

Global gap-analysis of amphipod barcode library

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In the age of global climate change and biodiversity loss there is an urgent need to provide effective and robust tools for diversity monitoring. One of the promising techniques for species identification is the use of DNA barcoding that in Metazoa utilizes the so called ‘gold-standard’ gene of cytochrome *c* oxidase (COI). However, the success of this method relies on the existence of trustworthy barcode libraries of the species. The Barcode of Life Data System (BOLD) aims to provide barcodes for all existing organisms, and is complemented by the Barcode Index Number (BIN) system serving as a tool for potential species recognition. Here we provide an analysis of all public COI sequences available in BOLD of the diverse and ubiquitous crustacean order Amphipoda, to identify the barcode library gaps and provide recommendations for future barcoding studies. Our gap analysis of 25702 records has shown that although 3835 BINs (indicating putative species) were recognised by BOLD, only 10 % of known amphipod species are represented by barcodes. We have identified almost equal contribution of both records (sequences) and BINs associated with freshwater and with marine realms. Three quarters of records have a complete species-level identification provided, while BINs have just 50%. Large disproportions between identification levels of BINs coming from freshwaters and the marine environment were observed, with three quarters of the former possessing a species name, and less than 40% for the latter. Moreover, the majority of BINs are represented by a very low number of sequences rendering them unreliable according to the quality control system. The geographical coverage is poor with vast areas of Africa, South America and the open ocean acting as “white gaps”. Several, of the most species rich and highly abundant families of Amphipoda (e.g. Phoxocephalidae, Ampeliscidae, Caprellidae), have very poor representation in the BOLD barcode library. As a result of our study we recommend stronger effort in identification of already recognised BINs, prioritising the studies of families that are known to be important and abundant

components of particular communities, and targeted sampling programs for taxa coming from geographical regions with the least knowledge.

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14 **Abstract**

15
16 In the age of global climate change and biodiversity loss there is an urgent need to provide
17 effective and robust tools for diversity monitoring. One of the promising techniques for species
18 identification is the use of DNA barcoding that in Metazoa utilizes the so called ‘gold-standard’
19 gene of cytochrome *c* oxidase (COI). However, the success of this method relies on the existence
20 of trustworthy barcode libraries of the species. The Barcode of Life Data System (BOLD) aims
21 to provide barcodes for all existing organisms, and is complemented by the Barcode Index
22 Number (BIN) system serving as a tool for potential species recognition. Here we provide an
23 analysis of all public COI sequences available in BOLD of the diverse and ubiquitous crustacean
24 order Amphipoda, to identify the barcode library gaps and provide recommendations for future
25 barcoding studies. Our gap analysis of 25702 records has shown that although 3835 BINs

26 (indicating putative species) were recognised by BOLD, only 10 % of known amphipod species
27 are represented by barcodes. We have identified almost equal contribution of both records
28 (sequences) and BINs associated with freshwater and with marine realms. Three quarters of
29 records have a complete species-level identification provided, while BINs have just 50%. Large
30 disproportions between identification levels of BINs coming from freshwaters and the marine
31 environment were observed, with three quarters of the former possessing a species name, and
32 less than 40% for the latter. Moreover, the majority of BINs are represented by a very low
33 number of sequences rendering them unreliable according to the quality control system. The
34 geographical coverage is poor with vast areas of Africa, South America and the open ocean
35 acting as “white gaps”. Several, of the most species rich and highly abundant families of
36 Amphipoda (e.g. Phoxocephalidae, Ampeliscidae, Caprellidae), have very poor representation in
37 the BOLD barcode library. As a result of our study we recommend stronger effort in
38 identification of already recognised BINs, prioritising the studies of families that are known to be
39 important and abundant components of particular communities, and targeted sampling programs
40 for taxa coming from geographical regions with the least knowledge.

41

42 **Keywords**

43 DNA barcoding, Crustacea, marine, freshwater, semi-terrestrial, taxonomic identification

44

45 **Introduction**

46 Nature in the age of Anthropocene is facing numerous global changes and challenges. One of the
47 drastic results of human associated activities is the acceleration of species extinctions, with one

48 million species estimated to be presently critically endangered (IPBES 2019). What is more,
49 although the rate of species discovery grows, large numbers of species remain undescribed and it
50 is believed many will not be recognized before they go extinct (Mora et al., 2011). This raises the
51 challenge of efficient environmental monitoring, which is crucial for biodiversity recognition and
52 preservation. Monitoring based on the taxonomic identification of organisms in samples is time-
53 consuming and requires knowledge of the studied group. In the time of the taxonomic
54 impediment (Ebach, Valdecasas & Wheeler, 2011), species identification methods offering an
55 alternative to morphology-based methods are of great interest. Utilization of DNA-barcoding
56 (identifying sequences of individual specimens), metabarcoding (high-throughput identification
57 of bulk samples) and the use of environmental DNA (e-DNA, identifying DNA of taxa directly
58 from water or soil sample, without collection of specimens) have been presented as promising
59 methods in monitoring and ecological studies (e.g., Hajibabaei et al., 2012; Cristescu, 2014;
60 Aylagas et al., 2018; Leese et al., 2018; Bush et al., 2019; Feio et al., 2020). The use of
61 metabarcoding in assessing the status of ecosystems has already received the new term
62 “Biomonitoring 2.0” (Bush et al., 2019). Such approaches require the existence of well-
63 established barcode fragment libraries, which allow accurate recognition of organisms in the
64 environment (Cristescu, 2014; Cowart et al., 2015; Oliveira et al., 2016; Múrria et al., 2020).
65 Recent studies indicate that although the use of barcoding in biomonitoring has great advantages
66 over morphological identification, the current gaps in barcode libraries may hinder their use
67 (Weigand et al., 2019; Feio et al., 2020; Hestetun et al., 2020; Leite et al., 2020; Múrria et al.,
68 2020).

69 There are two main repositories where DNA sequences are deposited: NCBI GenBank
70 (www.ncbi.nlm.nih.gov/genbank/, Sayers et al., 2020) and Barcode of Life Data System (BOLD,

71 www.boldsystems.org, Ratnasingham & Hebert, 2007). In contrast to GenBank, which
72 assembles nucleotide data of all genes, the primary aim of BOLD is to store data used for species
73 barcoding, which in the case of Metazoa is the cytochrome *c* oxidase (COI) gene. The
74 development of the BOLD database included the Barcode Index Number (BIN) system
75 implementation (Ratnasingham & Hebert, 2013) that intends to help in biodiversity assessments
76 by providing species-level taxonomic registry. Based on a molecular species delimitation
77 method, each Molecular Operational Taxonomic Unit (MOTU) recognized by BOLD receives a
78 unique alphanumeric code (BIN). Ideally, each BIN is associated with an accurate taxonomic
79 (preferably species) identification and links to the voucher stored in a recognised institution.
80 However, in practice this is not working well, and at the time of system implementation as many
81 as 46% of BINs lacked species names (Ratnasingham & Hebert, 2013). This issue has arisen for
82 a variety of reasons, which we investigate in this study using a particular faunal group, the
83 Amphipoda, as a model.

84 The Order Amphipoda are peracarid crustaceans belonging to the class Malacostraca. They are
85 very diverse components of aquatic environments. According to the World Amphipoda Database
86 (WAD, Horton et al., 2020, accessed on 17-07-2020) there are 10235 accepted amphipod
87 species, the majority of which (78%) inhabit the marine realm, around 20% are freshwater
88 species and just 2% are terrestrial taxa (Horton et al., 2020; Väinölä et al., 2008). The discovery
89 rate of new species has grown steadily since the first amphipod species description and has
90 particularly accelerated in the last six decades (Horton et al., 2020) with mean number of over
91 100 taxa annually described since the 1960s (Coleman, 2015). If the trend from the last sixty
92 years persists, we may expect to have ca. 8000 new species described by 2100. More
93 conservative estimates predict that 6100 new species will be described by that date (Arfianti,

94 Wilson & Costello, 2018). The use of molecular methods in the studies of Amphipoda has
95 revealed very high species diversity (e.g. Knox et al., 2012; Verheye, Backeljau & d'Udekem
96 d'Acoz, 2016; Jazdzewska & Mamos, 2019) and revealed the existence of cryptic species
97 complexes within widely distributed taxa (Witt, Therloff & Hebert, 2006; Mamos et al., 2014;
98 Wysocka et al., 2014; Havermans, 2016). Amphipoda are not only a species-rich group, but they
99 also often dominate the crustacean assemblages in which they occur (e.g., Corkum, 1989;
100 Humphries, Davies & Mulcahy, 1996; Vinogradov, Volkov & Semenova, 1996; Jazdzewski et
101 al., 2001; Väinölä et al., 2008; Frutos, Brandt & Sorbe, 2017; Brix et al., 2018; Havermans &
102 Smetacek, 2018). They can be found in both the benthos and the pelagic realm, presenting a
103 variety of states of mobility (from epibenthic clingers to fully mobile swimmers) and, as a result,
104 possess a wide variety of feeding habits including herbivory, detritivory, necrophagy, omnivory,
105 predation and ectoparasitism (Barnard & Karaman, 1991; Vinogradov, Volkov & Semenova,
106 1996; Dauby, Scailteur & De Broyer, 2001; Väinölä et al., 2008). Being diverse and abundant
107 they are important prey items for other invertebrates and vertebrates, including fish, birds and
108 mammals (e.g. Dalpadado et al., 2001; Dauby, Nyssen & De Broyer, 2003). Certain species of
109 Amphipoda are used in laboratory ecotoxicological studies (Hyne & Everett, 1998; Bundschuh et
110 al., 2011; Bundschuh et al., 2013, Major et al., 2013). Some amphipod species are well-adapted
111 to anthropogenic environments such as artificial structures used in coastal protection or are part
112 of fouling communities, and have shown a high invasion potential worldwide (e.g., Bij de Vaate
113 et al., 2002; Kelly et al., 2006; Cabezas et al., 2014; Rewicz et al., 2015; Beermann et al., 2020;
114 Sedano et al., 2020).

