# Analyzing Arthropod Biodiversity in Jinnah Garden, Lahore, Pakistan, Using DNA Barcoding

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Abstract:

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17 Previous difficulties in arthropod taxonomy (such as limitations in conventional morphological

18 approaches, the possibility of cryptic species and a shortage of knowledgeable taxonomists) has

19 been overcome by the powerful tool of DNA barcoding. This study presents a thorough analysis

of DNA barcoding in regards to Pakistani arthropods, which were collected from Lahore's Jinnah

Garden. The 88 percent (9,451) of the 10,792 specimens that were examined were able to

22 generate DNA barcodes and 83% (8,974) of specimens were assigned Barcode Index Numbers

(BINs). However, the success rate differed significantly between the orders of arthropods, from

77% for Thysanoptera to an astounding 93% for Diptera. Through morphological exams, DNA

25 barcoding, and cross-referencing with the Barcode of Life Data system (BOLD), the Barcode

Index Numbers (BINs) were assigned with a high degree of accuracy, both at the order (100%)

and family (98%) levels. Though, identifications at the genus (37%) and species (15%) levels

28 showed room for improvement. This underscores the ongoing need for enhancing and expanding

the DNA barcode reference library. This study identified 324 genera and 191 species, underscoring the advantages of DNA barcoding over traditional morphological identification

31 methods. Among the 17 arthropod orders identified, Coleoptera, Diptera, Hemiptera,

32 Hymenoptera, and Lepidoptera from the class Insecta dominated, collectively constituting 94%

of BINs. These results demonstrate that in Pakistani arthropods, DNA barcoding and BOLD are

an invaluable tool for improving taxonomic understanding and biodiversity assessment, opening

35 the door for further eDNA and metabarcoding research.

**Comment [WAP1]:** Title can be modified as "Assessing Arthropod Biodiversity with DNA Barcoding in Jinnah Garden, Lahore, Pakistan"

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37	Key Words: DNA Barcoding, Arthropods taxonomy, Biodiversity assessment, Barcode of Life
38	Data System (BOLD), Barcode Index Numbers (BINs)
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#### INTRODUCTION:

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42 The problems with arthropod taxonomy are not localized; they exist worldwide. When employed extensively, traditional morphological techniques—which have been utilized for many years to 43 identify species present a number of challenges. First off, it can be challenging to develop 44 consistent and trustworthy identification criteria due to the significant morphological variation 45 that some species can show within their populations (Hebert et al. 2003). Furthermore, it can be 46 47 particularly difficult to detect cryptic species that is, animals that share physical features but 48 have unique genetic characteristics—using conventional methods (William et al. 2006). Moreover, phenotypic plasticity—the ability of physical traits to alter in response to 49 environmental factors—is exhibited by many species. This might result in inconsistent 50 51 morphological identification, since members of the same species may display distinct 52 characteristics depending on their physical characteristics can change in response to environmental conditions. This can lead to inconsistencies in morphological identification, as 53 individuals within the same species may exhibit different traits based on their environmental 54 context (Moczek 2010). The lack of qualified taxonomists exacerbates these problems by 55 potentially impeding prompt and precise species identification. Examining preserved specimens 56 is often necessary for morphological identification; however, poor preservation practices, 57 specimen damage, or insufficient preservation can mask or change key morphological 58 characteristics, making identification more challenging (Cognato et al. 2020). Finally, it may 59 become more difficult to differentiate between closely related species based alone on 60 morphology due to convergent evolution, which is fueled by comparable environmental forces 61 62 and can result in the formation of similar features in various species (Montealegre et al. 2012).

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65 species identification. Researchers frequently use molecular techniques, such as DNA barcoding, in conjunction with morphological identification to overcome these challenges and 66 67 improve accuracy and reliability (Seberg et al. 2003). DNA barcoding is a widely used method in many different fields, such as: phylogenetic studies (Hajibabaei et al. 2007), 68 taxonomic analysis (Dewalt 2011), looking at biodiversity of insect communities (Hlebec et 69 70 al. 2022), examining genetic patterns (Zhou et al. 2010), phylogenetic analysis (Huang et al. 71 2016), and food authentication and safety (Dawan and Ahn 2022). Through the examination 72 of particular genetic markers, such as the cytochrome oxidase I (COI) gene, which is highly 73 conserved among species and resistant to homoplasy, DNA barcoding enables precise species 74 identification (Hajibabaei et al. 2007). Moreover, this 658-base pair sequence, often referred 75 to as the "DNA barcode," acts as a unique marker for species identification due to its 76 significant sequence variation, which helps differentiate species that are closely related (Jinbo 77 et al. 2011). DNA barcoding's rapid adoption in modern biodiversity research (Hebert et al. 2003) has been 78 79 powered by its use in specimen identification. This impressive efficacy of DNA barcoding in enabling thorough assessments of biodiversity is also demonstrated by the studies of Wilson et 80 al. 2017, Shashank et al. 2022, and D'Souza et al. 2021. DNA barcoding has significantly 81 advanced our comprehension of biological diversity by focusing on specific, consistent DNA 82 83 sequences like the internal transcribed spacer (ITS) region. This method delivers high accuracy 84 and dependability, even among species that are closely related, as evidenced by the research of 85 Tyagi et al. 2019. The study of Wilkinson et al. 2017 shows that development of nextgeneration sequencing technologies has expedited the identification and discovery of 86

