0.6	1	0	-		
Turkey and Middle East	F	0	0	0.2	0	0.6	-	
Northern and central Europe	G	0.6	1	1	0.2	1	0	-
Maghreb	Η	0.6	0.2	0	0	0	0	0
Pliocene		Α	В	С	D	Е	F	G
SE Iberian Peninsula and Mallorca	Α	-						
Centre and N Iberian Peninsula	В	1	-					
SE France, Italy and Sicily	С	0.6	1	-				
Corsica and Sardinia	D	0.6	0.6	1	-			
Balkans	Е	0	0.6	1	0.6	-		
Turkey and Middle East	F	0	0	0.6	0	1	-	

Northern and central Europe	G	0.6	1	1	0.6	1	0.6	
Maghreb	Η	1	0.6	1	1	0.6	0	0
Miocene (Late Tortonian)		А	В	С	D	Е	F	G
SE Iberian Peninsula and Mallorca	А	-						
Centre and N Iberian Peninsula	В	1	-					
SE France, Italy and Sicily	С	0.6	1	-				
Corsica and Sardinia	D	0	0.6	1	-			
Balkans	Е	0	0.6	1	0.6	-		
Turkey and Middle East	F	0	0	0.6	0	1	-	
Northern and central Europe	G	0.6	1	1	0.6	1	0.6	-
Maghreb	Η	0.6	0.2	0.6	0.6	0.2	0	0.2

**Table 2** Ancestral area reconstruction in the Lagrange analyses of the 1000 post-burn-in

trees. For each node present in the consensus tree (see Fig. 3) we give the number of trees

808 in which the node appears, the most frequent area or combined areas and the frequency of

the individual areas. In bold and with stars, areas with >90% of frequency; in bold, areas

810 between 70-90 % frequency. Area codes: A, south and east of the Iberian Peninsula

811 including Mallorca; B, centre and north of the Iberian Peninsula; C, Italy (including

812 Sicily) and southeastern France; D, Corsica and Sardinia; E, Balkan peninsula; F, Turkey

and Middle East; G, northern and central Europe and H, Maghreb.

814

NODE	n° trees	areas	А	В	С	D	Е	F	G	Н
1	1000	BDFH	5	93*	-	100*	9	100*	-	93*
2	1000	ABDH	56	93*	1	100*	18	41	-	89
3	488	В	-	100*	-	-	11	6	-	-
4	1000	В	48	100*	-	-	-	-	-	-
5	530	В	-	100*	-	-	-	-	-	-
6	1000	В	-	100*	-	-	-	-	-	-
7	272	AB	85	75	-	-	-	-	-	-
8	634	В	57	85	-	-	-	-	-	-
9	805	AB	99*	100*	-	-	-	-	-	-
10	1000	В	55	64	-	-	-	-	-	-
11	636	BEF	-	71	7	-	72	43	-	-
12	1000	BCG	-	100*	100*	-	-	-	100*	-
13	1000	CG	-	-	100*	-	-	-	100*	-
14	999	CG	-	-	100*	-	-	-	81	-
15	970	В	-	100*	-	-	-	-	-	-
16	418	EF	-	34	-	-	89	77	-	-
17	849	E	-	11	-	-	100*	28	-	-
18	1000	BEG	-	100*	-	-	100*	-	72	-
19	1000	EG	-	-	-	-	100*	-	100*	-
20	1000	ABDH	62	53	4	100*	-	-	-	95*
21	1000	ABDH	91*	73	14	87	-	-	-	100*
22	1000	ABCD	98*	80	54	81	1	-	-	21
23	1000	ABCD	100*	100*	100*	100*	-	-	-	-
24	1000	А	100*	-	-	-	-	-	-	1
25	1000	F	-	-	-	-	-	100*	-	-
26	1000	CEFG	-	30	100*	-	100*	100*	70	-
27	998	CEFG	-	34	100*	-	100*	100*	66	-
28	1000	BCEG	-	89	100*	-	100*	-	98*	-
29	907	BCEG	-	100*	100*	-	100*	-	100*	-
30	1000	F	-	-	-	-	-	100*	-	-
31	1000	F	-	-	-	-	-	100*	-	-
32	1000	F	-	-	-	-	-	100*	-	-
33	1000	F	-	-	-	-	-	100*	-	-
34	1000	F	-	-	-	-	-	100*	-	-

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817

819	FIGURE LEGENDS
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820	Figure 1 Distribution of the main lineages of <i>Deronectes</i> according to our phylogenetic
821	results (see Fig. 2).
822	
823	Figure 2 Phylogeny of <i>Deronectes</i> , as obtained with MrBayes with the combined nuclear
824	and mitochondrial sequence and a partition by gene. Numbers on nodes, Bayesian
825	posterior probabilities/Bootstrap support values in RAxML. Habitus photograph, D.
826	fosteri Aguilera & Ribera (from Millán et al., 2015).
827	
828	Figure 3 Ultrametric time calibrated tree obtained with Beast. Coloured branches show
829	ancestral distributions as estimated from the analysis of 1000 post-burn-in trees. Above
830	nodes (in brackets) the most frequent area or combined areas reconstructed as the
831	ancestral area of the node (see Table 2). Numbers inside nodes refer to Table 2.
832	
833	Figure 4 Lineage Through Time plot (LTT) of the western clade obtained from the
834	ultrametric tree in Fig. 3.
835	

837 Fig. 1





840 Fig. 2





850 Fig. 4