115 The combined factors of high diversity and the important role played by amphipods in the
116 aquatic ecosystem highlights the need for accurate species identifications which are required for

117 biological monitoring programs. The use of DNA-barcoding may speed up the identification
118 process, but it will only succeed if the barcode library is well-established and robust. Recent gap-
119 analyses of the barcode libraries in aquatic European environments showed very large
120 differences in the coverage between different taxonomic groups and geographic regions
121 (Weigand et al., 2019; Feio et al., 2020; Hestetun et al., 2020; Leite et al., 2020). These studies
122 used species lists restricted to particular geographic regions or chosen taxonomic groups. Basic
123 summaries concerning the extent of amphipod data in BOLD identified problems with lack of
124 taxonomic identification or detailed geographic information as well as contamination with
125 human or bacterial DNA and provided recommendations to improve the data (Radulovici &
126 Coleman, 2017; Coleman & Radulovici, 2020). However, to date there are no detailed analyses
127 that have been conducted on a single taxon on a global scale.

128 In this study we have conducted a gap-analysis of the barcode library of a single crustacean
129 order, the Amphipoda, on a global basis. In producing an up-to-date picture of the current state
130 of knowledge, we will provide researchers with a detailed understanding of the both the strengths
131 and the potential limitations of the use of DNA barcodes for identifications. We also propose
132 recommendations for future initiatives that involve molecular data and produce new barcodes to
133 fill the gaps in our knowledge of this taxon.

134

135 **Material and methods**

136 Data for the present study were retrieved from BOLD by searching the “Public Data Portal”
137 using the keyword “Amphipoda”. A combined dataset of all records was downloaded as an .xml
138 file on June 24th 2020.

139 All records of the barcoding fragment of the cytochrome c oxidase I (COI-5P in BOLD) were
140 extracted (29016 records). This extracted dataset was used for all further analyses conducted by
141 using various filtering options in an Excel spreadsheet. 2579 records, represented by sequences
142 shorter than 500 bp or having more than 1% ambiguous nucleotides for which BINs were not
143 ascribed, were removed from dataset. Continued analysis of the dataset revealed some duplicate
144 records (1468 records, 734 cases, Supplemental file 1). These derived from data harvested by
145 BOLD from GenBank and seemed to be associated with an update of the records GenBank. In
146 the dataset, these records had an identical sample ID that referred to a GenBank Accession
147 Number but with an additional ‘.1’ appended (e.g. KP713892 and KP713892.1) with an identical
148 identification provided. The differences were often linked with more detailed geographical
149 information in the case of one record from the pair. Only the more detailed entry was retained for
150 continued analysis. One sequence of *Niphargus novomestanus* S. Karaman, 1952 (KR858496,
151 BOLD:ADD1128) was removed from the dataset because it was deleted from GenBank by its
152 submitter (“This record was removed at the submitter's request because the source organism
153 cannot be confirmed.” GenBank website). The resulting dataset contained 25702 records (Fig. 1,
154 Supplemental file 2).

155 Each record in the dataset was then further refined by sorting into categories according to the
156 level of taxonomic identification. The following categories were used: order, family, subfamily,
157 genus and species. Where records were provided with a temporary species identification, i.e.
158 they are recognised as separate morphospecies but are not determined to correspond to a known
159 taxon - they were treated as a separate category. In the whole dataset ca. 2.5% of records (596
160 individuals, 145 BINs) had uncertain identification with “cf.” or “aff”. Because the majority of
161 them (417 records, 101 BINs) were associated with five species of one genus (*Gammarus*) for

162 simplification all such records were treated as final species identifications. However, it is
163 understood that the use of open nomenclature, when applied to identifications, provides an
164 indication of the level of uncertainty, and may be intended to indicate the presence of new
165 species or species complexes.

166 The data in BOLD come from wide variety of projects, some of which involve detailed
167 taxonomic study by specialists, others are focused on monitoring or other topics in which
168 taxonomic specialists are not involved. For the purposes of our analyses it was assumed that the
169 identification accuracy was equal throughout the whole dataset, regardless of its origin. In
170 several cases identification of the specimens within a single BIN varied strongly, with some
171 records remaining at order level while others were determined to the species level. BINs aim to
172 represent a putative species, so in the above example, the most detailed taxonomic information
173 was applied to all records within the single BIN. Sometimes multiple (most often two) species or
174 genus names were associated with a single BIN (87 cases). Each of these cases was checked
175 individually. Sometimes it was an obvious misidentification of a single individual within a large
176 group - if this was noted the misidentified record was added as an additional element to the
177 records identified to the lowest congruent level (e.g. if the genus name matched the BIN genus,
178 the misidentified taxon was added as an additional record identified to the genus level, if the
179 lowest congruent level was family it was added to the family records); and the taxon
180 identification of the majority of records was applied as correct. When it was impossible to judge
181 which name was correct, the name of the identifier was checked and identifications carried out
182 by taxonomists specializing in Amphipoda was prioritised over a that provided by a non-
183 specialist study. Where this process did not give a satisfactory conclusion, the BIN was allocated
184 an identification at a rank that was congruent for the different records. The list of taxa with

185 incongruent identifications together with an explanation of the final decision is presented in
186 Supplemental file 3.

187 Based on the taxonomic identification of the records the associated BINs were divided into the
188 following environmental categories:

- 189 a) marine
- 190 b) freshwater
- 191 c) terrestrial.

192 Taxa that inhabit both marine realm and brackish environments were allocated to the marine
193 category. Taxa from freshwater also occurring in brackish waters were allocated to the
194 freshwater category. All representatives of the family Talitridae were treated as terrestrial taxa.
195 Where taxonomic information was not detailed enough to provide environmental information
196 about the particular BIN, the geographic data (coordinates and/or locality description) of the
197 associated records were used to ascribe a particular BIN to one of the above categories. In some
198 cases, this necessitated checking the original publication. A small number of unallocated BINs
199 (18) and associated records (44) were removed from the dataset (Supplemental file 4).

200 In order to verify the correct environmental allocation of BINs, all BINs with records possessing
201 coordinates were plotted on a map using the software QGIS2.16.1 (QGIS Development Team,
202 2018). Cases where incongruence between the ascribed environment and the geographic position
203 appeared were checked individually. For those records without detailed geographic information
204 the country of origin was taken from either BOLD or the associated publication.

205 In order to verify the barcode coverage within the studied group a list of BINs associated with a
206 species name was compared with the list of accepted amphipod species names available in the
207 World Amphipoda Database (WAD, Horton et al., 2020, accessed on 17-07-2020). A barcode

208 quality assessment of the species represented in BOLD, based on the grading system proposed
209 by Oliveira et al. (2016) and slightly modified by Fontes et al. (2020) was applied. This system
210 consists of five grades: A – consolidated concordance (>10 sequences of a single morphospecies
211 grouped in a single BIN), B – basal concordance (same as grade A but between three and 10
212 sequences available in the library), C – multiple BINs (one morphospecies assigned to more than
213 one BIN), D – insufficient data (single species is assigned to single BIN but it is represented by
214 less than three sequences in the barcode library), E – discordant species assignment (more than
215 one species assigned to a single BIN). Fontes et al. (2020) provide an R-based application
216 (Barcode, Audit & Grad System – BAGS), and uses only those records possessing either
217 coordinates or the indication of the country of origin. Since this information is very often missing
218 in the data mined from GenBank, and our aim was to focus on all available barcode records, the
219 assessment was carried out manually. Additionally, as a result of initial treatment of the dataset,
220 misidentified species records or BINs with unclear species identification, were already removed,
221 so category E (discordant species assignment, Oliveira et al., 2016; Fontes et al., 2020) was not
222 recorded. For the purpose of the present study *Lysianassoidea incertae sedis* was treated as an
223 additional family. The amphipod families were divided into four categories depending on the
224 number of species in each: low species rich families (up to 10 species), moderately species rich
225 families (from 11 to 30 species), species rich families (31-100 species), very species rich families
226 (more than 100 species). This division allowed verification of pattern between the species
227 richness of the family and its representation in BOLD.