This emphasizes the need for different techniques to improve the efficiency and accuracy of

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Comment [WAP9]: Better to write "This highlights the requirement for various methods to raise the effectiveness and precision of species identification. To get around these issues and increase precision and dependability, researchers commonly combine morphological identification with molecular methods like DNA barcoding."

previously unknown species, significantly increasing the speed and efficiency of DNA barcoding. DNA barcodes are unique sequences that act like biological identification tags for species. These barcodes are central to the Barcode of Life Data System (BOLD), which is an openaccess platform that simplifies the tasks of classifying species, identifying unknown specimens, and discovering new species. BOLD is an abundant resource that provides a multitude of DNA barcode records from many taxonomic categories. By making a large database of barcode sequences easily accessible, this technology expedites the process of identifying species and enables scientists and researchers to compare and evaluate genetic data from different organisms (Ratnasingham and Hebert 2007). BOLD is an essential informatics platform for biodiversity and evolutionary research, offering a user-friendly interface that simplifies the management and analysis of genetic data. BOLD integrates molecular, morphological, and distributional data, bridging gaps in bioinformatics and supporting global research collaborations. By adhering to stringent data standards, BOLD ensures the quality and reliability of genetic information, making it an invaluable resource for the scientific community (Ratnasingham and Herbert 2007). A Barcode Index Number (BIN) is a unique identification number assigned to each species. Its species' unique DNA barcode sequence serves as the basis for its BIN. The cytochrome c oxidase subunit I (COI) gene, a standardized region of the genome with notable species variation, serves as the foundation for everything. This is where what makes starts, because that special BIN can be created with just a few hundred base pairs. As to the findings of Ratnasingham and Hebert (2013) each BIN is assigned to a particular species or genus, making

the complex subject of taxonomy a little easier to navigate and comprehend.

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BINs have vast scope beyond their significance in the discovery of new species. Ren et al. 2018 demonstrated that they are useful tools for tracking the locations of current species. Moreover, BINs are also useful for estimating species abundances within large samples, as Andújar et al. 2018 and Braukmann et al. 2019 have shown. It's similar to using a microscope to enlarge on the populations of various species concealed in large samples, allowing us to learn more about the complex web of life. In addition, the use of BINs for DNA barcoding has made it easier for researchers to examine museum collections and learn more about past biological assemblages (Pentinsaari et al. 2020). Furthermore, scientists can assess the degree of similarity or dissimilarity across those populations by comparing the BIN profiles of fauna from other locations and the world at large (Ashfaq et al. 2017), which advances our knowledge of trends in global biodiversity.

Keeping in view the importance of DNA barcoding, the current study significantly expands the scope of DNA barcoding for Pakistani insects, thereby advancing our understanding of the country's taxonomic biodiversity and laying the foundation for future eDNA and metabarcoding investigations.

#### MATERIALS AND METHODS

### Sample collection and preparation

#### 1- Collections

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Specimens were collected at various sites across Jinnah garden, Lahore, Pakistan. Jinnah garden is a public park that covers an area of approximately 16 hectares (39 acres). In terms of topography, Jinnah Garden features a mix of flat and hilly areas, with several walking paths and walkways that crisscross the park.

Between 2019 and 2021, collections were conducted using various methods such as light traps, Malaise traps, sweep nets, and hand collections for larvae (Fig 1). The collected specimens were euthanized in cyanide jars, placed in paper envelopes, and then relaxed, pinned, labeled, and stored at the Entomological Laboratory within Lahore College for Women University.

## 2- Specimen identification

College for Women University, with valuable insights provided by various taxonomic experts.

To enhance accuracy, morphological identifications were cross-referenced whenever feasible by comparing the Pakistani specimens' barcode records with pre-existing records on BOLD.

The images, Collected data and specimen details were submitted to BOLD and can be accessed through the dataset DS-GMPJA Malaise trap Jinnah Garden Lahore.

