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1 Molecular phylogeny reveals distinct evolutionary lineages of the banded krait,

2 Bungarus fasciatus (Squamata, Elapidae) in Asia

3 Lal Biakzuala¹, Hmar T. Lalremsanga^{1,*}, Vishal Santra^{2,3}, Arindam Dhara², Molla T. Ahmed²,

4 Ziniya B. Mallick², Sourish Kuttalam^{2,4}, A. A. Thasun Amarasinghe^{5,*} & Anita Malhotra^{4,*}

⁵ ¹Developmental Biology and Herpetology Laboratory, Department of Zoology, Mizoram

6 University, Aizawl, Mizoram 796004, India. ²Society for Nature Conservation, Research

⁷ and Community Engagement, Nalikul, Hooghly, West Bengal 712407, India. ³Captive and

8 Field Herpetology, 13 Hirfron, Anglesey LL65 1YU, Wales, UK. ⁴Molecular Ecology and

9 Evolution at Bangor, School of Natural Sciences, College of Environmental Sciences and

10 Engineering, Environment Centre Wales, Bangor University, Bangor LL57 2UW, UK.

⁵Department of Biology, Faculty of Mathematics and Natural Sciences, Universitas

12 Indonesia, Kampus UI, Depok 16424, Indonesia. email: htlrsa@yahoo.co.in;

13 thasun.amarasinghe@ui.ac.id; a.malhotra@bangor.ac.uk

14 Abstract

15 The banded krait, *Bungarus fasciatus* is a widespread elapid snake, likely to comprise 16 several distinct species in different geographic regions of Asia. Therefore, based on 17 molecular phylogenetics and comparative morphology data, we present an overview of the 18 systematic composition of the species to delimit potential biogeographic boundaries. Our 19 phylogenetic analyses, based on four mitochondrial genes, reveal the existence of at least

20 three evolutionary lineages within *B. fasciatus*, corresponding to Indo-Myanmar, Sundaic

21 and eastern Asian lineages. We are convinced that there are at least three taxonomic

22 entities within the nomen *B. fasciatus* and restrict the distribution of *B. fasciatus* sensu

stricto to the Indo-Myanmar region. We also provide additional natural history data of the

taxon from eastern India. Finally, we advocate further studies to establish the degree of

25 reproductive isolation among these diverging evolutionary lineages and to reassess the

systematic status of this species complex especially the Sundaic and eastern Asian

27 lineages.

28 Introduction

Aside from its taxonomical importance, recognition and ascertainment of independently

30 evolving lineages is crucial for understanding the evolutionary processes affecting the origin

- of population structure and species diversification [1]. Because of the growing availability of
- 32 genetic methods for species delineation [2], numerous studies have uncovered cryptic

diversity within the widespread vertebrate species including in tropical and sub-tropical Asia; 33 for instance, among fishes [3–5], amphibians [6–8], birds [9–11], and mammals [12–14]. 34 Moreover, recent phylogeographical and molecular studies have refined our understanding of 35 cryptic speciation across biogeographic boundaries or within biogeographic regions [15,16], 36 and even propounded the suitability of reptiles in particular as biogeographic indicators 37 [17,18]. Recent studies focussing on widespread reptilian species have also established the 38 existence of previously unnoticed cryptic diversity, including in lizards [19-22] and snakes 39 23-30]. 40

Bungarus Daudin, 1803, collectively known as kraits, are venomous elapid snakes 41 42 which inhabit the Asian subcontinent [31]. Most of the nominal Bungarus species are poorly understood. However, recent study on the diversification and evolution of elapid snakes have 43 highlighted that the diversification of kraits occurred around 30-25 million years ago, and 44 45 are close relatives of other Australasian elapid genera and sea snakes [32]. Bungarus 46 fasciatus (Schneider, 1801), commonly known as the banded krait, is a nocturnal and 47 conspicuous krait that grows up to 2,250 mm in total length and is morphologically characterized by its yellow (or cream) and black banded body [33]. It occurs in various 48 49 habitat types such as primary forests, agricultural lands as well as domestic gardens up to 2,300 m above sea level [33,34]. So far, B. fasciatus has been reported from eastern India, 50 51 Nepal, Bhutan, Bangladesh, and Myanmar, extending southwards through Thailand, Malaysia and Singapore into the Indonesian archipelago, and eastwards through Laos, 52 53 Vietnam and China [35,36]. The species is currently listed as a Least Concern (LC) species 54 in the IUCN Red List [35]. Despite its wide distribution, studies have so far been conducted mainly on its potential medical significance [37], ecological importance [38,39], or 55 characterization of venom [40–45]. 56