228 **Results**

229 Of the 25702 amphipod COI records, 46.5% (11958 records) were freshwater, 43.5% (11169
230 records) were from the marine realm, and 9.8% (2531 records) were terrestrial taxa. Of the 3835

231 recognized BINs in total, 45% (1726 BINs) belonged to freshwater taxa, 50% (1920 BINs) were
232 marine, and 4.5% (171 BINs) were from terrestrial taxa. 44 records (0.2%) and their associated
233 18 BINs (0.5%) could not be ascribed to the above environmental categories (Fig. 2A, B).

234 More than half (57.5%) of the records available in BOLD possessed coordinates, and 20% had
235 information about the country of origin. Geographic information about the remaining 22.5% was
236 provided only in the original publication and this information was added manually. Geographic
237 information is more comprehensive for marine taxa, where 71% of records possessed coordinates
238 (compared to 47% for freshwater, and 50% for terrestrial taxa). Molecular studies of freshwater
239 Amphipoda are focused mainly in the Northern hemisphere (particularly European countries,
240 Russia and United States) while in the Southern hemisphere, Australia, New Zealand and
241 Argentina are well studied (Fig. 3A). There is a complete lack of records (amphipod sequences)
242 from Brazil, equatorial America and vast areas of Africa. Similar patterns of data coverage were
243 seen for marine amphipods, which have greater numbers of records along European, North
244 American and East Asian coasts. In the Southern hemisphere, Australia, New Zealand and
245 Antarctica had larger numbers of barcode records (Fig. 3B). However, vast areas of the deep sea
246 and the Arctic Ocean remain undersampled. Terrestrial Amphipoda in Europe, North America,
247 China, Australia and Chile were the best represented (Fig. 3C), but sampling gaps were seen in
248 the continents of South America and Africa.

249 The majority of records with environmental information (69.8%, 17922 recs.) had a complete
250 species-level identification. Of the remaining 30.2% of records, 5.6% (1433 recs.) had received
251 temporary names (open nomenclature), 11.3% (2902 recs.) remained identified at the genus
252 level, 0.2% (40 recs.) at subfamily, 5.0% (1285 recs.) at family, and 8.1% (2076 recs.) at the
253 order level. Levels of identification varied according to the environment, with marine taxa

254 having greater proportions of taxa identified only to higher taxonomic ranks (Fig. 4A). The
255 majority of BINs with geographic affiliation (3817) were associated with species names (55.7%,
256 2126 BINs). These were followed by BINs identified to the order level (13.3%, 506 BINs),
257 generic or family level (10.7%, 407 BINs each) and those with a temporary name (9.4%, 359).
258 BINs with only a subfamily name constituted just 0.3% (12). Greater variations between
259 environments were seen for the BINs, with 74% (1284) of freshwater BINs having a species
260 level identification, compared to only 39% (751) of marine BINs (Fig. 4B). More than 20% (444,
261 23%) of the BINs for marine taxa remained identified at the order level.

262 Regardless of the environmental origin, the majority of BINs were represented by a single
263 sequence (Fig. 5). BINs represented by five or fewer sequences constituted around two thirds
264 (67%, 114 terrestrial BINs to three quarters, 78%, 1488 marine BINs) of BINs recorded in a
265 particular environment. Freshwater taxa had 41 BINs (2.4%) represented by more than 50
266 sequences, compared to 28 (1.5%) for marine taxa, and eight (4.7%) for terrestrial taxa. When
267 only those BINs with complete species-level identifications are considered, the proportion of
268 sequences representing a particular MOTU does not change, with freshwater taxa having 78% of
269 BINs (1016) represented by five or fewer sequences. Almost three quarters of marine BINs
270 (71%, 525 BINs) had five or fewer sequences in BOLD, while this proportion was 61% (56
271 BINs) for terrestrial taxa. Freshwater taxa had 35 BINs (3%) represented by more than 50
272 sequences, compared to 27 (4%) for marine taxa, and 6 (7%) for terrestrial taxa. The best
273 represented BIN in BOLD (801 sequences) belonged to the terrestrial species *Orchestoidea*
274 *tuberculata* Nicolet, 1849 (BOLD:ACQ3380), followed by the marine species *Gammarus*
275 *oceanicus* Segestråle, 1947 (BOLD:AAA1262, 553 sequences), and the freshwater species
276 *Diporeia hoyi* (S.I. Smith, 1874) (BOLD:AAA1473, 512 sequences). A further 26 BINs were

277 represented by more than 100 sequences, including 17 freshwater, seven marine and two
278 terrestrial BINs (Supplemental file 5).

279 Out of the 3817 studied BINs, just over half (55.7%, 2126) were associated with a species-level
280 identification, representing 1001 species. Freshwater BINs with species identification reached
281 1284, associated with 453 species, while 751 marine BINs were determined to 496 species. Of
282 the 91 terrestrial BINs, 52 species were identified. Generally, a single morphological species was
283 associated with each BIN (68%, 680 cases, 288 in freshwater, 359 marine, 33 terrestrial). 17% of
284 the identified species were associated with two different BINs (72 freshwater, 82 marine and 14
285 terrestrial) (Fig. 6). There were however 19 cases when one single morphological species was
286 represented by more than 10 BINs (17 freshwater, one marine and one terrestrial) (Supplemental
287 file 6). The greatest number of BINs was recorded for the freshwater species *Gammarus*
288 *balcanicus* Schäferna, 1923 represented by 143 BINs (45 BINs were identified as “cf.” or “aff.”)
289 followed by another freshwater taxon *Hyalrella azteca* (Saussure, 1858) (62 BINs) and
290 *Gammarus fossarum* Koch, 1836 (51 BINs; 19 BINs identified as “cf.” or “aff.”). Among
291 terrestrial taxa the highest molecular variation (12 BINs) was recorded for *Morinoia japonica*
292 (Tattersall, 1922) (present in BOLD under former generic name *Platorchestia*), while *Apohyale*
293 *stebbingi* Chevreux, 1888 (with 11 BINs recognized) was the most diverse among marine
294 species.

295 Of the 239 accepted families of Amphipoda (238 families and *Lysianassoidea incertae sedis*),
296 105 (44%) were represented by at least one species in BOLD (Table 1). The largest number of
297 families had up to 20% of species barcoded, while only ten families had more than half of the
298 known species barcoded (Supplemental file 7). Thirteen families lacking barcoded species had at

299 least one barcoded taxon identified at the genus level, a further five families had a taxon
300 identified at the family level.

301 Just under ten percent (999 spp., 9.7%) of the 10330 accepted species of Amphipoda (Horton et
302 al., 2020) had barcodes. Of the nominal species possessing barcodes almost 500 (496 spp.) are
303 marine, 451 spp. are freshwater and 52 spp. are terrestrial taxa. The data coverage of the majority
304 of species, no matter their environmental origin, is not sufficient for the barcodes to be trusted
305 according to the quality control system (Table 2) (Oliveira et al., 2016; Fontes et al., 2020).
306 Additionally, a large group of taxa is represented by multiple BINs; only 10% of species
307 represent consolidated concordance of available barcodes.

308 The breakdown of amphipod families according to the assigned categories of richness and their
309 respective representation in BOLD can be seen in Table 3. Almost every one of the very species
310 rich families had at least one species barcoded (31 families out of 32), and 22 of 30 species rich
311 families are represented in BOLD. For both moderately low and low species rich families 26
312 possessed at least one representative in BOLD constituting respectively 48% and 21% of all
313 families each (Supplemental file 7). The mean coverage of barcodes for species in each of the
314 above groups was around 10% with the highest observed for low species rich families (12%) and
315 the lowest (8%) recorded for families grouping from 30 to 100 species. However, if the families
316 without any molecular information were removed from the study these numbers considerably
317 change. The low species rich families (1-10 spp.) had a barcode coverage at the level of 49%,
318 moderately species rich families (11-30 spp.) reached 21% of coverage, while the rich and very
319 rich amphipod families (more than 30 spp.) had only 9-10% of species studied.

320 A third of families (34) have at least one species characterized by consolidated concordance of
321 available barcodes (category A of the quality grading system). Another third of families (38) do

322 not have any species in categories A or B, indicating that the species already studied represent a
323 potential cryptic diversity or the available data are insufficient (Table 4).

324 Within the very species rich families, the best representation in BOLD was recorded for
325 Niphargidae (36.5% of known species represented with a barcode), Gammaridae (31%) and
326 Crangonyctidae (16%). Only the family Stegocephalidae did not have any representative with
327 species level identification (although barcodes belonging to this family but identified at genus
328 level were present). The least studied families within this group (but having at least one species
329 barcode) were: Phoxocephalidae (1% of the species with a barcode), Dexaminidae,
330 Liljeborgiidae and Maeridae (ca. 2% of the species with a barcode). Among species rich families
331 41% of the species from Pseudoniphargidae had barcodes, while the Epimeriidae and
332 Pontogammaridae had 20% and 19%, respectively. The best represented moderately species rich
333 families were Metacrangonyctidae, Oxycephalidae and Hyperiididae with 55%, 50% and 48% of
334 the associated species represented with a barcode respectively. Within low species rich families
335 four (Baikalogammaridae, Crymostygidae, Cyllopodidae and Tryphanidae) had all known
336 species represented with barcodes, but other than Cyllopodidae (two species) the families are
337 monotypic (Supplemental file 7).