Morphological identifications were carried out at the Entomological Laboratory at Lahore

#### 3- DNA barcoding / Molecular analysis

A total of 10792 insects were subjected to barcoding in Jinnah Garden, Lahore, following the established protocols (deWaard et al. 2019a, 2019b). Briefly, for larger specimens, a leg was

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carefully removed using sterile forceps and transferred to a well containing 30 ml of 95% ethanol. Smaller specimens, already on plates, were prepared for analysis, with vouchers retrieved after DNA extraction (Porco et al. 2010). At the Canadian Centre for DNA Barcoding (CCDB), we followed well-established procedures for DNA extraction, PCR amplification, and sequencing. These methods were described in previous publications (Ivanova et al. 2006, Hebert et al. 2018, deWaard et al.,2019b). Depending on the specific experiment, we used either six or twelve milliliters of material for the PCR reactions, as outlined in Hebert et al. 2013.

Using an Applied Biosystems 3730XL DNA Analyzer and the BigDye Terminator Cycle Sequencing Kit (v3.1), specimens were subjected to Sanger sequencing. Afterward, CodonCode Aligner was used to assemble, align, and modify the sequences before being submitted to BOLD. All DNA extracts are stored within the DNA archive facility at Centre for Biodiversity Genomics (CBG), Guelph, Canada.

## 4- Data analysis

The final dataset consisted of 8974 records, which received BINs and taxonomy assignments according to the workflow proposed by deWaard et al. 2019b. This involved a two-step process, where first, the barcode data was uploaded onto BOLD, and then each record underwent taxonomic assignment and verification. Morphological study by taxonomic specialists was also conducted alongside the molecular analysis to enhance species delimitation. Prior studies have shown the benefit of integrating both approaches, such as those by Silva-Brandão et al. 2009 on Lepidoptera, Blagoev et al. 2013 on Araneae (Theridioidea), and Naseem et al. 2019 on thrip specimens collected from cotton fields in Pakistan. These studies demonstrate the advantages of combining molecular and morphological techniques for accurate species identification. Only

sequences that met the criteria of quality were either assigned to already existing BINs or used to 170 create new ones, following the approach outlined by Ratnasingham and Hebert 2013. 171 To delineate new Barcode Index Numbers (BINs), the protocol necessitated adherence to 172 stringent quality criteria. Eligibility for BIN classification required sequences to span at least 500 173 base pairs of the barcode region, specifically between positions 70 and 700 on the alignment of 174 BOLD contain less than 1% ambiguous bases, and be devoid of stop codons or contamination 175 indicators (Ratnasingham and Hebert 2013). Additionally, sequences of shorter length (300-495 176 base pairs) that met the quality standards—lacking ambiguous bases and stop codons—and 177 demonstrated high similarity to an established BIN were consolidated under the corresponding BIN (deWaard et al. 2019a). Comprehensive BIN data, inclusive of specimen records and their 178 179 images where available, are accessible through the BOLD interface at DS-GMPJA Malaise trap 180 Jinnah Garden Lahore. 181 Complete BIN data, including sample records and representative 182 A "BIN discordance" analysis was employed to ascertain the proper BIN assignments within BOLD. Unassigned specimens underwent scrutiny via the BOLD Identification Engine 183 184 (http://www.boldsystems.org/index.php/IDS\_OpenIdEngine). Ensuing assignments underwent 185 corroboration through the taxon ID tree to ensure accuracy. Sequences identified as 186 contamination were consequently flagged, cataloged as such on BOLD, and excised from both 187 the analysis and their associated BIN entries. 188 The "BIN discordance" report leveraged the comprehensive suite of functions within MS Excel 189 to compute summary statistics. Furthermore, indices of species richness and evenness were 190 evaluated using the "Diversity measure" function provided by BOLD.

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## RESULTS

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194 remaining 12% were excluded from additional investigation because they were either unable to 195 amplify or produced undesirable sequences (such as contamination, NUMTs, stop codons, or 196 endosymbionts) (Supp. File 1 & 2). Sequence recovery varied widely among orders with 100 or 197 more specimens, ranging from 77% for Thysanoptera to 93% for Diptera. There was significant 198 difference in the sequence recovery for the other four major orders of insects, which are the 199 Coleoptera (83%), Hymenoptera (81%), Hemiptera (80%), and Lepidoptera (72%).(Supp. Table 200 S1) 201 Among the 9,541 successfully barcoded records, 8,974 (95%) were assigned to BINs (Barcode 202 Index Numbers), leaving 477 records that did not qualify for BINs. These 477 sequences, not 203 meeting BIN criteria, underwent analysis using the stand-alone version of the RESL algorithm 204 (via the 'Cluster Sequences' function on BOLD), revealing 386 Operational Taxonomic Units 205 (OTUs), potentially representing distinct species. Of these, only 18 OTUs (encompassing 24 206 records) were free from contamination or stop codons (Supp. File 3). Each of these 18 OTUs was 207 cross-referenced with the BOLD ID Engine, revealing no matches to known BINs and 208 suggesting they are novel to BOLD, as further supported by "taxon ID tree" analysis on BOLD. (Supp. File 4) 209 210 The 8,974 barcodes successfully assigned were distributed across 1,361 BINs. Notably, 191 211 unique BINs (14%) were exclusively identified at the Jinnah Garden site, while the remaining 212 1,170 BINs (86%) were shared with other locations, both within and outside Pakistan. In terms 213 of taxonomic classification, 98% of the barcodes (9,255) belonged to the Class Insecta, followed by Class Arachnida (99 barcodes, 1%), Class Collembola (91 barcodes, 0.96%), and Class 214