Although there are no studies specifically on the molecular systematics of this 57 species, several previous studies have highlighted intra-specific or geographical variability 58 based on genetic barcoding [46–48]. Accurate species delimitation is crucial in view of the 59 60 variability in snake venom composition [49] and its potential effects on antivenom efficacy [50]. Most of the existing taxonomic and systematic literature on *Bungarus* have apparently 61 overlooked the intraspecific diversity of *B. fasciatus* [51–58]. Therefore, in this study we fill in 62 63 the inherent knowledge gaps by providing comparative morphological evidence and molecular phylogeny based on four mitochondrial genes (COI, CYTB, ND4 and 16S rRNA) 64 based on sequences from east and northeast India, Indochina, and the Greater Sunda islands. 65 66 Moreover, given the minimal knowledge on the natural history, reproductive behaviour, and

- ecology, which are important for assessing the population status of the species [34,59], we
- also provide natural history data for the populations of *B. fasciatus* from India.
- 69

70 Materials and methods

Sampling. For this study, we collected both morphological and genetic data for *Bungarus* 71 fasciatus, which we compared to publicly available or unpublished data. We collected 72 morphological data for the *B. fasciatus* population represented by 15 specimens from 73 74 northeastern India between the years 2007–2022. We surveyed during the day and night, collected individuals by hand, and euthanized them with MS-222 following the standard 75 procedure [60] in compliance with the American Veterinary Medical Association (AMVA) 76 guidelines and approved by the Institutional Animal Ethics Committee (IAEC) (Permission 77 No. MZU-IAEC/2018/12). We then fixed the specimens in 10% buffered formalin solution 78 overnight, prior to their storage in 70% ethanol. We preserved liver tissue samples for DNA 79 80 analysis in 95% ethanol, which were stored at -20 °C. Vouchered specimens were deposited at the Departmental Museum of Zoology, Mizoram University (MZMU). Additional blood 81 samples from the caudal sinus were collected from the West Bengal (WB) populations and 82 preserved in EDTA-Tris buffer; these specimens were subsequently released after taking 83 necessary scale counts. Our study is reported in accordance with the ARRIVE 2.0 guidelines 84 85 (Animal Research: Reporting of In Vivo Experiments) [61]. The distribution map was prepared using QGIS v3.16.2 and the digital elevation model (DEM) was downloaded from 86 87 Open Topography (https://opentopography.org/). DNA extraction, amplification and molecular analyses. Liver tissue or blood was used to 88 extract genomic DNA using DNeasy (Qiagen[™]) blood and tissue kits following the 89 manufacturer's instructions. Fragments of four mitochondrial (mt) markers (16S, COI, ND4 90 and CYTB) were amplified in a 20 µL reaction volume, containing 1X DreamTaq PCR 91 Buffer, 2.5 mM MgCl₂, 0.25 mM dNTPs, 0.2 pM of each gene primer pair, approximately 92 3.0 ng of extracted DNA, and 1 U of Taq polymerase. A negative control with reagent grade 93 water instead of DNA template was always included. Target mt gene sequences were 94 95 amplified using the thermal profiles and primers given in Supplementary Table S1. PCR products were checked using gel electrophoresis on a 1.5% agarose gel containing ethidium 96 bromide. The PCR products were cleaned using ThermoFisher ExoSAP-IT PCR product 97 cleanup reagent and subsequently sequenced using the Sanger dideoxy method using the 98 99 ABI 3730xl DNA Analyzer at Barcode BioSciences, Bangalore, India. The generated partial gene sequences were deposited on the NCBI repository (GenBank accession numbers are 100

