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# **An annotated reference library for supporting DNA metabarcoding analysis of aquatic macroinvertebrates in French freshwater environments**

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2 **aquatic macroinvertebrates in French freshwater environments**

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33 **Abstract**

34 Freshwater ecosystems are increasingly threatened by human activities, leading to  
35 biodiversity loss and ecosystem degradation. Effective biodiversity monitoring,  
36 particularly through the use of aquatic macroinvertebrates as bioindicators, is crucial  
37 for assessing ecological health. While traditional morphological methods face  
38 limitations, DNA metabarcoding offers higher accuracy and efficiency in species  
39 identification using environmental DNA. However, the success of metabarcoding is  
40 contingent on the quality of reference libraries, which are often incomplete or biased.  
41 This study aimed to construct a comprehensive COI-based DNA barcode library for  
42 freshwater macroinvertebrates in France, specifically targeting short gene regions  
43 amplified with fwhF2/fwhR2N primers, suitable for degraded DNA. A list of species  
44 occurring in French freshwater ecosystems was established from official national  
45 checklists and Alpine lake surveys. The resulting library was analyzed for taxonomic  
46 completeness, barcode coverage and genetic diversity. The checklist consisted of  
47 2,841 species across 10 phyla, for which 56% had at least one COI-5P sequences  
48 available in the Barcode of Life Data System (BOLD). Alignment challenges with the  
49 primers were identified for certain taxa, particularly among Coleoptera, Diptera, and  
50 Malacostraca. The genetic diversity approached by the number of haplotypes per  
51 species highlighted that most of the species have limited diversity, with only 3 species  
52 having more than 100 haplotypes. Finally, this study showed that a total of 57  
53 haplotypes were shared among 116 distinct species. This work emphasized the need  
54 for expanded sequencing efforts to improve barcode coverage and highlighted the  
55 pitfalls associated with the use of these primers for further biodiversity assessment of  
56 macroinvertebrates with degraded DNA.

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58 **Keywords:** freshwater, COI, fwhF2 primers, macroinvertebrates, gap analysis,  
59 metabarcoding, haplotype diversity.

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72 **Introduction**

73 Freshwater ecosystems face escalating anthropogenic pressures threatening their  
 74 functionality (Reid et al. 2019) and leading to significant biodiversity loss (Young et al.  
 75 2016, Borgwardt et al. 2019). In this context, biodiversity monitoring is crucial to  
 76 provide relevant ecological diagnoses and support the management and conservation  
 77 of these ecosystems and the vital services they provide. Aquatic macroinvertebrates  
 78 are widely used as bioindicators due to their abundant presence, high species diversity  
 79 and sensitivity to both anthropogenic and natural disturbances (Hering et al. 2006).  
 80 These organisms primarily inhabit the littoral zone and play pivotal roles in community  
 81 assembly and food web dynamics, contributing significantly to beta diversity and  
 82 metacommunity structure across diverse aquatic habitats (May 2019). Consequently,  
 83 they serve as key indicators in environmental monitoring programs, facilitating  
 84 assessments of the ecological status of aquatic ecosystems as mandated by the EU  
 85 Water Framework Directive (WFD) and national legislation (Mondy et al. 2012). The  
 86 standardized methodology for macroinvertebrate biomonitoring in Europe is based on  
 87 morphological identification through binocular microscopy (ANFOR, 2010; CEN 2019).  
 88 This methodology encounters significant challenges, including the labor-intensive and  
 89 expensive nature of collecting and sorting individual benthic invertebrates, which  
 90 hinders its broader adoption in routine biomonitoring (Blackman et al. 2019; Bonada et  
 91 al. 2006). Moreover, routine morphological identification poses drawbacks such as  
 92 high expertise demands and lower taxonomic resolution (Leese et al. 2016, 2018,  
 93 Bean et al. 2017, Hering et al. 2018). Notably, many taxa can only be identified at  
 94 limited taxonomic resolution (Caesar et al. 2006), without a timely effort of highly skilled  
 95 taxonomists. This is the case for Chironomids, a common group in freshwaters,  
 96 frequently classified at the family level due to challenges in species or genus  
 97 identification (Beermann et al. 2018). Additionally, immature life stages lacking  
 98 diagnostic morphological traits hinder accurate species identification and therefore limit  
 99 the proper quality assessment of freshwater ecosystems (Sweeney et al. 2011). These  
 100 challenges underscore the need for innovative approaches to enhance the efficiency  
 101 and accuracy of macroinvertebrate biomonitoring practices. The recent advent of DNA  
 102 metabarcoding techniques, integrating amplicon barcoding with high-throughput  
 103 sequencing, represents a promising advancement in biomonitoring applications  
 104 (Deiner et al. 2017, Pawlowski et al. 2018, Carraro and Altermatt 2024), providing a  
 105 valuable complement to morphology-based approaches for species identifications.  
 106 Metabarcoding finds application in analyzing environmental DNA (eDNA) samples  
 107 obtained from water, biofilm or sediment (Sakata et al. 2020; Valentini et al. 2016,  
 108 Rivera et al. 2021, Gauvin et al. submitted) or DNA extracts derived from a  
 109 homogenate of the sample's fauna (Taberlet et al. 2012). These molecular innovations  
 110 enable simultaneous processing of multiple samples, identification of small taxa,  
 111 immature or larval stages, and offer increased sensitivity and specificity, often  
 112 revealing hidden diversity, while enhancing time and cost effectiveness (Shokralla et  
 113 al. 2012, Pochon et al. 2015, Holman et al. 2019). These advantages facilitate direct  
 114 comparison among sites and studies and enable higher spatial-temporal frequency in  
 115 monitoring due to increased throughput (Bush et al. 2019). The efficiency of  
 116 macroinvertebrates DNA metabarcoding relies heavily on primer sets' effectiveness  
 117 across a broad taxonomic spectrum. The recovery rate of taxa using metabarcoding is