DNA barcoding analysis of 10,792 specimens yielded successful results for 9,451 (88%), the

215 Malacostraca (5 barcodes, 0.05%). The Class Arachnida specimens were further categorized into 216 four orders (Araneae, Mesostigmata, Sarcoptiformes, and Trombidiformes), encompassing 17 217 families, 14 genera, and 9 species. Collembola included two orders (Entomobryomorpha and 218 Symphypleona), with the former yielding 3 families, 3 genera, and 2 species. Malacostraca 219 featured only the order Isopoda, with four barcodes across one species. 220 In the class Insecta, specimens were assigned across 10 orders, with 98% falling into 149 221 families (as detailed in Tables 1). The majority (92%) belonged to three orders: Diptera (66%), 222 Hymenoptera (16%), and Hemiptera (10%), as shown in Figure 2. Other orders like Coleoptera, 223 Lepidoptera, and Thysanoptera each had over 50 specimens, while Neuroptera, Odonata, 224 Orthoptera, and Psocodea had just fewer. 225 Among 10792 specimens, 97% of specimens (N = 10,448) were accompanied by images. Most 226 sequences (95%) received a BIN assignment, cumulating in 1,361 BINs. Over half (51%) of the 227 1,361 Barcode Index Numbers (BINs) were represented by a minimum of two or more 228 sequences, while the remaining 49% were represented by only a single specimen. Notably, the 229 proportion of these single-specimen BINs exceeded 40% in the orders of Coleoptera, Diptera, 230 Hemiptera, and Hymenoptera, with the highest occurrence in Hymenoptera (58%, N=283). 231 Additionally, a significant majority of specimens (97%, N=10,448) were documented with images. The assignment of BINs varied across different orders, with the order Araneae (Class 232 233 Arachnida) showing an 82% success rate, and the order Entomobryomorpha (Class Collembola) 234 achieving 85%. Within Class Insecta, the distribution of BIN assignments was as follows: 235 Diptera and Hymenoptera both at 79%, Coleoptera at 75%, Lepidoptera at 64%, and Hemiptera 236 at 52% (refer to Table 1). Together these five orders contributed to 94% of the BINs and 81% of

the families identified (as shown in Table 1, Figures 2 and 3).

the Shannon Index is 5.77, suggesting high species richness and evenness (Table 2, Fig 5).

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(Supp. File 5)

## DISCUSSION

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261 Pakistan's arthropod biodiversity has been estimated to contain anywhere from approximately 262 5,000 to 20,000 species, according to various sources (Ministry of Climate Change, Pakistan 263 2019, Hasnain 1998). Nevertheless, these previous estimates have been deemed insufficient by recent studies (Baig and Al-Subaiee 2009, Rana et al 2019). This research aimed to provide a 264 265 more accurate assessment by utilizing DNA barcoding and the BIN system. A thorough 266 examination was conducted on more than 10,792 specimens gathered from Jinnah Garden in 267 Lahore, Pakistan. This exhaustive endeavor aimed to establish a comprehensive DNA barcode 268 library of the region's arthropod fauna. 269 Although the success rate (88%) for recovering DNA barcodes was good, it differed significantly 270 between orders from 77% for thysanoptera to 93% for Diptera (with specimens more than 100). 271 In other research, similar variance in barcode recovery across several arthropod taxa has been 272 documented.Park et al. 2011 reported sequence recovery rates ranging from 75% to 80% in 273 Hemiptera. Ashfaq et al. (2012) reported similar trends of variation in sequence recovery rates in 274 various insect orders and in a broader scope, across Canada, DNA barcoding was performed on 275 1,500,003 animal specimens across diverse taxonomic levels, resulting in 38% of specimens 276 being assigned taxonomically at the species level, with higher proportions at the genus and 277 family levels (deWaard et al. 2019a). However, low sequence recovery rates in DNA barcoding can be attributed to various factors. 278 279 One significant factor is the failures in primer binding, which can result from genetic variation 280 within the primer binding sites or mismatches between the primers and the targeted DNA, 281 leading to reduced sequence recovery (Elbrecht et al. 2018, Wilson et al. 2017). Such primer 282 binding failures can especially affect the amplification of DNA from diverse or taxonomically