given in Supplementary Table S2). In this study, a total of one COI, six 16S, six ND4, and 101 nine CYTB sequences were generated and were combined withpublished sequences of B. 102 fasciatus obtained from the NCBI database; database sequences of B. caeruleus, B. 103 candidus, B. ceylonicus, B. sindanus, and B. multicinctus were used as outgroups. The four 104 mt gene alignments were concatenated in SequenceMatrix [62]. Using the CYTB dataset, 105 106 the uncorrected p-distance was estimated in MEGA X using the complete deletion option for the treatment of gaps/missing data [63]. Prior to the Bayesian analysis, PartitionFinder 107 v2.1 [64] was utilized to search the best partitioning schemes and the best fitting model 108 109 through Bayesian Information Criterion (BIC) (Supplementary Table S3). Bayesian 110 phylogeny (BI) was recon-structed using the selected models in Mr.Bayes v3.2.5 [65]. The MCMC was run with four chains (one cold and three hot chains) for 20 million generations 111 and sampled every 5000 generations. Tracer v1.7 [66] was used to check the convergence of 112 likelihood and the burn-in cut-off. The diagnosis of topological convergence and MCMC 113 114 and mixing of chains was done in R-Studio [67] using the package, R We There Yet (RWTY) [68]. The BI tree was further illustrated using web-based tree annotator iTOL 115 116 software v5 [69]. The Maximum Likelihood (ML) tree was reconstructed in IQ-TREE [70] using 10,000 Ultrafast Bootstrap (UFB) [71] based on the dataset partitioned by codon 117 118 positions with the most appropriate model selected for each partition using ModelFinder [72] integrated in IQ-TREE [70]. The CYTB dataset, partitioned by codon, was utilized for 119 performing BI and ML based PoissonTree Processes (PTP) species delineation analyses [73] 120 implemented in iTaxoTools v0.1 [74]. For the input file of PTP, a non-ultrametric tree was 121 produced in IQ-TREE [70] with 10,000 UFB replicates [71] using the models selected for 122 CYTB partitions. Only the CYTB dataset was selected for the species delimitation analysis 123 as it contains more samples from different geographical regions compared to the other three 124 125 genes.

126 Morphology. We obtained morphometric (mensural and meristic) data for species

127 comparisons, and distribution data from examined specimens (Java (JV), Mizoram (MZ)

and WB) and published literature [54,75–77]. We measured the following characters to the

nearest millimetre with a Mitutoyo digital caliper and Leica M50 (Leica Microsystems Inc.)

130 dissecting microscopes: eye diameter (ED, horizontal diameter of orbit); eye–nostril length

131 (EN, distance between anteriormost point of eye and middle of nostril); snout length (ES,

distance between anteriormost point of eye and snout); head length (HL, distance between

133 posterior edge of mandible and tipof snout); head width (HW, maximum width of head);

snout-vent length (SVL, measured from tip of snout to anterior margin of vent); tail length

(TaL, measured from anterior margin of vent to tail tip). Meristic characters were taken as 135 follows: supralabials (SL) and infralabials (IL) (first labial scale to last labial scale 136 bordering gape); dorsal scale rows (DSR, counted around the body from one side of ventrals 137 to the other in three positions, on one head length behind neck, at midbody and at one head 138 length prior to cloacal plate); when counting the number of ventral scales (Ve), we scored 139 values according to the method described by Dowling [78]. We counted subcaudal scales (Sc) 140 from the first subcaudal scale meeting its opposite to the scale before the tip of the tail, the 141 terminal scute is excluded when counting. Sex of the specimens was identified by examining 142 143 everted hemipenes or by ventral tail dissection. We evaluated the relative size of the nuchal 144 band, the number of the black cross bands of each individual. The number of cross bands on the body (BB) were counted from the first band posterior to the nuchal band on the nape up 145 146 to the level of cloaca, the count on the tail from the level of cloaca to the tip of tail (BT), and number of vertebral scales covering the nuchal band (NBW). In addition, the number of 147 148 vertebral scales covering the first cross band is also considered a reliable character for adult individuals. Values for bilateral head characters are given in left/right order. We followed 149 150 Keogh [79] for hemipenial terminology, and the extent of inverted hemipenis in terms of percentage of subcaudal scales (HpR). 151