118 contingent upon various factors, including the taxonomic resolution of the gene marker  
 119 employed (e.g. COI or ribosomal markers like 16S, e.g, Elbrecht et al. 2016), amplicon  
 120 length (Meusnier et al. 2008), primer universality, and the number of primer pairs  
 121 utilized to amplify target taxonomic groups (Elbrecht and Leese 2015, Gibson et al.  
 122 2015). Specifically, a segment of the cytochrome c oxidase subunit I (COI) gene has  
 123 emerged as the standard DNA barcoding marker for most animal groups (Hebert et al.  
 124 2003a), with over 95% of species in diverse animal groups exhibiting distinctive COI  
 125 sequences in test assemblages (Hebert et al. 2003b, 2004). Given the considerable  
 126 phylogenetic diversity among macroinvertebrates, the high taxonomic resolution and  
 127 existing reference data for the COI marker, make it a judicious choice for  
 128 metabarcoding freshwater macroinvertebrate communities (Ratnasingham and Hebert  
 129 2007, Andújar et al. 2018). Furthermore, well-established barcoding gaps for the COI  
 130 marker in freshwater macroinvertebrates underscore its suitability for such applications  
 131 (Zhou et al. 2009). Numerous primers of COI gene of varying base pair sizes are  
 132 available in the literature for DNA amplification of macroinvertebrates communities.  
 133 Environmental DNA released by target taxa can degrade rapidly (Seymour et al. 2018).  
 134 Consequently, for amplifying degraded DNA, that is extracted from water samples for  
 135 example, targeting a short COI marker region is suggested to enhance amplification  
 136 success (Barnes et al. 2014, Jelger Herder et al. 2014). The effectiveness of DNA  
 137 metabarcoding programs relies on open, comprehensive and accurate reference  
 138 sequence libraries (Briski et al. 2016, Oliveira et al. 2016, Weigand et al. 2019),  
 139 ensuring precise taxonomic assignment (Richardson et al. 2018, Rodriguez-Ezpeleta  
 140 et al. 2021). In that sense, incomplete DNA barcode libraries, as in poorly represented  
 141 species, pose a significant challenge, often leading to false negatives and  
 142 compromising biodiversity assessments (Ardura 2019, Leite et al. 2020, Duarte et al.  
 143 2021). Therefore, evaluating these gaps and the quality of sequence data in reference  
 144 libraries is imperative for the effective implementation of DNA-based tools in  
 145 biodiversity assessments (Duarte et al. 2020). Given the significance of reference  
 146 libraries completeness, primarily those that are pertinent in the context of the WFD,  
 147 numerous studies have been conducted to assess their representativeness by  
 148 comparing them with lists of described species (Trebitz et al. 2015, Weigand et al.  
 149 2019, Duarte et al. 2020, Leite et al. 2020, Specchia et al. 2020, Csabai et al. 2023).  
 150 Despite efforts to assess their representativeness, biases in taxonomic coverage  
 151 persist within reference libraries (Li et al. 2019, Weigand et al. 2019). Indeed,  
 152 numerous studies have emphasized the construction of tailored reference libraries to  
 153 suit the geographic context of the research area (Ficetola et al. 2021, Mugnai et al.  
 154 2023) and emphasized the importance of such databases to be refined with custom  
 155 sequences specific to local study areas and free from unexpected taxa. Additionally,  
 156 research by Abad et al. 2016 and Schenekar et al. 2020 has highlighted the value of  
 157 possessing DNA barcodes for local species. These studies have demonstrated that  
 158 DNA barcodes from local organisms can improve the accuracy of taxonomic  
 159 assignments and reveal previously unrecognized biodiversity, leading to adjustments  
 160 in taxonomic classifications among species. Our study is in line with the  
 161 recommendations proposed by Blackman et al. 2023, which advocate for compiling a  
 162 comprehensive list of target species within the study area and assembling accurate  
 163 sequences pertinent to those species. Our first objective is to assemble a list of  
 164 macroinvertebrate species known to be present in French freshwater ecosystems. The

165 second objective is to construct a DNA barcode library utilizing COI primers,  
 166 particularly focusing on a short region of this gene which is highly effective when  
 167 targeting degraded DNA (fwhF2/fwhR2N, 205pb, Vamos et al. 2017), to identify  
 168 freshwater macroinvertebrate fauna in France using metabarcoding techniques and to  
 169 assess its genetic completeness. Our third objective is to identify the gaps of this new  
 170 reference library, thereby providing insight into its limitations and facilitating data  
 171 interpretation for futures users.