338 **Discussion**

339 **Extent of barcode library of Amphipoda**

340 One of the aims of establishing the BOLD database was to store and publish barcodes, based on
341 records uploaded by its users and supplemented by the data harvested from GenBank
342 (Ratnasingham & Hebert, 2007). Together with the BIN system, that groups similar sequences in
343 clusters representing putative species (Ratnasingham & Hebert, 2013), the BOLD database aids
344 in recognising and quantifying biodiversity. The extent of data in BOLD expresses the activity of

345 researchers studying particular groups using molecular methods. The number of available
346 sequences of Amphipoda in BOLD is comparatively large. At the time of download (end of June
347 2020) Amphipoda in BOLD were represented by almost 26 000 records (3835 BINs), and by the
348 end of August there were more than 34 000 public sequences (3914 BINs) (BOLD accessed on
349 20-08-2020), indicating the great intensity of molecular studies involving this crustacean group,
350 and that the data in BOLD are actively growing. Among other crustacean groups only Decapoda
351 is represented by a higher number of records (64 281 records). Copepoda are represented by 18
352 511, Thecostraca by 15 554, Isopoda by 13 858 and Branchiopoda by 12 326 sequences. The
353 large number of identified BINs within the Amphipoda also places this group second only to
354 Decapoda (with 6056 BINs). Isopods and copepods are represented by 1853 and 1804 BINs,
355 respectively, while 969 BINs were identified within Branchiopoda. Within Thecostraca only 545
356 BINs were identified (boldsystems.org, accessed on 20-08-2020).

357 Ratnasingham & Hebert (2013) when presenting the BIN system indicated that overall 12% of
358 the sequences available in BOLD at that time were lacking a family name, 19% a genus name
359 and 40% a species name. A comparison of these numbers with the present data on Amphipoda
360 looks optimistic, where only 8% of sequences are without family indication, 13% without genus
361 and 29% species identification. However, the global analysis of Ratnasingham & Hebert (2013)
362 also identified 10% of BINs lacking family names, almost 24% lacking generic names and 46%
363 lacking species names. These numbers are almost identical for amphipod BINs known presently
364 (13%, 23%, 43% of BINs lacking family, genus and species information, respectively). Among
365 all known species of Amphipoda, almost 80% of species are marine, some 20% live in
366 freshwaters, while 2% may be considered as terrestrial (Horton et al., 2020; Väinölä et al., 2008).
367 The above proportions are expressed neither in the number of records nor the number of

368 recognized BINs that are more or less evenly distributed between freshwater and marine taxa.
369 This demonstrates that in terms of amphipod crustaceans freshwater taxa are much better studied
370 than the marine taxa. These disproportions are even more striking when the level of
371 identification of sequences and BINs is considered. Although the majority of data present in
372 BOLD possess species-level identifications, marine amphipods are? less thoroughly identified.
373 This is especially clear for marine BINs, of which only 39% had species-level identifications,
374 while as much as one fifth are identified only as “Amphipoda”. The fact that freshwater
375 amphipods are better studied is not surprising considering the easier access to this environment.
376 In the case of marine fauna, obtaining samples suitable for molecular analysis can be
377 challenging, especially when extreme habitats (polar regions, deep-sea, hydrothermal vents etc.)
378 are considered (e.g., Riehl et al., 2014; Jazdzewska & Mamos, 2019;). Additionally, rarity is a
379 common feature of numerous marine species (particularly in the deep-sea environment, see
380 Kaiser et al., [2007]), where many taxa are known only from their original descriptions and type
381 localities (Jazdzewska & Mamos, 2019). The question of how many of the BINs not associated
382 with a species identification actually belong to already known species is also of concern. In these
383 cases, it is highly advisable to put every effort to identify the already available material – this
384 will relatively efficiently improve data usability. Taxa that are associated with a BIN, yet are
385 known to be new to science are another cause for concern. This is particularly evident for marine
386 taxa collected during recent deep-sea exploration programs (e.g., Brandt et al., 2007;
387 Jazdzewska, 2015; Brandt et al., 2019; Brix et al., 2020). It is imperative that full scientific
388 descriptions of new species are produced to reduce the current proliferation of ‘dark taxa’ (Page,
389 2016).

390 The geographic distribution of available amphipod sequence records shows clear sampling gaps.
391 In particular the African continent, the northern part of South America and the Coral Triangle in
392 Asia are complete “white spots” when freshwater and terrestrial taxa are considered. For marine
393 species, the coasts of Africa and South America, the Coral Triangle, and large parts of the deep
394 sea of all oceans, lack coverage. Considering the known high species diversity of these regions it
395 will be necessary to establish targeting sampling programs before we can consider that we have
396 adequate global coverage of the molecular diversity of the Amphipoda.

397 Our study shows that globally the barcoding coverage of amphipod species is only about 10%. In
398 comparison, just over 20% of all species registered in the European Register of Marine Species
399 (ERMS) and almost 50% of species listed in the AZTI Marine Biotic Index (AMBI) have been
400 barcoded (Weigand et al., 2019). The percentage of barcoded European freshwater invertebrates
401 used in environmental monitoring reaches 64.5%, and when considering only Peracarida, 24% of
402 ERMS species, 45% of AMBI and 82% of freshwater monitored taxa have been barcoded
403 (Weigand et al., 2019). It has to be emphasized however, that only ERMS lists all marine
404 invertebrates from European region, while both other datasets studied by Weigand et al. (2019)
405 consists of a subset of species from this area. More specific studies of Iberian macroinvertebrates
406 revealed that ca. 40% of amphipod species possess barcodes (Leite et al., 2020; Múrria et al.,
407 2020;). Hestetun et al. (2020) conducted a barcode library gap-analysis of the benthic
408 macrofauna of one region of the North Sea, which indicated the barcode coverage varying from
409 42.4% to 61% (depending on the calculation method). This indicates that for smaller subset of
410 taxa and specified geographic region it is much easier to produce good barcode coverage. It can
411 be concluded that although Amphipoda are an actively studied taxonomic group where scientists

412 increasingly use molecular methods, this diverse and abundant macrofaunal taxon is still
413 insufficiently represented in the BOLD barcode library.

414 **Quality of amphipod barcode library**

415 In order to provide a trusted barcode for a particular species, at least one good quality sequence
416 associated with a species-level identification provided by taxonomic specialist is required as an
417 absolute minimum. However, a single sequence cannot provide information about intraspecific
418 variation, and overlooked contamination of the sample will mean the sequence cannot be
419 validated. As such, it is advisable to provide a small number of sequences to characterise each
420 taxon. The recently proposed barcode quality auditing system suggests providing at least three
421 sequences to enable proper barcode evaluation (Oliveira et al., 2016; Fontes et al., 2020).
422 Unfortunately, as we have shown in the case of Amphipoda, globally more than half of BINs are
423 represented by only 1-2 sequences in BOLD. This low number of sequences places them in
424 category D of the Oliveira et al. (2016) system, indicating the existing data is insufficient for use
425 as trusted barcodes. Similar observations for a restricted amphipod dataset are made by Fontes et
426 al. (2020).

427 Due to methodological differences it is impossible to make direct comparisons of our data with
428 the results of the gap analysis of aquatic organisms in European waters (Weigand et al., 2019),
429 but re-calculation of their data shows much improved barcode coverage. Among all freshwater
430 invertebrates 65% of taxa barcoded are represented by more than five sequences, while this
431 percentage rises to 77% when considering only freshwater Peracarida. This proportion of high
432 quality datasets diminishes when marine taxa are considered; with 52% of the marine species
433 from the AMBI list and 45% those listed in ERMS having at least five barcodes available. These
434 numbers do not change when considering only marine Peracarida (52% and 46% of the ones

435 presented in AMBI and ERMS lists, respectively). Our analysis of Amphipoda shows opposite
436 pattern with about 1/4-1/3 of BINs represented by more than five sequences but the good
437 barcode coverage observed by Weigand et al. (2019) may be biased by the fact that they targeted
438 the species used in water quality assessment programs. Because of their practical use such taxa
439 receive more scientific interest and it may be assumed that their barcoding is prioritized by
440 different institutions.

441 The amphipod BINs that have the largest numbers of sequences in BOLD are often the result of
442 detailed studies of targeted species, which have produced large numbers of sequences as a
443 secondary aim of the study. For example, 750 out of the 801 sequences in BOLD of terrestrial
444 *Orchestoidea tuberculata* come from a single study by Brante et al. (2019); 406 records out of
445 411 sequences in BOLD of freshwater *Dikerogammarus haemobaphes* (Eichwald, 1841) come
446 from Jazdzewska et al. (2020); while 232 records of 235 sequences in BOLD of marine *Caprella*
447 *scaura* Templeton, 1836 come from Cabezas et al. (2014). The disproportional representation
448 between the few species that are very thoroughly studied and the remaining majority of species
449 that are represented only by a single, or a low number of sequences emphasises the need for
450 more targeted sampling of less common species.

451 **Best studied families and cryptic diversity**

452 Almost half of the 239 known amphipod families are represented in BOLD. However, only ten
453 of these families have more than 50% of their associated species sequenced. It is important to
454 underline that there are 18 families in BOLD that do not have species-level identifications, but
455 have records left at the family or genus level. A small effort to provide trusted species-level
456 identifications for these taxa will greatly improve barcode coverage of the Amphipoda,
457 particularly if they represent species already known to science.