152 Statistical analyses. The morphological information was obtained from three different populations examined by us: recent and long-term preserved specimens from JV in 153 154 Indonesia (n = 15), live specimens from WB(n = 8) and live, recent and long-term preserved specimens from MZ (n = 15) states in India. Before performing any further analyses, the 155 156 meristic data were standardized to zero mean and unit standard deviation to avoid potential bias due to difference in the range of measurement among variables; for mensural data, the 157 158 combination of characters with the highest R-squared score obtained through linear 159 regression was selected as the best log transformation model to make linear relationship 160 with body size. Since we do not have gender information from the WB population, the meristics of the remaining populations (JV and MZ) were first tested using separate one-161 way analysis of variance (ANOVA) using sex and locality as factors along with Levene's 162 test [80] to test the homogeneity of variances; if the assumption of homoscedascity was 163 violated, Brown-Forsythe test [81] was utilised as an alternative approach. For mensurals 164 (TaL, HL, and HW), a two-way analysis of covariance (ANCOVA) was carried out with 165 snout-vent length (SVL) as a covariate. The meristic variables identified with no sexual 166 dimorphism were utilised for multiple comparison among the three populations by pooling 167 sexes using one- way ANOVA using locality as a factor, and post-hoc was performed with 168

applying Bonferroni correction. In addition, a potential observer difference was screened by 169 repeating measurements on the same specimens and then tested using one-way ANCOVA. 170 The variable characters among lineages identified through the univariate analyses were 171 utilized further for Principal Component Analysis (PCA) to visualize the clustering of the 172 different populations. The correlation matrices between all pairs of the morphological 173 174 variables, variance explained by each eigenvalue as well as the correlations of each variable to the first two components are explored. Specimens with missing characters were excluded in 175 the multivariate analysis. Statistical analyses were performed using the SPSS v.25.0 statistical 176 177 package (Armonk, NY: IBM Corp.).

178 **Results**

Phylogenetic relationship. The first 25% of trees from the BI analysis were discarded as 179 burn-in, and the standard deviation of split frequencies were < 0.005 when analyses 180 terminated. The graphs created using RWTY in R-Studio also indicated satisfactory 181 topological mixing. The inferred concatenated trees from BI and ML analyses were congruent 182 with each other. The BI tree, created using Mr.Bayes v3.2.5 [65] and further illustrated using 183 iTOL software v5 [69], is show in Fig. 1, with Bayesian posterior probabilities from the BI 184 analysis and UFB values from the ML analysis. The CYTB dataset consisted of a total of 185 186 1047 aligned characters, with 97 variable sites (excluding outgroups).

Molecular phylogenetic based on the concatenated aligned matrix for four 187 188 mitochondrial genes (16S, COI, ND4, and CYTB; 2850 bp in length), recovered a monophyletic clade consisting of three lineages within Asia. Both the phylogenetic analyses 189 190 and the single-locus-based PTP species delineation approach significantly support these three distinct clades which we describe as, (i) B. fasciatus from the Sundaic region, especially 191 192 from Great Sunda islands which we describe as the Sundaic lineage (Clade I; Fig. 1); (ii) B. fasciatus from Indo-Myanmar (Clade II; Fig. 1), and (iii) B. fasciatus from mainland 193 194 Sundaland including southern China, here described as east Asian lineage (Clade III; Fig. 1). The overall mean intra-specific divergence across all lineages of B. fasciatus 195 (uncorrected p-distance) was 3.5%. Furthermore, 0.4% intra-clade genetic divergence was 196 observed within Clade I (between two locations in JV), 0.0%–1.3% within Clade II (between 197 India and Myanmar), and 0.0%-6.5% within Clade III (among China, Vietnam, Thailand, and 198 an unknown locality). The mean inter-clade genetic divergence is 5.0% between Clade I 199 200 (Sundaic) and Clade II (Indo-Myanmar), 5.3% between Clade II (Indo-Myanmar) and III