284 Furthermore, co-amplification of pseudogenes, which are non-functional DNA sequences that 285 resemble the target genes, can introduce errors and reduce the accuracy of species identification 286 (Leite 2012). The presence of endosymbionts, like Wolbachia, in the host species can interfere 287 with DNA extraction and amplification, impacting the success of barcoding (Jones et al., 2011). 288 Recent speciation events and incomplete lineage sorting can lead to genetic similarity among 289 closely related species, making it challenging to distinguish them using a single barcode marker 290 (Soria-Carrasco et al. 2014, Yasuda et al. 2015). 291 The combination of morphological examination and barcode matching on BOLD (deWaard et al. 292 2019a, 2019b) proved highly effective in assigning BINs to an order level with 100% efficacy 293 and 98% to the family level. However, just 37% of BINs were identified to the genus level, and 294 15% to the species level. This resulted in the identification of 324 genera and 191 species, 295 demonstrating improved parameterization of the barcode reference library. This was especially 296 true for the two most diverse orders (Hymenoptera: 8% and Diptera: 12%), where species 297 assignments were less than 15%. Notably, similar studies conducted in other regions have 298 achieved considerably higher assignment success rates; Canada (38%) and Germany (34%) (e.g., Geiger et al. 2016, deWaard et al. 2019a), suggesting that further optimization of the DNA 299 300 barcode reference library may be necessary to improve identification accuracy. 301 Although the reference database used in the current analysis was limited (Virgilio et al. 2010), 302 the study still managed to identify representatives from a significant number of genera and 303 species. Specifically, 324 genera and 191 species successfully identified through the use of the 304 global reference library known as BOLD. These findings highlight the advantages of using DNA 305 barcoding over traditional morphological identification methods (Marshall et al. 2009).

complex groups, contributing to underestimations of species richness (Wilson et al. 2017).

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306 Furthermore, the analysis yielded 1361 Barcode Index Numbers (BINs), indicating a high level 307 of species richness in the fauna of Jinnah Garden in Lahore, Pakistan. However, It is crucial to 308 emphasize that these estimates are derived from a restricted sample collection and limited 309 geographic coverage, so more comprehensive efforts and broader sampling would likely result in 310 even higher estimates of species richness. 311 Of the 17 arthropod orders identified in the study, five orders (Coleoptera, Diptera, Hemiptera, 312 Hymenoptera and Lepidoptera), from the class Insecta were the most abundant, collectively 313 making up a substantial portion of 94% BINs. This finding corroborates earlier investigations 314 employing both morphological (Stork 2018) and molecular techniques (Ashfaq et al. 2022; 315 Pentinsaari et al. 2020). The dominance of Diptera and Hymenoptera can be attributed to the 316 Malaise trap collection method, which preferentially captures low-flying insects, including these 317 orders (Cooksey and Barton, 1981, deWaard et al. 2019b). Comparable patterns have been 318 documented in other studies, such as those conducted in Canada where Diptera constituted 319 approximately 57% of collections (deWaard et al. 2019b). 320 With 100 or more specimens, 16 of the 362 families dominated, and the BIN diversity mirrored 321 this trend. The fact that 59 families were represented by a single BIN and 33 families by a single 322 specimen lends credence to the survey's inconsistent detection of families. It's interesting to note that of the 21 families with the highest BINs, eight were dipterans, and the greatest BIN: 323 specimen ratio was found in the family Cecidomyiidae (Fig 4). 324 325 The analysis of species richness extrapolation, utilizing the Preston log-normal species 326 distribution model, suggest that a comprehensive sampling effort of the fauna at the Jinnah 327 Garden as it could uncover a significantly higher number of Barcode Index Numbers (BINs) than 328 what has been observed to date. A similar results were documented by (deWaard et al. 2019).

The Barcode of Life Data System (BOLD) houses an extensive database, and is a dependable platform for evaluating faunal overlap through BINs, with over nine million DNA barcode records for over 760,000 animal species. The results emphasize the importance of compiling local biodiversity inventories. This study represents a significant advancement in establishing an inventory of the arthropod fauna in Lahore, Pakistan.