458 Another concern that has arisen as part of this study relates to the format of temporary names in
459 GenBank and BOLD, the different requirements by users for their input, and how this has
460 changed following development of the databases. In GenBank, the incorporation of temporary
461 names or codes is allowed (referred to as placeholder names in GenBank). In 2010, a large
462 amount of COI data was incorporated into the BOLD database. The identifications associated
463 with each of these imported sequences were included verbatim from GenBank. BOLD users,
464 however, were originally able to use temporary names in the database only in private
465 projects/dataset and when opening their data for public they were expected to provide the
466 identification to the lowest taxonomic level possible (e.g. genus) and to provide the temporary
467 name (e.g. incorporating “cf.” or “aff.”) as a taxonomy note (that has happened to the authors of
468 the present paper). However, in BOLD a taxonomy note is only visible when the specimen page
469 is open, and not in a general search. Recently, we have learnt that open nomenclature identifiers
470 (such as ‘cf.’ and ‘aff.’) are accepted by BOLD, but it may be assumed that numerous records
471 remain at a higher taxonomic level, with more detailed identifications available that are hidden
472 from general searches. This discrepancy in dealing with temporary names has become apparent
473 when analysing the whole dataset as part of this study. In particular, the inconsistent use of
474 temporary names in these databases mean that it is very difficult to differentiate between
475 temporary names which are being used to refer to species that are new to science, and those
476 which have remained at a higher taxonomic level because they were simply not identified further
477 (which could be for a variety of reasons). Molecularly well-defined temporary names for new
478 species are likely to become more abundant and therefore critical to our knowledge of
479 biodiversity in the coming years, and we need to ensure they are managed carefully and
480 consistently. Recommendations for the use of open nomenclature have been proposed recently to

481 attempt to standardise and overcome these issues (Sigovini, Keppel & Tagliapietra, 2016; Horton
482 et al., 2021) and it is hoped that these standard formats will be considered for use in both BOLD
483 and GenBank.

484 Barcode coverage of families varies depending on the species richness. For species rich families
485 it is around 10%, while coverage is increased for moderate and low species rich taxa. This is not
486 surprising considering it is much easier to receive better coverage for monotypic families or
487 those represented by only a few species. The best studied families are the ones that remain under
488 the interest of large working groups who focus on studying specific families (e.g. Hou, Fu & Li,
489 2007; Hou, Sket & Li, 2014; Mamos et al., 2014; Wysocka et al., 2014; Delić et al., 2017a; Delić
490 et al., 2017b; Fišer et al., 2017; Copilaş-Ciocianu, Sidorov & Gontcharov, 2019). It is worth
491 noting that providing barcodes is generally more a “by-product” of other analyses than the goal
492 per se. Another issue that should be emphasized is that species rich families are proportionally
493 under studied. This is important because they usually do not only group many species but very
494 often the species from these families constitute the majority of amphipods characterizing
495 different assemblages. This is clearly shown by the Phoxocephalidae (1% of the 367 known
496 species are barcoded), Ampeliscidae (7% of 312 of the known species barcoded) or
497 Oedicerotidae (10% of the 246 known species barcoded), all constitute very large and important
498 components of marine benthic communities worldwide (Brandt, 1993; Weissappel &
499 Svavarsson, 1998; Frutos & Sorbe, 2017; Brix et al., 2018;). Another example is provided by the
500 Caprellidae (6% of the 443 known species are barcoded) which are an important part of many
501 fouling communities (e.g., Ros, Vázquez-Luis & Guerra-García, 2013; Ros et al., 2013;) and
502 where proper species identifications are crucial in the context of growing transport with their

503 resulting potential alien species invasions (op. cit.). The studies of these families should be
504 prioritized in order to support marine monitoring programs based on barcode libraries.

505 The analysis of the amphipod BINs with a species-level identification showed that there were
506 only a few cases where multiple names were associated with a single molecular unit. A quarter
507 of these cases resulted from the misidentification of single individuals within a taxon. In some
508 cases different names were associated with the description of new species (present in the
509 database under both former and newly established name). Problems with morphological
510 identification of cryptic species and the lack of well established diagnostic characters within
511 closely related species may also be the reason of the presence of multiple names for single BIN.

512 The above problems have been recognized within *Gammarus ochridensis* Schäferna, 1926
513 species complex that is the group of morphologically very similar species of which two
514 *Gammarus cryptosalemaai* Grabowski, Wysocka & Mamos, 2017 and *Gammarus*
515 *cryptoparechiniformis* Grabowski, Wysocka & Mamos, 2017 are recognizable only based on
516 molecular data (Wysocka et al., 2013; Grabowski, Wysocka & Mamos, 2017). This indicates that
517 generally BOLD may be considered a trusted tool for species identification. Our analyses
518 showed that in the majority of cases, a single BIN was characterising a single species, which is
519 congruent with the results of other similar studies (Fontes et al., 2020; Leite et al., 2020). Some
520 morphologically identified species were represented by two or even three BINs, which can
521 indicate overlooked diversity. It has been noted however, that sometimes due to the methodology
522 used during BIN-identification and the threshold used (2% of similarity, Ratnasingham &
523 Hebert, 2013) some valid species may be split into two or more BINs (Lörz, Jazdzewska &
524 Brandt, 2018; Jazdzewska & Mamos, 2019). This happens more frequently when the sample size
525 is small and the intraspecific variation range cannot be adequately assessed. In such cases, the

526 use of additional genes or other data analysing methods may help to decide the proper species
527 delineation. The present study revealed 19 morphological species that were represented by 11 or
528 more BINs. This multi-BIN representation was much more common in freshwater environments,
529 where 17 species with potential cryptic diversity were observed. The existence of such high
530 cryptic diversity especially in European waters was recognized by authors of the original works
531 (Witt, Threlhoff & Hebert, 2006; Bauzà-Ribot et al., 2011; Major et al., 2013; Mamos et al., 2014;
532 Wysocka et al., 2014; Delić et al., 2017a; Delić et al., 2017b, Fišer et al., 2017; Tomikawa et al.,
533 2017) most recently confirmed also by Wattier et al. (2020). A detailed study of the available
534 barcodes and cryptic diversity of the Gammaridae and other representatives of the superfamily
535 Gammaroidea is in preparation (Mamos pers. comm.). The large representation of freshwater
536 taxa forming cryptic species complexes (especially in Europe) can be partly explained by the
537 geological events that shaped the European freshwater system (Wysocka et al., 2014; Mamos et
538 al., 2016; Wattier et al., 2020). Presence of marine cryptic or pseudo-cryptic species have also
539 been reported (Havermans, 2016; Verheye, Backeljau & d'Udekem d'Acoz, 2016;), but the
540 extent of molecular studies of amphipods from this realm is much smaller and as a result cryptic
541 species may have been overlooked. A study of the marine genus *Apoehyale* showed high
542 diversification of species within the genus, and confirms that more studies are required to
543 correctly identify species diversity and uncover cryptic diversity in marine taxa (Desiderato et
544 al., 2019). Cases of highly diverse nominative species usually come from studies based in a
545 single research group that was already aware of the high diversity within the taxon. There are,
546 however, cases where multiple BINs have received the same identification but this was carried
547 out by different authors at different times (without comparison of the material) and it is difficult
548 to judge if the observed diversity is a result of the existence of a cryptic species or of

549 misidentification of the species. In such cases it is impossible to decide which of the BINs
550 represents the known species and which are cryptic/new species that require more detailed study
551 (Jazdzewska et al., 2018). A detailed analysis of the species represented by several BINs was not
552 the focus of the present study, but it should be a priority for BOLD to identify such cases and
553 inform users about the presence of possible cryptic taxa. Users of BOLD who seek to obtain
554 identification for their own sequence should be notified that the specimen they have may belong
555 to a group of cryptic species so that the taxonomic identification can be treated with caution. It is
556 also highly recommended that mistakes or problematic issues that are found in the database are
557 corrected and published e.g. the case of *Hyperiella antarctica* Bovallius, 1887/*H. dilatata*
558 Stebbing, 1888 which was recently clarified by Havermans et al. (2019).

559 **Conclusions and Future recommendations**

560 We have conducted a gap-analysis of the barcode library using a single crustacean order, the
561 Amphipoda, as a model. The high diversity and the important role played by amphipods in the
562 aquatic ecosystem combine to highlight the need for accurate species identifications which are
563 required for biological monitoring programs. DNA-barcoding may speed up the identification
564 process, but success is dependent on the barcode library coverage and quality. Our gap analysis
565 has shown that although a large number of BINs (indicating putative species) was recognized by
566 BOLD still only 10 % of the amphipod species are represented by barcodes. Moreover, the
567 majority of BINs is represent by a very low number of sequences that make them unreliable
568 according to the quality control system. The geographical coverage is poor with vast areas of
569 Africa, South America and the open ocean acting as “white gaps”, also the level of barcoding
570 effort is skewed depending on the environment.