172

## 173 **Material and methods**

### 174 **Checklist constitution for French metropolitan freshwater macroinvertebrates** 175 **species**

176 In order to establish a checklist of macroinvertebrates taxa of freshwater aquatic  
 177 habitats in metropolitan France, three checklists were compiled. The first one comes  
 178 from PERLA, an interactive tool managed by a French regional environmental agency  
 179 (DREAL Auvergne-Rhône-Alpes) accessible at [http://www.perla.developpement-](http://www.perla.developpement-durable.gouv.fr/index.php)  
 180 [durable.gouv.fr/index.php](http://www.perla.developpement-durable.gouv.fr/index.php). PERLA serves as a national comprehensive checklist for  
 181 water managers and encompasses larvae, nymphs and adults across various  
 182 taxonomic groups, including insects, molluscs, crustaceans, and worms, among  
 183 others, found in rivers and aquatic ecosystems. In the manuscript, we will refer to this  
 184 checklist as the 'French aquatic ecosystems checklist'. The second checklist is coming  
 185 from macroinvertebrate surveys conducted in four Alpine lakes (Lake Geneva, Lake  
 186 Annecy, Lake Bourget and Lake Tignes) from 2015 to 2022. In the manuscript, we will  
 187 refer to this checklist as the 'French Alpine lakes checklist'. The third checklist we used  
 188 was from a French NGO, Opie-benthos ([https://www.opie-benthos.fr/opie/monde-des-](https://www.opie-benthos.fr/opie/monde-des-insectes.html)  
 189 [insectes.html](https://www.opie-benthos.fr/opie/monde-des-insectes.html)), studying freshwater insect's taxonomy and diversity. Currently, Opie-  
 190 benthos leads comprehensive inventories across France, specifically targeting insect  
 191 species undergoing a part of their life cycle within diverse aquatic ecosystems. In the  
 192 manuscript, we will refer to this checklist as the 'French aquatic insect's species list'.  
 193 These three checklists were merged into a single one. When the taxonomic resolution  
 194 of taxa was limited to a level above species (genus, family, class, phylum), we  
 195 consulted the taxonomic referential of the National Inventory of Natural Heritage  
 196 (TAXREF v17.0 2024, available at  
 197 <https://inpn.mnhn.fr/telechargement/referentielEspece/taxref/17.0/menu>) to select  
 198 species from those ranks. The phylum Acanthocephala and Rotifera, the classes  
 199 Copepoda and Ostracoda (Arthropoda, Crustacea) and the families Succineidae  
 200 (Arthropoda, Insecta, Diptera) and Hydrachnidae (Arthropoda, Arachnida,  
 201 Trombidiformes), initially listed in French aquatic ecosystems checklist, were omitted  
 202 from our search from TAXREF v17.0 2024 as they are not considered as freshwater  
 203 macroinvertebrates according Tachet et al. 2000. Once those macroinvertebrates  
 204 species were retrieved for each taxonomic levels without species identification, several  
 205 filters from TAXREF were applied to select only French metropolitan freshwater  
 206 species from this taxonomic referential. First, we selected only the species occurring  
 207 in the following habitats: freshwater, marine and freshwater, brackish water,  
 208 continental (land and freshwater). Then, from this updated list, only species occurring

209 in metropolitan France were kept. Finally, species characterized by one those 8 status  
210 over 15 status in total were selected (present (native or undetermined), endemic, sub-  
211 endemic, cryptogenic, introduced, invasive introduced, non-established introduced  
212 (including cultivated / domestic), occasional). The checklists described above and the  
213 summarized species list resulting from these compilations, inclusive of all species from  
214 the mentioned inventories, is referred as the “French metropolitan freshwater  
215 macroinvertebrates species list” and all together are available at  
216 <https://doi.org/10.57745/LMOXEW> in the ‘Checklists composition data’ file for future  
217 reference.

218

## 219 Sequences origin and cleaning steps

220 All the sequences for those species listed in the French metropolitan freshwater  
221 macroinvertebrates species list, whatever the gene, were downloaded from October  
222 2023 to May 2024, from the Barcode of Life Data System v4 -Bold- repository  
223 (<https://boldsystems.org/index.php/databases>). We opted to utilize Bold as reference  
224 library instead of GenBank due to concerns regarding the latter's unverified submission  
225 process, which frequently results in misannotated sequences (Kozlov et al. 2016,  
226 Locatelli et al. 2020, Steinegger and Salzberg 2020). The presence of stop codons  
227 were checked among the retrieved sequences and none were found. The absence of  
228 sequences for a given species may indicate either that the search in BOLD returned  
229 "Unmatched terms" or that while the species was found, no sequences were publicly  
230 available, rendering them private sequences (regardless of the reason for their  
231 absence, they were all defined as "not available sequences" for this analysis). Before  
232 concluding that a species lacked records in BOLD for the species from the French  
233 aquatic ecosystems checklist and French Alpine lakes checklist, synonyms were  
234 searched using INPN. The details of taxa resulting from this taxonomic harmonization  
235 are available in Table S1. From this file containing all the sequences retrieved  
236 (whatever the gene) for the queried species of the French metropolitan freshwater  
237 macroinvertebrates species list, the COI-5P sequences belonging to the COI gene  
238 were selected. All the remaining sequences were aligned with the fwhF2/fwhR2N  
239 primer pair using MAFFT version 7 (Auto) (<https://mafft.cbrc.jp/alignment/server/>), by  
240 genus or species group (as in all sequences assigned to the same genus or specie  
241 were aligned together). Then, sequences characterized by gaps, insufficiencies in  
242 length relative to the COI barcode standard (shorter than 205 pb), or identical (same  
243 genetic sequence for one species) from the sequences file, were removed using  
244 Jalview (<https://www.jalview.org/>). Due to the possibility of identical sequences being  
245 shared among different species, certain species may have been excluded during this  
246 stage, as a result of sequence overlap with other taxa. To address this issue, the list  
247 of species obtained from this step was cross-referenced with the initial species list  
248 following the alignment process. This comparison identified any missing species,  
249 whose sequences were then realigned individually to ensure their accurate  
250 representation. This final reference library containing only genetic sequences capable  
251 of being aligned by the fwhF2/fwhR2N primer pair is named ‘Aligned DNA library’  
252 hereafter.