- 201 (east Asia); 5.7% between Clade I (Sundaic) and III (east Asia). Combined *B. fasciatus*
- 202 (Clades I + II + III) shows the least inter- specific genetic divergence (19.5%-19.8%) with B.

- 203 *candidus*, while inter-specific distances among other species (*B. sindanus*, *B. caeruleus*, *B.*
- 204 *candidus*, *B. ceylonicus*, and *B. multicinctus*) range from 3.0% (between *B. candidus* and *B.*
- 205 *multicinctus*) to 19.0% (between these two species and *B. ceylonicus*) (also see
- 206 Supplementary Table S4).

Morphometric analysis. In this study, despite limited sampling, morphometric analyses 207 208 were performed to identify taxonomically informative characters among the examined populations (WB, MZ and JV). Only the mensurals such as TaL (p < 0.001), HW (p < 0.05) 209 210 and HL (p < 0.05) showed significantly dimorphic characters between males and females within JV and MZ populations. For meristic characters, inter-population differences were 211 212 statistically significant (p < 0.001) for Ve (MZ vs. JV), BB, BT, and NBW (the latter three characters are tested among three populations), all of which showed a higher number in the 213 MZ population; for mensural characters, inter-population differences were also statistically 214 significant for TaL (p < 0.05) and HL (p < 0.001) (Table 1). Post-hoc tests conducted among 215 the three populations for BB, BT, and NBW showed that, except for BT between MZ and 216 WB populations (p > 0.05), significant differences are seen for all characters: BB (p < 0.001217 across all the populations), NBW (p < 0.001 in MZ vs. WB, and JV vs. WB; p < 0.05 in MZ 218 vs. JV), and BT (p < 0.001 in MZ vs. JV; p < 0.01 in JV vs. WB). Comparison was also 219 made based on the identified variable meristic characters among the three populations using 220 221 a PCA. The correlation matrix showed weak correlations between pairs of variables (r < 0.7); thus, all variables were retained for this analysis. The first two components accounted 222 for 84% of the total variation of the data, with PC1, PC2 and PC3 representing 64%, 20% 223 and 11%, respectively. The loadings of all variables are high on the first axis, while only Ve 224 loads considerably highly on the second axis, with Ve having less effect on PC1 than PC2 225 226 (Supplementary Table S5). The representation of the first two components depicts substantial separation of the Javanese and the Indian populations on the first axis (PC1), and 227 marginal separation of the WB and MZ populations on the second axis (PC2) (Fig. 2). Given 228 that the samples from the three populations (WB, MZ and JV) were examined by different 229 recorders, we also tested for potential recorder bias between the East Indian and northeast 230 231 Indian specimens; however, no significant differences were seen after re-examination of the same specimens (p > 0.05). 232 **Systematics.** We present diagnostic morphological, morphometric, and meristic data taken 233

- for *Bungarus fasciatus* Clade II from east and northeast India (Supplementary Table S6).
- 254 for *Durigurus fusciarus* erade ir from east and northeast mana (Supplementary Tuble So
- The examined specimens of *B. fasciatus* from India are morphologically distinguishable
- from the Sundaic population (see Table 2). Based on the present study, we postulate the