571 As such, we make the following recommendations (in order of priority), which will improve the
572 data currently held within BOLD, and we outline steps that are needed to provide a more equal
573 coverage of the sequence data within the Amphipoda, and thus improve the utility of the
574 database for a variety of applications, including species identification and biomonitoring.

575 1. Morphological identification of the already recognised BINs (that are missing species ID) if
576 the voucher specimens are available.

577 2. Analysis of the nominal species that are represented by more than one BIN, especially if
578 identifications represented by different BINs were produced by separate working teams.

579 3. Prioritised barcoding of representatives from families that are known to be important and
580 abundant components of communities; Phoxocephalidae, Ampeliscidae, Oedicerotidae, and
581 Caprellidae should be prioritised.

582 4. Targeted sampling programs for taxa coming from geographical regions with the least
583 knowledge.

584 5. Targeted sampling to obtain more sequences for taxa present in BOLD but represented by
585 small numbers of sequences (especially singletons), from different parts of the species' range if
586 possible.

587 6. Targeted programs to sequence type specimens stored in musea or to collect and study fresh
588 individuals from type localities if types are unsuitable for analyses.

589

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911 **Figure captions**

912 Figure 1. PRISMA 2020 work-flow diagram (Page et al., 2021). Summary of the data download,
913 identification and screening before analysis. All record removals were done by the leading
914 author of the paper.

915 Figure 2. Environmental origin of the amphipod records (A) and BINs (B) in BOLD database.

916 Figure 3. Geographic distribution of amphipod records expressed by sequences present in BOLD
917 (A – freshwater, B – marine, C – terrestrial). Dots indicate records with exact coordinates, for
918 records without latitude and longitude the country of origin was checked. Background color
919 of the country indicates this number per country.

920 Figure 4. Proportion of records (A) and BINs (B) with different level of identification within
921 freshwater, marine and terrestrial amphipod taxa.

922 Figure 5. Number of BINs represented by given number of sequences. Upper set (A, B, C) – all
923 BINs, lower set (D, E, F) – only BINs with complete species-level identification considered.

924 A, D – freshwater, B, E – marine, C, F – terrestrial taxa.

925 Figure 6. Number of nominal species represented by given number of BINs.

926

927 **Supplemental files list**

928

929 **Supplemental file 1**

930 - File format .xlsx

931 - Title: List of the doubled records indentified in the original dataset

932 - Description: The file consists of records that appeared to be doubled in the BOLD database.

933

934 **Supplemental file 2**

935 - File format .xlsx

936 - Title: The list of records analysed (after removal of doubled records).

937 - Description: The file consists of all records that was basis of the present study. The colors of
938 the cell of recordID indicates the environment: green - marine (including brakishwater and
939 fully marine taxa), red - freshwater (including freshwater and brakishwater taxa), yellow -
940 terrestrial, blue - environment not recorded.

941

942 **Supplemental file 3**

943 - File format .xlsx

944 - Title: List of BINs possessing more than one ID variant with notes on the identification

945 - Description: The file presents BINs that have received different identifications with details of
946 the ID and comments concerning the final identification used in the study.

947

948 **Supplemental file 4**

949 - File format .xlsx

950 - Title: List of BINs for which the environment was not able to be assessed.

951 - Description: The file presents BINs for which the available data did not allow to specify the
952 environment.

953

954 **Supplemental file 5**

955 - File format .xlsx

956 - Title: List of BINs with the largest number of records (blue - freshwater, green - marine, yellow
957 - terrestrial taxa).

958 - Description: The file presents BINs that are represented by the largest number of records. The
959 information about environmental origin of associated species are provided.

960

961 **Supplemental file 6**

962 - File format .xlsx

963 - Title: Nominal species with the largest number of BINs identified.

964 - Description: The file presents nominal species for which the largest number of BINs has been
965 identified. Environmental origin of species is also provided.

966

967 **Supplemental file 7**

968 - File format .xlsx

969 - Title: List of amphipod families with number of accepted and barcoded species as well as
970 information of the barcoding coverage within family. Families within each category with the
971 highest barcoding coverage indicated in bold.

972

973 **Supplemental file 8**

974 - File format .docx

975 - PRISMA 2020 checklist

Table 1 (on next page)

Representation of amphipod families in BOLD.

* in parentheses the number of families without barcoded species but with at least one BIN identified to the genus (g) or family (f) level.

- 1 Table 1. Representation of amphipod families in BOLD. * in parentheses the number of families
- 2 without barcoded species but with at least one BIN identified to the genus (g) or family (f) level.

Number of families	
without any barcoded species	117 (+ 13g, 5f)*
with up to 10% barcoded species	47
with 11-20% barcoded species	24
with 21-50% barcoded species	24
with >50% barcoded species	10

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Table 2 (on next page)

Number of amphipod species in each realm with indication of their barcode quality according to grading system from Fontes et al. (2020).

A - consolidated concordance, B - basal concordance, C - multiple BINs for single morphospecies, D - insufficient data; for more detailed explanation of grading system, see Material and methods section.

- 1 Table 2. Number of amphipod species in each realm with indication of their barcode quality
- 2 according to grading system from Fontes et al. (2020). A – consolidated concordance, B – basal
- 3 concordance, C – multiple BINs for single morphospecies, D – insufficient data; for more
- 4 detailed explanation of grading system, see Material and methods section.

	A	B	C	D	all species
All species	100	155	276	468	999
Freshwater spp.	31	55	140	225	451
Marine spp.	58	92	120	226	496
Terrestrial spp.	11	8	16	17	52

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Table 3(on next page)

Number of accepted families and species of Amphipoda (according to WAD accessed on 17-07-2020), number of families with representation in BOLD, number of species present in BOLD and mean coverage of barcodes in amphipod families represented in BOLD.

- 1 Table 3. Number of accepted families and species of Amphipoda (according to WAD accessed
- 2 on 17-07-2020), number of families with representation in BOLD, number of species present in
- 3 BOLD and mean coverage of barcodes in amphipod families represented in BOLD.

	No. of families	No. of species	No. of families with species representation in BOLD	No. of species present in BOLD	Mean barcode coverage [%] of those families with representation in BOLD
Very species rich families (>100 spp.)	33	7302	32	714	8
Species rich families (31-100 spp.)	30	1633	22	127	10
Moderately species rich families (11-30 spp.)	53	979	26	107	21
Low species rich families (<10 spp.)	123	416	26	51	49

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Table 4(on next page)

Percent of families with species belonging to different quality grading categories (Fontes et al., 2020).

A - consolidated concordance, B - basal concordance, C - multiple BINs for single morphospecies, D - insufficient data; for more detailed explanation of grading system, see Material and methods section.

- 1 Table 4. Percent of families with species belonging to different quality grading categories
 2 (Fontes et al., 2020). A – consolidated concordance, B – basal concordance, C – multiple BINs
 3 for single morphospecies, D – insufficient data; for more detailed explanation of grading system,
 4 see Material and methods section.

	% of families				
	All families	Very species rich families (>100 spp.)	Species rich families (31-100 spp.)	Moderately species rich families (11-30 spp.)	Low species rich families (<10 spp.)
At least one sp. in the category A	32.4	65.6	31.8	16	7.7
At least one sp. in the category B	31.4	28.1	36.4	28	34.6
At least one sp. in the category C	10.5	0	13.6	24	7.7
At least one sp. in the category D	25.7	6.3	18.2	32	50
Number of families	105	32	22	25	26

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Figure 1

PRISMA 2020 work-flow diagram (Page et al., 2021).

Summary of the data download, identification and screening before analysis. All record removals were done by the leading author of the paper.

Figure 1. PRISMA 2020 work-flow diagram (Page et al., 2021). Summary of the data download, identification and screening before analysis. All record removals were done by the leading author of the paper.

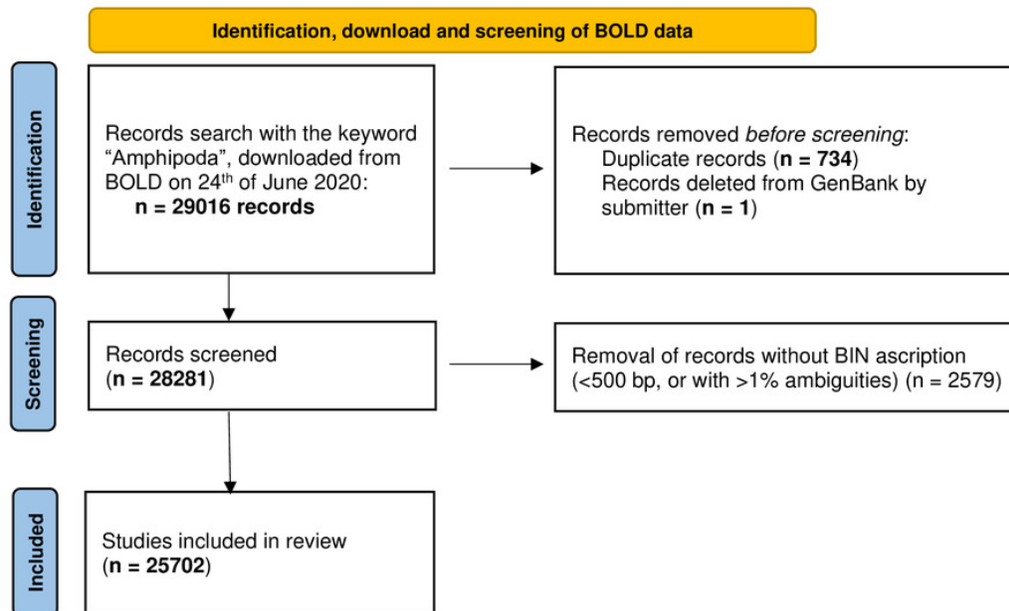


Figure 2

Environmental origin of the amphipod records (A) and BINs (B) in BOLD database.