253

254 **Graphical analysis**

255 We firstly analysed the taxonomic composition of the French metropolitan freshwater  
 256 macroinvertebrates species list. Secondly, to highlight any gaps in the availability of  
 257 sequences in our list of species, the taxonomic coverage of barcodes at various  
 258 taxonomic levels was assessed. Thirdly, we highlighted the taxonomic composition of  
 259 species with COI-5P sequences that did not align with fwhF2/fwhR2N primers to reveal  
 260 which species and taxonomic groups could be misrepresented in metabarcoding  
 261 studies using those primer pair. Then, to explore the species haplotype diversity, the  
 262 number of different haplotypes available relative to the number of species by phylum  
 263 was examined. The categorization of unique barcodes ranging from limited (<5  
 264 barcodes) to moderate (5-25 barcodes) to good (>25 barcodes), was adopted from  
 265 Trebitz et al. 2015, and the species with the highest number of unique sequences  
 266 (haplotype) exceeding 100 were identified. Finally, in order to demonstrate the ability  
 267 of the primers used in the aligned DNA library to discriminate each species by a unique  
 268 barcode, groups of taxa sharing the same haplotype were identified.

269

270 **Results and discussion**

271 **Taxonomic composition of the French metropolitan freshwater**  
 272 **macroinvertebrates species list and barcoding coverage**

273 The French metropolitan freshwater macroinvertebrates species list is composed by  
 274 10 phyla, 16 classes, 50 orders, 222 families, 670 genera, and 2841 species (Table 1).

Phylum	Number of				
	Class	Order	Family	Genus	Species
Annelida	1	2	6	18	40
Arthropoda	4	20	135	508	2469
Bryozoa	2	2	6	7	11
Cnidaria	1	2	3	3	7
Entoprocta	1	1	1	1	1
Mollusca	2	11	24	52	195
Nematoda	2	9	42	68	99
Nemertea	1	1	1	1	1
Platyhelminthes	1	1	3	7	12
Porifera	1	1	1	5	6

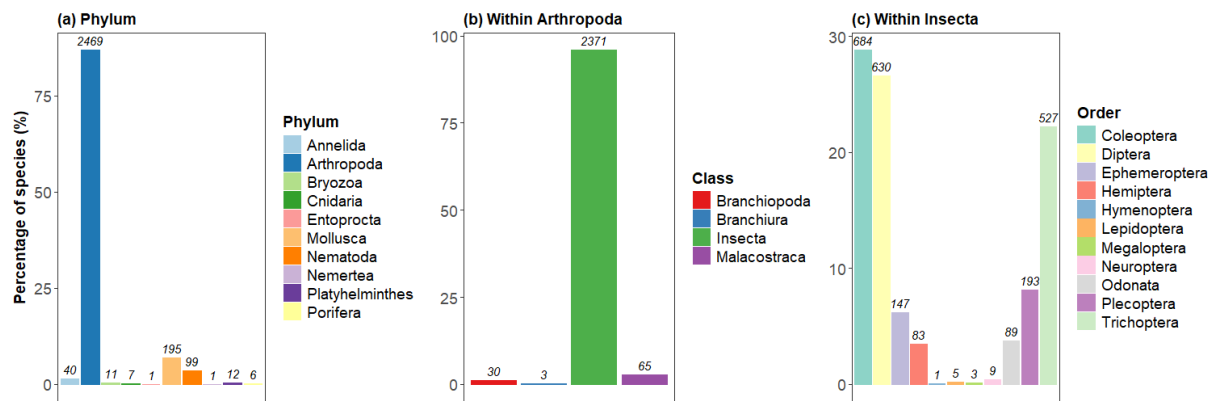
275 **Table 1.** Summary of taxonomic composition for each phylum of the the French  
 276 metropolitan freshwater macroinvertebrates species list.

277 Figure 1 provides insights into the taxonomic composition, highlighting that the highest  
 278 diversity is observed within the phylum Arthropoda, followed by Mollusca and  
 279 Nematoda, with 2469, 195, and 99 species, respectively. On the other hand, taxa such  
 280 as Entoprocta and Nemertea are represented by only one species (Figure 1, Table 1).