AD, Young MR, Zakharov EV, Hebert PDN, Wilson JJ. 2019b. Expedited assessment of 368 terrestrial arthropod diversity by coupling Malaise traps with DNA barcoding. Genome 369 62(3):85-95. https://doi.org/10.1139/gen-2018-0093. 370 371 deWaard JR, Ratnasingham S, Zakharov EV, Borisenko AV, Steinke D, Telfer AC, Perez 372 KHJ, Sones JE, Young MR, Levesque-Beaudin V, Sobel CN, Abrahamyan A, Bessonov K, Blagoev G, deWaard SL, Ho C, Ivanova NV, Layton KKS, Lu L, Manjunath R, McKeown 373 JTA, Milton MA, Miskie R, Monkhouse N, Naik S, Nikolova N, Pentinsaari M, Prosser 374 375 SWJ, Radulovici AE, Steinke C, Warne CP, Hebert PDN. 2019a. A reference library for the identification of Canadian invertebrates: 1.5 million DNA barcodes, voucher specimens, 376 and genomic samples. Scientific Data 6(1):308. https://doi.org/10.1038/s41597-019-0320-377 2. 378 379 DeWalt RE. 2011. DNA barcoding: a taxonomic point of view. Journal of the North 380 381 American Benthological Society. 30(1):174-81. https://doi.org/10.1899/10-021.1 382 383 D'Souza ML, Van der Bank M, Shongwe Z, Rattray RD, Stewart R, Van Rooyen J, Govender D, Hebert PD. 2021. Biodiversity baselines: tracking insects in Kruger National 384 256:109034. Park with DNA barcodes. Biol. 385 Conserv. https://doi.org/10.1016/j.biocon.2021.109034 386 387 388 Elbrecht V, Hebert PD, Steinke D. Slippage of degenerate primers can cause variation in amplicon length. Sci Rep. 8(1):10999. https://doi.org/10.1038/s41598-018-29364-z 389 390 Geiger M, Moriniere J, Hausmann A, Haszprunar G, Wägele W, Hebert PDN, Rulik B. 2016. Testing the global malaise trap program-how well does the current barcode reference 391 392 library identify flying insects in Germany? Biodiversity Data Journal 4:e10671 393 https://doi.org/10.3897/BDJ.4.e10671. 394 Government of Pakistan. 2000. Biodiversity action plan for Pakistan. Available at 395 https://portals.iucn.org/library/efiles/documents/2000-081.pdf.

deWaard JR, Levesque-Beaudin V, deWaard SL, Ivanova NV, McKeown JTA, Miskie R,

Naik S, Perez KHJ, Ratnasingham S, Sobel CN, Sones JE, Steinke C, Telfer AC, Young

366

- 396 Hajibabaei M, Singer GA, Hebert PD, Hickey DA. 2007. DNA barcoding: how it
- complements taxonomy, molecular phylogenetics and population genetics. TiG. 23(4):167-
- 398 172. https://doi.org/10.1016/j.tig.2007.02.001
- 399 Hajibabaei, M., Singer, G. A., Hebert, P. D., and Hickey, D. A. (2007). DNA barcoding: how
- 400 it complements taxonomy, molecular phylogenetics and population genetics. TRENDS in
- 401 Genetics, 23(4), 167-172.
- 402 Hasnain T. 1998. Implementation of convention on biological diversity in Pakistan: policy
- brief series # 2. Islamabad, Pakistan: Sustainable Development Policy Institute (SDPI).
- 404 Hausmann A, Godfray HCJ, Huemer P, Mutanen M, Rougerie R, van Nieukerken EJ,
- 405 Ratnasingham S, Hebert PDN. 2013. Genetic patterns in European geometrid moths
- revealed by the barcode index number (BIN) system. PloS one. 8:e84518
- 407 https://doi.org/10.1371/journal.pone.0084518.
- 408 Hebert PDN, Braukmann TWA, Prosser SWJ, Ratnasingham S, deWaard JR, Ivanova NV,
- Janzen DH, Hallwachs W, NaikS, Sones JE, Zakharov EV. 2018. ASequel to Sanger:
- 410 amplicon sequencing that scales. BMC Genom, 19:219. https://doi.org/10.1186/s12864-
- 411 018-4611-3.
- Hebert PDN, Cywinska A, Ball SL, deWaard JR. 2003. Biological identifications through
- 413 DNA barcodes. Proc. R. Soc. B: Biol. Sci. 270:313–321.
- 414 https://doi.org/10.1098/rspb.2002.2218.
- 415 Hlebec D, Sivec I, Podnar M, Kučinić M. 2022. DNA barcoding for biodiversity
- assessment: Croatian stoneflies (Insecta: Plecoptera). PeerJ. 2022 Apr 20;10:e13213.
- 417 <u>https://doi.org/10.7717/peerj.13213</u>
- Huang Z, Yang C, Ke D. 2016. DNA barcoding and phylogenetic relationships in Anatidae.
- 419 Mitochondrial DNA Part A. 27(2):1042-4. https://doi.org/10.3109/19401736.2014.926545
- 420 Ivanova NV, deWaard JR, Hebert PDN. 2006. An inexpensive, automation-friendly
- 421 protocol for recovering high quality DNA. Mol. Ecol. Notes. 6(4):998–1002.
- 422 https://doi.org/10.1111/j.1471-8286.2006.01428.x.
- 423 Jinbo U, Kato T, Ito M. 2011. Current progress in DNA barcoding and future implications
- 424 for entomology. Entomol. Sci. 14(2):107-24. https://doi.org/10.1111/j.1479-