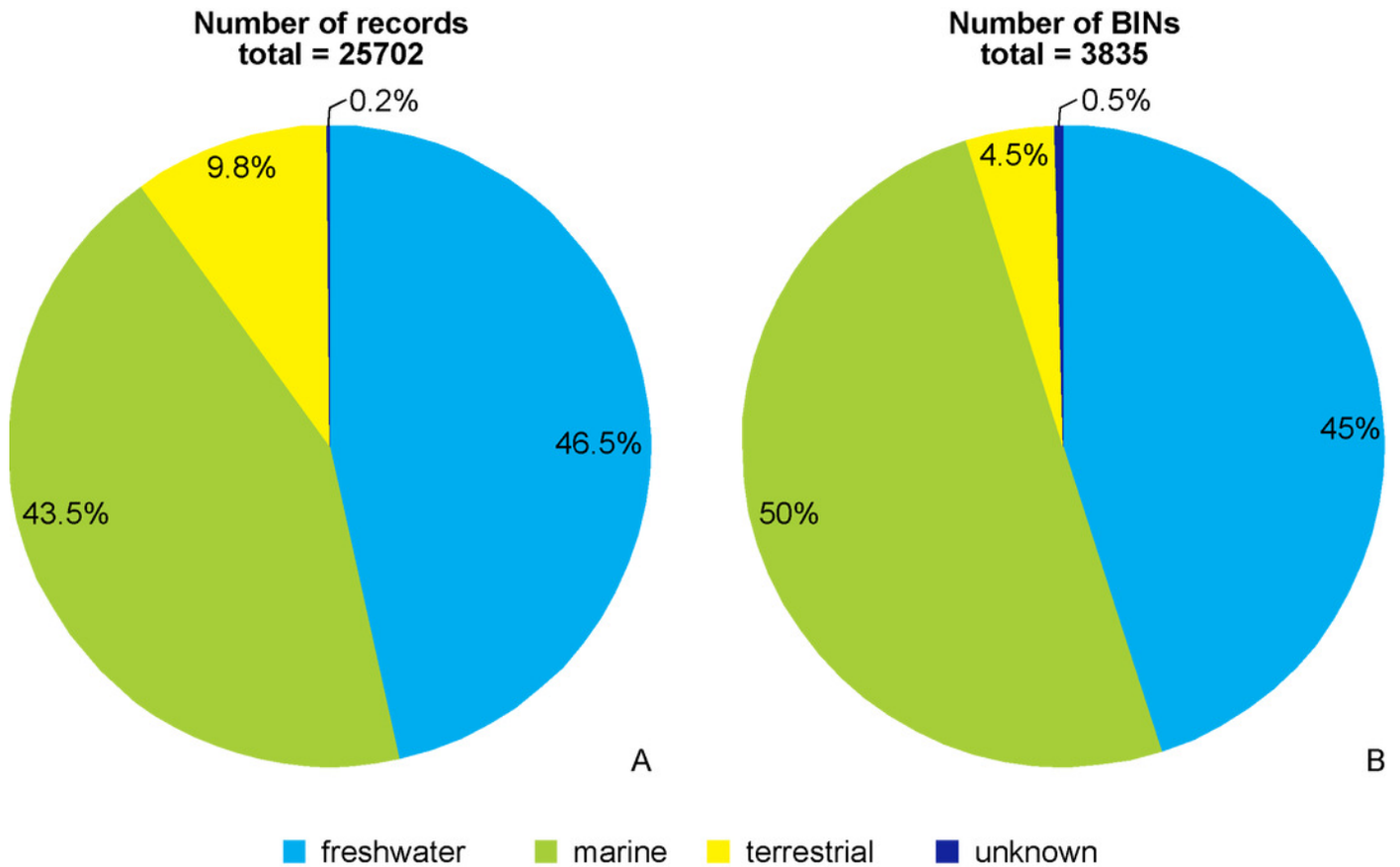


Figure 3

Geographic distribution of amphipod records expressed by sequences present in BOLD (A - freshwater, B - marine, C - terrestrial).

Dots indicate records with exact coordinates, for records without latitude and longitude the country of origin was checked. Background color of the country indicates this number per country.

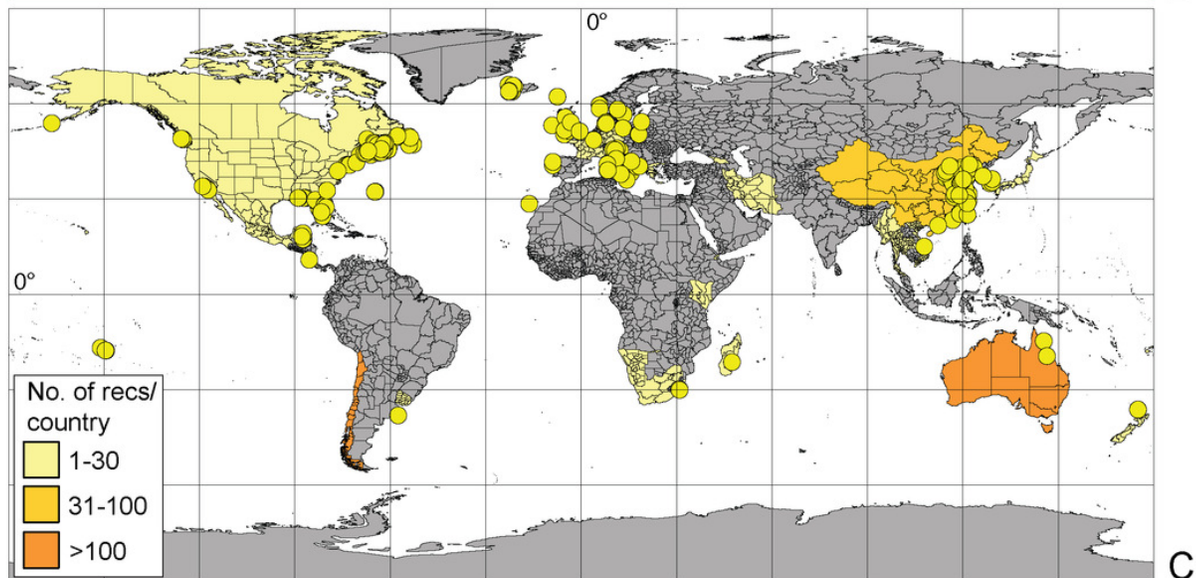
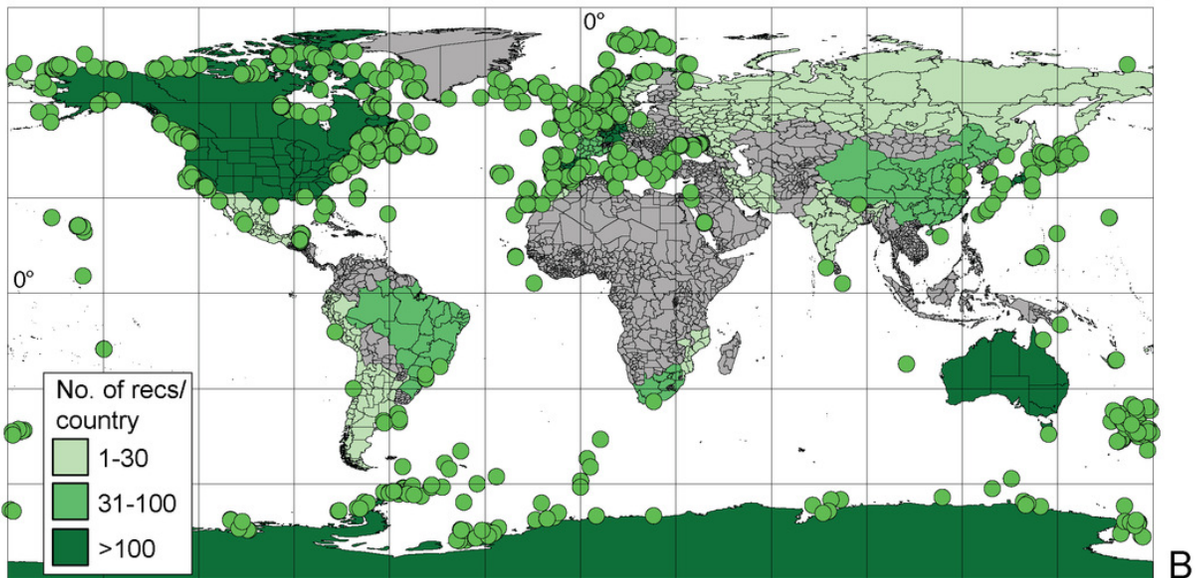
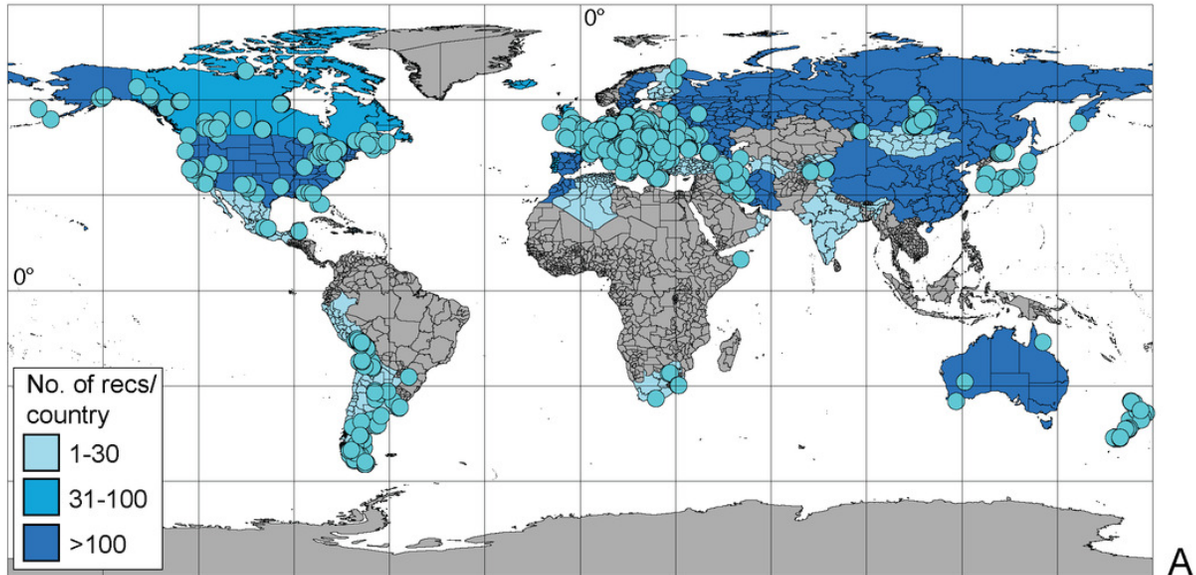