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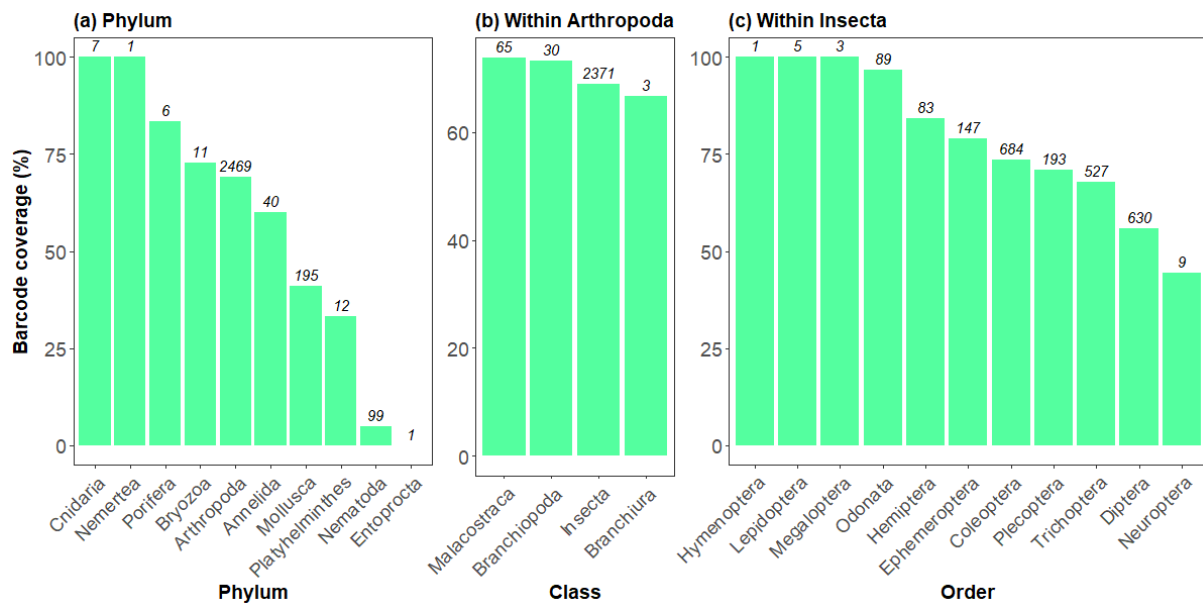
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283 **Figure 1.** Taxonomic composition of the French metropolitan freshwater  
 284 macroinvertebrates species list. Percentage of the number of species according to  
 285 phylum (a), class within the arthropod phylum (b) and order within the insect class (c).  
 286 The number above each bar represents the total number of species affiliated to each  
 287 taxonomic rank.

288 Among the insects, Coleoptera dominate, followed by Diptera and Trichoptera, with  
 289 684, 630, and 527 species, respectively. The predominant phyla with the highest  
 290 number of species in our checklist matches with those identified in a study by Specchia  
 291 et al. 2020, who conducted a gap analysis of DNA barcodes available in international  
 292 repositories using the aquatic macroinvertebrate species checklist of a south-eastern  
 293 Apulia region in Italy. Furthermore, our findings align with the prevailing understanding  
 294 that Arthropoda, and thus insects, represent the most diverse group of animals,  
 295 exerting dominance in freshwater ecosystems inventories (Choudhary & Ahi, 2015;  
 296 Dijkstra et al. 2014; Grosberg et al. 2012; Yeates & Wiegmann, 1999). Moreover,  
 297 Diptera emerge as the most species-rich group utilized in biomonitoring across Europe,  
 298 with chironomids recognized for their prevalence and diversity in freshwater habitats  
 299 (Pinder 1986). Alongside Diptera, Coleoptera (beetles) stand out for their exceptionally  
 300 high species numbers across various ecoregions and countries in Europe and serving  
 301 as the most abundant group of aquatic insects (Jäch and Balke 2008, Short 2018).  
 302 Following these orders, Trichoptera (caddisflies), Plecoptera (stoneflies) and  
 303 Ephemeroptera (mayflies), collectively known as EPT, emerge as the next three orders  
 304 in terms of species richness from our checklist. These organisms spend their immature  
 305 stages in freshwater and are widely employed as biological indicators for freshwater  
 306 quality assessment (Hering et al. 2004, Sweeney et al. 2011) and ecological  
 307 investigations, demonstrating robust responses to pollution or climate change  
 308 (Álvarez-Troncoso et al. 2015). Additionally, Nematoda emerges as a highly diverse  
 309 group, ranking third in species richness among all phyla listed in our checklist. This  
 310 outcome may be attributed to the the interest in this phylum is its utility in ecological  
 311 assessments, as nematodes are used for this purpose since a long time (e.g. Bongers,  
 312 1990; Moreno et al. 2011). From all the sequences retrieved based on the species of  
 313 the French metropolitan freshwater macroinvertebrates species list, 85.4% were  
 314 belonging to COI-5P gene (Figure S1). This result was expected as BOLD is the main  
 315 repository for COI sequences (Ratnasingham and Hebert 2007). Overall, 56% of the  
 316 2841 species listed in the French metropolitan freshwater macroinvertebrates species  
 317 list possessed at least one COI-5P genetic sequences publicly available in BOLD  
 318 database (Figure 2).



319 **Figure 2.** Barcoding coverage of the French metropolitan freshwater  
 320 macroinvertebrates species list. The barcoding coverage gives for each phylum, for  
 321 each class within Arthropoda and for each order within Insecta the total percentage of  
 322 species with available COI-5P public sequence in BOLD. The number above each bar  
 323 represents the total number of species listed in the Freshwater macroinvertebrate  
 324 checklist.