426	Jinbo, U., Kato, T. and Ito, M. (2011). Current progress in DNA barcoding and future
427	implications for entomology. Entomological Science, 14(2), 107-124.
428	
429	1Jones M, Ghoorah A, Blaxter M. 2011. MOTU and taxonerator: turning DNA barcode
430	sequences into annotated operational taxonomic units. PLoS one. 6(4):e19259.
431	https://doi.org/10.1371/journal.pone.0019259
432	
433	Leite LA. 2012. Mitochondrial pseudogenes in insect DNA barcoding: differing points of
434	view on the same issue. Biota Neotrop. 12:301-8. https://doi.org/10.1590/S1676-
435	06032012000300029
436	Marshall, S., Paiero, S. and Buck, M. 2009. Point Pelee National Park species list. Available
437	at http:// www.uoguelph.ca/debu/pelee_specieslist.htm (accessed 20 June 2020).
438	Ministry of Climate Change, Pakistan. 2019. Pakistan's sixth national report to the United
439	Nations Convention on Biological Diversity. Convention on Biological Diversity, Montreal,
440	Canada. Available at https://www.cbd.int/doc/nr/nr-06/pk-nr-06-en.pdf.
441	Moczek AP. 2010. Phenotypic plasticity and diversity in insects. Philos. Trans. R. Soc.
442	Lond., B, Biol. Sci. 365(1540):593-603. https://doi.org/10.1098/rstb.2009.0263
443	Montealegre-Z F, Jonsson T, Robson-Brown KA, Postles M, Robert D. 2012 Convergent
444	evolution between insect and mammalian audition. Sci. 338(6109):968-71.
445	https://doi.org/10.1126/science.1225271
446	Naseem MT, Ashfaq M, Khan AM, Rasool A, Asif M, Hebert PD. 2019. BIN overlap
447	confirms transcontinental distribution of pest aphids (Hemiptera: Aphididae). Plos one.
448	14(12):e0220426. https://doi.org/10.1371/journal.pone.0220426
449	Park DS, Suh SJ, Hebert PD, Oh HW, Hong KJ. 2011. DNA barcodes for two scale insect
450	families, mealybugs (Hemiptera: Pseudococcidae) and armored scales (Hemiptera:
451	Diaspididae). Bull. Entom. Res., Lond. 101(4):429-34.
452	https://doi.org/10.1017/S0007485310000714

Pentinsaari M, Blagoev GA, HoggID, Levesque-Beaudin V, Perez K, Sobel CN,

425

453

8298.2011.00449.x

- 454 Vandenbrink B, Borisenko A. 2020. DNA barcoding survey of an Arctic arthropod
- 455 community: implications for future monitoring. Insects. 11(1):46.
- 456 https://doi.org/10.3390/insects11010046
- 457 Porco D, Rougerie R, Deharveng L, Hebert P. 2010. Coupling non-destructive DNA
- 458 extraction and voucher retrieval for small soft-bodied arthropods in a high-throughput
- 459 context: the example of Collembola. Mol. Ecol. Resour. 10(6):942–945.
- 460 https://doi.org/10.1111/j.1755-0998.2010.2839.x.
- 461 Rana, N, Saleem M, Majeed W, Jalal F, Ehsan N, Nargis S. 2019. Diversity of arthropods
- 462 regarding habitat specialty in agro-ecosystem of Faisalabad, Pakistan. GSC
- biol. pharm. sci. 6(2):01-08. https://doi.org/10.30574/gscbps.2019.6.2.0008
- 464 Ratnasingham S, Hebert PD. 2013 A DNA-based registry for all animal species: the
- 465 Barcode Index Number (BIN) system. PloS one.8(7):e66213.
- 466 <u>https://doi.org/10.1371/journal.pone.0066213</u>
- 467 Ratnasingham S, Hebert PD. 2017. BOLD: The Barcode of Life Data System (http://www.
- 468 barcodinglife. org). Mol. Ecol. Notes. 7(3):355-64. https://doi.org/10.1111/j.1471-
- 469 8286.2007.01678.x
- Ren J, Ashfaq M, Hu X, Ma J, Liang F, Hebert PDN, Lin L, Germain JF, Ahmed MZ. 2018.
- 471 Barcode index numbers expedite quarantine inspections and aid the interception of
- 472 nonindigenous mealybugs (Pseudococcidae). Biological Invasions 20:449-460.
- 473 https://doi.org/10.1007/s10530-017-1546-6.
- 474 Seberg O, Humphries CJ, Knapp S, Stevenson DW, Petersen G, Scharff N, Andersen NM.
- 475 2003. Shortcuts in systematics? A commentary on DNA-based taxonomy. Trends
- 476 Ecol. Evol. 18(2):63-65. https://doi.org/10.1016/S0169-5347(02)00059-9
- 477 Seberg, O., Humphries, C. J., Knapp, S., Stevenson, D. W., Petersen, G., Scharff, N., and
- 478 Andersen, N. M. (2003). Shortcuts in systematics? A commentary on DNA-based
- taxonomy. Trends in Ecology & Evolution, 18(2), 63-65.
- 480 Shashank PR, Naveena NL, Rajgopal NN, Elliott TA, Sreedevi K, Sunil S, Meshram NM.
- 481 DNA barcoding of insects from India: Current status and future perspectives. Mol. Biol.