Figure 4

Proportion of records (A) and BINs (B) with different level of identification within freshwater, marine and terrestrial amphipod taxa.

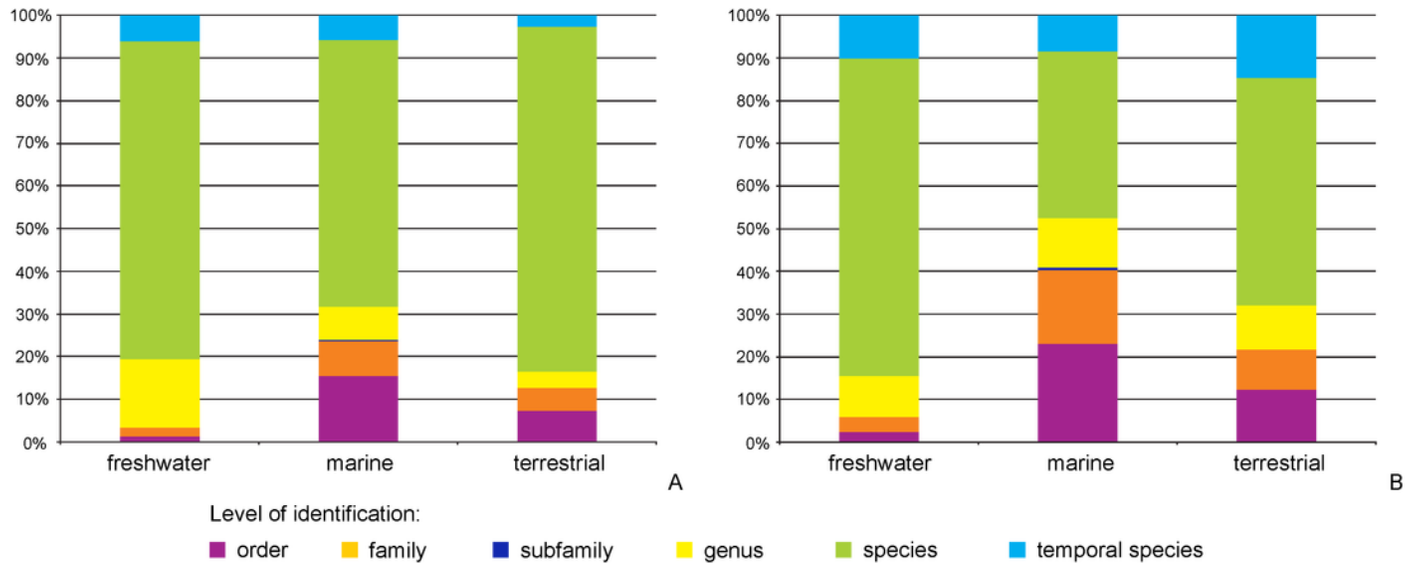


Figure 5

Number of BINs represented by given number of sequences.

Upper set (A, B, C) - all BINs, lower set (D, E, F) - only BINs with complete species-level identification considered. A, D - freshwater, B, E - marine, C, F - terrestrial taxa.

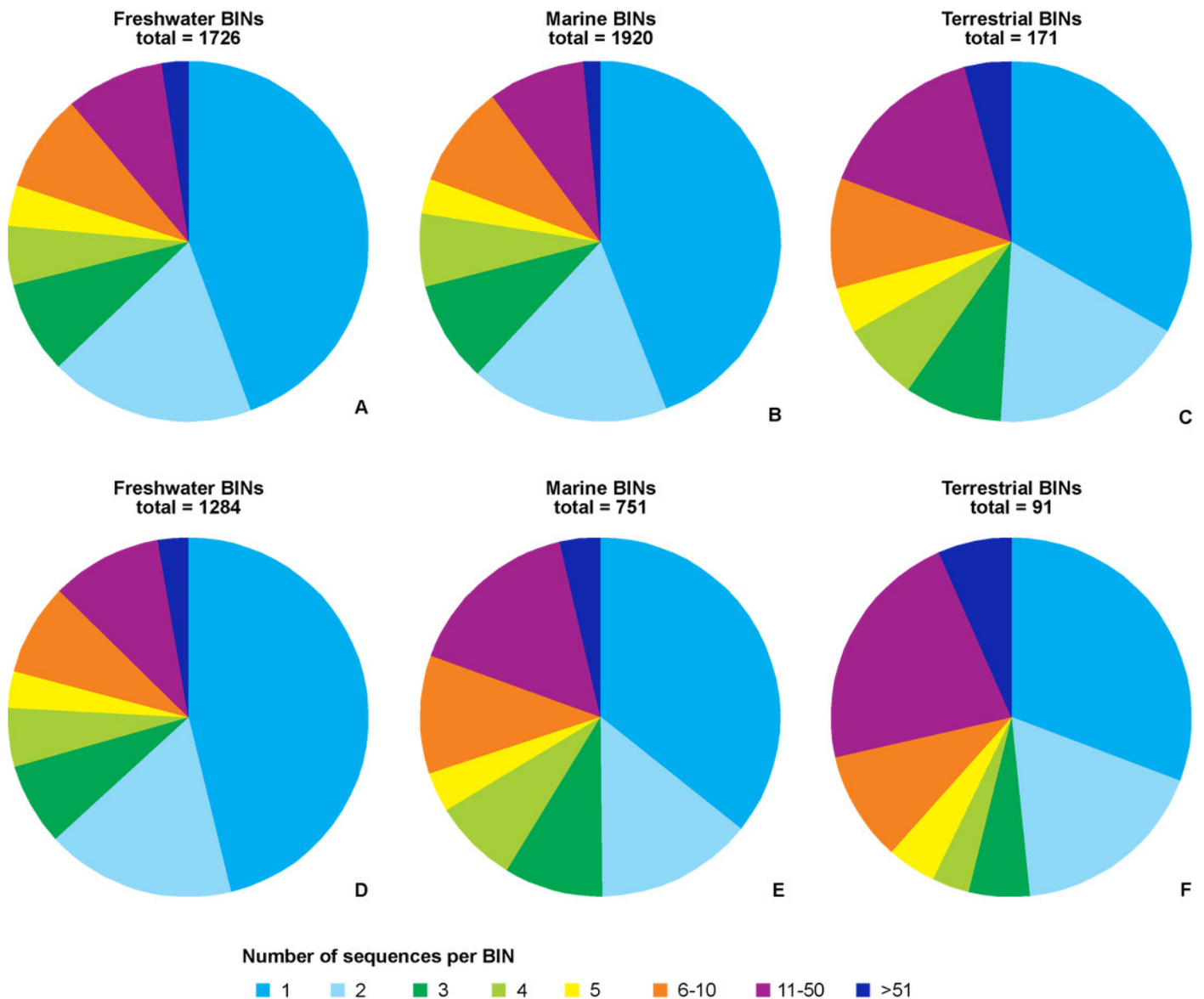


Figure 6

Number of nominal species represented by given number of BINs.