325

326 There are some exceptions with Nemerta, which have a barcode coverage of 100%  
 327 and the Entoprocta which have 0% coverage (both phylum with only species  
 328 referenced). Mollusca (195 species) and Nematoda (99 species), which are the second  
 329 and third most diverse phylum in terms of species in the French metropolitan  
 330 freshwater macroinvertebrates species list, have one the worst barcode coverage of  
 331 all phyla (41% and 5.1%, respectively). Within Arthropoda, most of the class have a  
 332 barcode coverage above 60%, with the highest being the Malacostraca (65 species)  
 333 (73.8%) and the lowest the Branchiura (3 species) (66.7%). Finally, within insects,  
 334 Hymenoptera (1 specie), Lepidoptera (5 species), Megaloptera (3 species), Odonata  
 335 (89 species) and Hemiptera (83 species) have the highest barcode coverage (>80%).  
 336 The three most species diverse insects orders (Coleoptera (684 species), Diptera (630  
 337 species), Trichoptera (527 species)) have a barcode coverage of 73.4%, 55.9% and  
 338 67.9%, respectively (Figure 2). Compared to previous gap analyses carried out in other  
 339 countries, our analysis in freshwater ecosystems in France revealed a slightly lower  
 340 barcoding coverage for freshwater macroinvertebrates (56%). For instance,  
 341 investigations in specific regions such as the Apulia Region of Southeast Italy reported  
 342 DNA barcode availability for 58% of listed aquatic Macroinvertebrate species  
 343 (Specchia et al. 2020), while a study in Atlantic Iberia documented coverage for 63%  
 344 of macroinvertebrates (Leite et al. 2020). Similarly, a comprehensive assessment of  
 345 4502 freshwater invertebrate species utilized in ecological quality assessments  
 346 indicated that 60% possessed one or more barcodes (Weigand et al. 2019). Our  
 347 findings of the lowest barcode coverage at the phylum level align with previous studies,  
 348 which also reported very low barcode coverage for freshwater Platyhelminthes, with

349 only three species having sequences deposited in examined databases (two in our  
 350 study). The low taxonomic coverage for Mollusca in our study can be attributed to the  
 351 high number of DNA barcodes deposited in GenBank rather than BOLD. As our study  
 352 exclusively utilized BOLD for sequence retrieval, this likely accounts for the observed  
 353 low barcode coverage of Mollusca, despite it being the second most listed phylum in  
 354 our checklist. Additionally, although nematodes are taxonomically diverse and  
 355 ecologically significant, they are generally overlooked in existing surveys (Weigand et  
 356 al. 2019 and references therein). Insects exhibited the highest number of available COI  
 357 sequences, consistent with previous studies highlighting the overrepresentation of  
 358 Arthropoda, particularly insects (Megléc, 2023). The barcode coverage for Insecta  
 359 was 69%, aligning with other findings such as Weigand et al. (2019), who reported that  
 360 66% of monitored insect species were barcoded, and Trebitz et al. (2015), who found  
 361 approximately 70% representation in BOLD for Great Lakes fauna. Within Insecta,  
 362 Diptera had the lowest coverage on BOLD at 55.9%, while Odonata and Hemiptera  
 363 were the best covered, with over 80% of species barcoded in each group, similar to  
 364 Weigand et al. (2019). The large number of DNA barcodes accessible within public  
 365 databases often reflects the intensity of dedicated studies and associated barcoding  
 366 projects. Certain taxonomic groups, such as Arthropoda or Mollusca receive  
 367 disproportionate attention, resulting in a heightened focus and increased deposition of  
 368 sequences within genetic databases (Briski et al. 2011, 2016, Ardura 2019).  
 369 Consequently, the absence of comprehensive reference databases may lead to false-  
 370 negative outcomes (Klymus et al. 2017), while inaccuracies within these databases  
 371 can engender false-positive identifications, as evidenced by instances of misreporting  
 372 species presence (Port et al. 2016). Furthermore, the false assignment of sequences  
 373 to closely related species may ensue when references for the true species are absent  
 374 (Schenekekar et al. 2020, Couton et al. 2022). Resolving this issue necessitates to  
 375 sequence new specimens of target taxa and their subsequent integration into  
 376 reference libraries.

377

### 378 **Gap analysis of the aligned DNA library**

379 The Figure 3 represents the taxonomic composition of the 27 species from the French  
 380 metropolitan freshwater macroinvertebrates species list which cannot be aligned with  
 381 the primer pair fwhF2/fwhR2N.