- existence of at least three different taxonomic entities within the nomen B. fasciatus, and also 237 confirm that populations in eastern India (e.g. Odisha, WB, etc.) and northeastern India (e.g. 238 MZ, Assam, etc.) are conspecific. Based on the original description of *Pseudoboa fasciata*, 239 minimum three specimens were available or referable to Schneider [82]; hence syntypes. 240 Among these syntypes two specimens (ZMB 2771,2772) have been deposited at ZMB from 241 242 the collection of Marcus Bloch (fide Bauer [83]). In addition, one of syn-types was depicted in Russell [84] (page 3, plate 3) as the "Bungarum Pamah", an adult from "Mansoor Cottah" 243 (now Gobalpur, Odisha (Orissa), India), specimen is now lost (fide Bauer [85]). So far, the 244 245 only existing name-bearing type specimens are the two syntypes in the collection of Berlin Zoological Museum (ZMB 2771–72) originating from "Indien" (=India) fide ZMB 246 catalogue [36] a detailed taxonomic revision will be published elsewhere (Amarasinghe et 247 al. in preparation). We affirm that the specimen used by Russell [84] for his illustration is the 248 same specimen (syntype) housed in the ZMB, thus we adhere with the type locality given by 249 250 Russell [84]. Therefore, here we postulate the Indo-Myanmar populations (Clade II) as B. fasciatus sensu stricto, while considering the populations from Sundaic region, especially 251 252 from Greater Sunda Islands (Clade I) and mainland Sundaland including southern China (Clade III) as *B. fasciatus* sensu lato. Consequently, we redescribe the *B. fasciatus* sensu 253 254 stricto, including hemipenis morphology, based on MZ population, from where a large 255 number of samples are available.
- 256 Bungarus fasciatus (Schneider, 1801) sensu stricto
- 257 (Tables 1, 2; Figs. 3A–E, 4A–B, 5)
- 258 [English: Banded krait; Bengali: Sankhamuti/Sankhini/Chamorkasa; Mizo:
- 259 Chawnglei/Tiangsir]
- 260 Pseudoboa fasciata Schneider, 1801
- 261 Bungarus annularis Daudin, 1803.
- 262 Bungarus fasciatus bifasciatus Mell, 1929.
- 263 Bungarus fasciatus insularis Mell, 1930.
- 264 Examined materials. Males (*n*=7; MZMU 933, 1314, 1320, 1417, 1421, 1883, 2935) and
- Females (*n*=8; MZMU 1319, 1321, 1550, 1562, 1561, 1548, 1572, 2481) collected from
 Mizoram, northeast India.
- 267 Species redescription. Based on the overall examined MZ materials with combined sexes,
- adults SVL 444.0–1220.0 mm, tail length 47.0–133.0 mm; head elongate (HL 2.0–3.5% of
- SVL), wide (HW 71.8–92.1% of HL), slightly flattened, indistinct from neck; snout elongate

(ES 22.8–40.1% of HL), moderate, flat in dorsal view, rounded in lateral profile, rather 270 depressed. Rostral shield large, flat, slightly visible from above, pointed posteriorly; 271 interorbital width broad; internasals subtriangular; nostrils rather large, nasals large, divided, 272 and elongated, in anterior contact with rostral, and internasal and prefrontal dorsally, 1st and 273 2nd supralabial ventrally, preocular posteriorly; no loreal; prefrontal rather large, broader 274 than long, and pentagonal; frontal large, hexagonal, short, slightly longer than width; 275 276 supraoculars narrow, elongate, subrectangular, posteriorly wider; parietals large, elongate, 277 butterfly wing-like in shape, bordered by supraoculars, frontal, upper postocular anteriorly, 278 anterior and upper posterior temporals, and five or six nuchal scales posteriorly; one preocular, vertically slightly elongated, hexagonal, in contact with prefrontal and posterior 279 nasal anteriorly, supraocular dorsally, and 2nd and 3rd supralabials ventrally; eye moderate 280 (ED 10.7–21.7% of HL), round, about half of the size of snoutlength (ED 41.7–69.9% of 281 ES), pupil rounded; two postoculars, subequal or upper one larger, pentagonal, upper 282 postocular in broad contact with supraocular, parietal and anterior temporal, lower 283 postocular in contact with anterior temporal and 5th supralabials; temporals 1 + 2, large, 284 285 slightly elongated, subrectangular or pentagonal; anterior temporal larger than posterior temporal, in contact with parietal and both postoculars dorsally, and 5th and 6th 286 287 supralabial ventrally; lower posterior temporal in contact with 6th and 7th supralabials ventrally. Supralabials seven (on both sides), 5th–7th largest in size; 1st supralabial in contact 288 with rostral anteriorly, nasals dorsally, 2nd with posterior nasal and preocular dorsally, 3rd 289 with preocular and orbit dorsally, 4th with orbit; 5th with orbit, lower postocular, and 290 291 anterior temporal dorsally, and 6th with anterior and lower posterior temporals dorsally, 7th 292 with lower posterior temporal dorsally and scales of the neck posteriorly.