- 482 Rep. 49(11):10617-10626. https://doi.org/10.1007/s11033-022-07628-2
- 483 Shashank, P. R., Naveena, N. L., Rajgopal, N. N., Elliott, T. A., Sreedevi, K., Sunil, S., &
- 484 Meshram, N. M. (2022). DNA barcoding of insects from India: Current status and future
- perspectives. *Molecular Biology Reports*, 49(11), 10617-10626.
- 486 Silva-Brandão KL, Lyra ML, Freitas AV. 2009. Barcoding Lepidoptera: current situation
- and perspectives on the usefulness of a contentious technique. Neotrop. entomol. 38:441-51.
- 488 https://doi.org/10.1590/S1519-566X2009000400001
- 489 Smith MA, Bertrand C, Crosby K, Eveleigh ES, Fernandez-Triana J, Fisher BL, Gibbs J,
- 490 Hajibabaei M, Hallwachs W, Hind K, Hrcek J, Huang D-W, Janda M, Janzen DH, Li Y,
- 491 Miller SE, Packer L, Quicke D, Ratnasingham S, Rodriguez J, Rougerie R, Shaw MR,
- 492 Sheffield C, Stahlhut JK, Steinke D, Whitfield J, Wood M, Zhou X. 2012. Wolbachia and
- 493 DNA barcoding insects: patterns, potential, and problems. Plos One 7:e36514
- 494 https://doi.org/10.1371/journal.pone.0036514.
- 495 Soria-Carrasco V, Gompert Z, Comeault AA,et al. 2014. Stick insect genomes reveal natural
- 496 selection's role in parallel speciation. Science. 344(6185):738-42.
- 497 https://doi.org/10.1126/science.1252136
- 498 Stork NE. 2018. How many species of insects and other terrestrial arthropods are there on
- 499 Earth?. Annu. Rev. Entomol. 63:31-45. https://doi.org/10.1146/annurev-ento-020117-
- 500 043348

- 501 Virgilio M, Backeljau T, Nevado B, De Meyer M. 2010. Comparative performances of
- 502 DNA barcoding across insect orders. BMC bioinform. 11(1):1-10.
- 503 https://doi.org/10.1186/1471-2105-11-206
- Wilkinson MJ, Szabo C, Ford CS, Yarom Y, Croxford AE, Camp A, Gooding P. 2017.
- 505 Replacing Sanger with Next Generation Sequencing to improve coverage and quality of
- reference DNA barcodes for plants. Sci Rep. 7(1):46040. https://doi.org/10.1038/srep46040
- Williams HC, Ormerod SJ, Bruford MW. 2006. Molecular systematics and phylogeography
- of the cryptic species complex Baetis rhodani (Ephemeroptera, Baetidae)
- 510 Mol. Phylogenet. Evol. 40(2):370-382. https://doi.org/10.1016/j.ympev.2006.03.004

511	Wilson JJ, Sing KW, Floyd RM, Hebert PD. 2017. DNA barcodes and insect biodiversity.
512	J. Insect Biodivers. Syst. 575-592. <a href="https://doi.org/10.1002/9781118945568.ch17">https://doi.org/10.1002/9781118945568.ch17</a>
513	Yasuda N, Taquet C, Nagai S. Fortes M, Fan TY, Harii S, Yoshida T, Sito Y, Nadaoka K.
514	2015. Genetic diversity, paraphyly and incomplete lineage sorting of mtDNA, ITS2 and
515	microsatellite flanking region in closely related Heliopora species
516	(Octocorallia). Mol. Phylogenet. Evol. 93: 161-171.
517	https://doi.org/10.1016/j.ympev.2015.07.009
518	
519	Zhou X, Jacobus LM, DeWalt RE, Adamowicz SJ, Hebert PD. 2010. Ephemeroptera,
520	Plecoptera, and Trichoptera fauna of Churchill (Manitoba, Canada): insights into
521	biodiversity patterns from DNA barcoding. Journal of the North American Benthological
522	Society. 29(3):814-837. https://doi.org/10.1899/09-121.1