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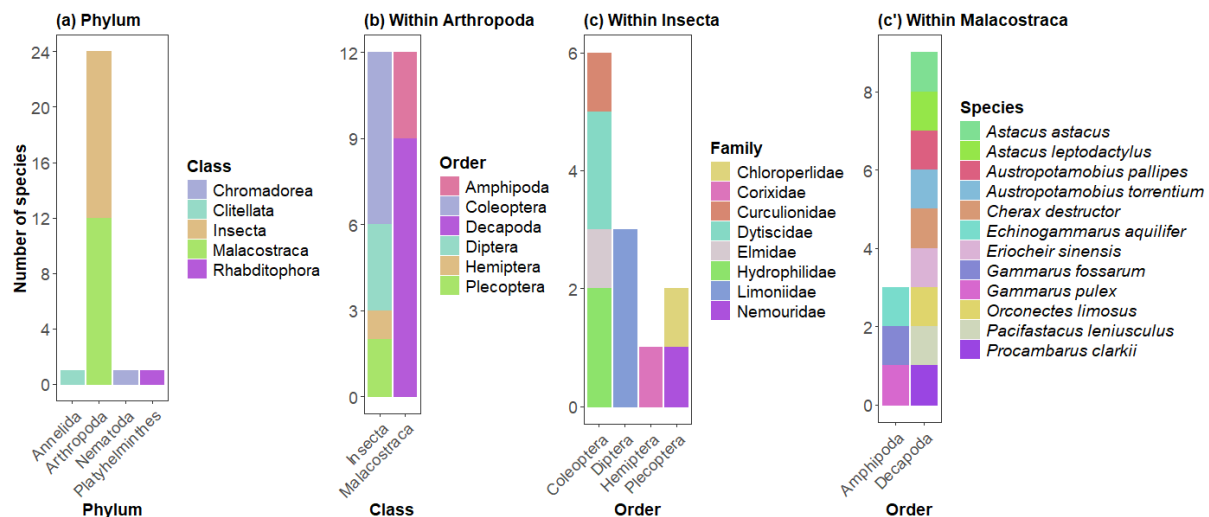
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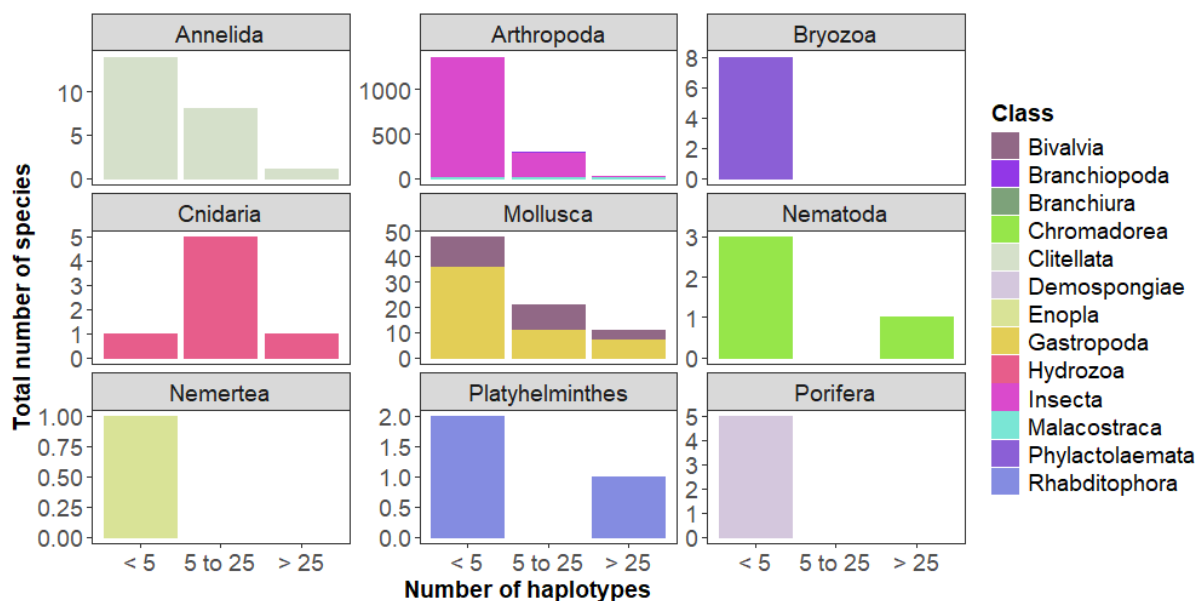
390 **Figure 3.** Gap analysis of the aligned DNA library. Number of species associated with  
 391 COI-5P genetic sequences not alignable with fwh2 primer pair at the phylum rank (a),  
 392 within the Arthropoda phylum (b) and within the Insecta and Malacostraca class (c, c').

393

394 Arthropoda emerged as the predominant taxonomic groups facing alignment difficulties  
 395 at the phylum level. Within insects (Arthropoda), Coleoptera and Diptera had the  
 396 species with the most alignment issues. Specifically, within Malacostraca (Arthropoda),  
 397 the gammarid (Gammaridae) and crayfish (Astacidea) families had the highest number  
 398 of species with alignment issues (Figure 3). Several reasons could explain these  
 399 results. Firstly, although COI-5P sequences were available, some were found to lie  
 400 outside the primer pairs' intended region during the alignment process, either entirely  
 401 or partially. In the latter case, where the sequence in question was shorter than the  
 402 desired amplified barcode, it was subsequently discarded. Another possible  
 403 explanation is the poor sequence annotation, such as mislabelling (e.g. COI-3P instead  
 404 of COI-5P) or misidentification of specimens, leading to incorrect species assignments.  
 405 Misidentification of voucher specimens has been highlighted as a major factor  
 406 contributing to erroneous records, as morphological identifications of closely related  
 407 species can be challenging. This issue has been noted by Leite et al. 2020; Paz &  
 408 Rinkevich, 2021; Pentinsaari et al. 2020, emphasizing its impact on subsequent  
 409 species identifications using databases like BOLD. This was evident in cases when  
 410 some species of insects failed to align with the primer pair, while others belonging to  
 411 the same genus did. Despite BOLD being a curated database with verification  
 412 procedures during sequence deposition and a reported error rate of less than 1% for  
 413 Metazoan sequences at the genus level (Leray et al. 2019), problematic records may  
 414 still exist, as evidenced in various studies on marine macroinvertebrates (Radulovici et  
 415 al. 2021) where up to 39% of sequences were considered ambiguous. For the  
 416 Decapoda species (Malacostraca, Arthropoda) that failed to align with the primers pair  
 417 fwhF2/fwhR2N, comprising eight crayfish species and one crab species, their absence  
 418 in the mock community during primer design could account for this discrepancy  
 419 (Elbrecht and Leese 2015, Elbrecht et al. 2017). Although Gammarids (Amphipoda,  
 420 Arthropoda) were included in primer design, the failure of some species to align with  
 421 the primers could be due to their vast diversity within aquatic environments (Horton et

422 al. 2023). Molecular studies on Amphipoda have revealed extensive species diversity  
 423 and the presence of cryptic species complexes (Jażdżewska and Mamos 2019),  
 424 suggesting a high genetic variability within species (Lefébure et al. 2006) that may not  
 425 be compatible with the primer pair. Furthermore, Vamos et al. 2017 indicated that their  
 426 analysis of the efficiency of the primers pair displayed higher penalty scores for certain  
 427 taxa of Turbellaria, Mollusca, Trichoptera, and Isopoda, indicating potential  
 428 underrepresentation due to primer bias. This aligns with findings from other studies  
 429 suggesting preferential detection of taxonomic groups by different markers and primers  
 430 (Leduc et al. 2019). Consequently, the incorporation of multiple genetic regions in DNA  
 431 metabarcoding to ensure the broadest possible taxonomic detection may prove to be  
 432 a good solution (Duarte et al. 2021). This could be particularly important to monitor the  
 433 noble crayfish, *Astacus astacus* and the amphipod *Gammarus roeselii* which cannot  
 434 be detected with fwhF2/fwhR2N primers pair, (Figure 3), whereas they are the most  
 435 frequently monitored species of the malacostracans in Europe (Weigand et al. 2019).  
 436 Analysis of the number of haplotypes relative to the number of species available within  
 437 different phyla (Figure 4) provides important information about the potential to detect  
 438 taxa in natural samples.

439



440 **Figure 4.** Taxonomic composition at the class rank for the species from the aligned  
 441 DNA library. Each box at the phylum level represents the distribution of COI-5P  
 442 haplotypes relative to the number of species.

443

444 In our reference library, for most phyla, the majority of species have less than five  
 445 haplotypes, except cnidarians, which have the majority of their species with 5 to 25  
 446 haplotypes. Only three species presented more than 100 unique barcodes: one  
 447 trichopteran (*Agraylea multipunctata*), one isopod (*Asellus aquaticus*) and one  
 448 gasteropod (*Physella acuta*). These findings are consistent with the observations of  
 449 Trebitz et al. 2015, suggesting a noteworthy exception to the prevailing low barcoding  
 450 rate for invertebrates, with some species being exceptionally well genetically

451 represented. This phenomenon may be attributed to the scientific significance and  
 452 ecological relevance of these species, such as being acknowledged as a reliable  
 453 indicator. Indeed, *Asellus aquaticus* is a common species monitored in European  
 454 countries (Weigand et al. 2019), known as bioindicator for metal pollutants detection  
 455 (e.g. O’Callaghan et al. 2019). *Physella acuta* is also intensively studied as it is an  
 456 invasive aquatic Gasteropoda with worldwide distribution (Banha et al. 2014, Vinarski  
 457 2017). The analysis on the ability of the fwhF2/fwhR2N primers to discriminate each  
 458 species by a unique sequence demonstrated that 57 identical sequences were shared  
 459 by 116 distinct species (Table S2). Among those sequences, 10 were shared by two  
 460 or three different genus and 47 were shared by two or three different species. These  
 461 results could be explained by the fact that some taxa could have been mislabelled or  
 462 that, for this length of barcode (205 pb), no genetic variability is found between two  
 463 related taxa, therefore this barcode is not suitable to decipher those species. Several  
 464 authors showed the necessity to have multiple sequences for each species to cover  
 465 correctly their haplotypic diversity (Leite et al. 2020, Keck et al. 2023). This imperative  
 466 arises from the recognition that the absence of intraspecific variants can pose  
 467 significant challenges. In instances where a single sequence is available for a species  
 468 exhibiting high genetic variability, the accurate identification of all haplotypes may be  
 469 compromised. Moreover, inadequate representation of closely related species within  
 470 reference libraries can lead to the erroneous assignment of multiple species to a single  
 471 taxonomic entity (e.g. Jackman et al. 2021), potentially resulting in erroneous  
 472 assessments of species diversity.

473

## 474 **Conclusions**

475 Our study underscores a widespread absence of reference barcodes for numerous  
 476 extant invertebrate species. Although barcoding offers advantages over morphological  
 477 identification in biomonitoring, existing gaps in barcode libraries may hinder their  
 478 effectiveness (Duarte et al. 2020, Feio et al. 2020, Hestetun et al. 2020, Vieira et al.  
 479 2021). Substantial efforts are necessary to sequence new individuals for species  
 480 absent from in the reference barcoding library, and for species with a low number of  
 481 sequences. This will enable a more efficient and reliable species identification and  
 482 biodiversity assessment, since the effectiveness and reliability of DNA barcoding  
 483 identification are linked to the thoroughness of taxonomic curation and completeness  
 484 of the reference barcoding libraries (Geiger et al. 2021) (Keck et al. 2023).  
 485 Nevertheless, our work has established a reference barcoding library for freshwater  
 486 macroinvertebrates at the French level. This database can be utilized for biodiversity  
 487 studies employing environmental DNA with short COI primers (fwhF2/fwhR2n)  
 488 designed for samples containing partially degraded DNA. Furthermore, we have  
 489 provided insights into the various biases associated with the utilization of this library,  
 490 that future users will have to take into account to interpret their results. Looking ahead,  
 491 an important future development for the fwhF2/fwhR2N reference library would be to  
 492 assign a confidence level to the identification of each taxon (e.g. species). This  
 493 confidence could be determined based on several factors, including the number of  
 494 reference sequences available (with higher sequence counts providing greater  
 495 confidence), geographical coverage (broader coverage being preferable), species

496 delimitation (with monophyletic groups offering more robust identification compared to  
497 paraphyletic ones), and the availability of metadata associated with each sequence  
498 (e.g. sampling date and location, habitat, sequence quality). These criteria align with  
499 recommendations proposed by (Fontes et al. 2021).

500

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507

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