Mental large, triangular, blunt posteriorly; first infralabial pair larger than mental plate and in broad contact with each other, in contact with anterior chin shields posteriorly; seven infralabials, 1st-4th in contact with anterior chin shields, 4th infralabial largest in size in contact with both anterior and posterior chin shields; 4th-7th infralabials in contact with gular scales; two larger anterior chin shields, and two slightly smaller posterior chin shields; anterior chin shields in broad contact between them; posterior chin shields bordered posteriorly by seven gular scales.

Body robust, elongate and subcylindrical; dorsal scales in 15 midbody rows, all smooth and pointed posteriorly; 222–228 ventrals in males and 224–231 in females; cloacal plate divided. Tail comparatively short, TaL 8.9–10.4% of total length in males and 13.5– 17.1% of total length in males, robust and thick; subcaudals 35–37 in males and 32–36 in

304 females, divided.

Coloration. In preservative, dorsum and venter white or yellow; 22–27 black cross bands 305 306 along the body and 4 or 6 on the tail; cross bands complete laterally, and reaching the 307 ventrals except the nuchal band; the bands on the tail distinct; the nuchal band on the nape 308 anteriorly inverted V-shaped covering 15-20 vertebral scales; nuchal band starts from mid frontal; snout, anterior head, and lateral head black making remaining the white dorsal color 309 310 an inverted V-shaped marking; first black band on the body covering 6 or 7 vertebral scales; inter-band width covers with 3-5 vertebral scales; lower parts of the supralabials white; 311 312 ventral head white until the first black band; tail tip black dorsally, white ventrally.

In life (Fig. 4A), same color as in preservative, but the white body color may vary from white, cream, pale yellow to bright yellow. One juvenile with cream and black body bands was encountered in Saikhawthlir, MZ (Fig. 4B), but the snake escaped before recording morphological data.

Variation. Except the anomalous specimen (MZMU1321) which had three postoculars on
left and two on right, and temporals 1 + 2 on the left and 2 + 2 on the right, all other meristic
and morphometric characters obtained so far did not show any significant variation between
the examined populations, and also correspond to the conventional taxonomical characters
provided in previously published literature [77,86,87].

Hemipenis. Based on MZMU2935, the organ is single and subcylindrical, relatively short, 322 323 robust, and capitate; inverted hemipenis extends to 4th-7th subcaudal level (i.e. 11.1-20% from the total number of Sc); sulcus spermaticus bifurcate below the crotch, shallow and 324 325 centripetal; apical lobe less evident with only slight apical flaring; calyculate organ with a 326 complex ornamentation of retiform ridges, papillate flounces, and spines; spines on the upper 327 basal areas enlarged and decreasing the size towards the proximal portion; apical region 328 sharply separated from the basal portion by a well-defined demarcation, so the apex is free 329 and the apical part of the hemipenis is richly capitate (Fig. 5).

330

Distribution. Within India, *B. fasciatus* has been reported from Uttar Pradesh (Gorakhpur,

fide Masson [88]; also see Anwar [89] and Das et al. [90]) in the north and central

333 Maharashtra in the west [91–93], extending across Telangana (Hyderabad, fide Kinnear

[94], Andhra Pradesh [95], Chhattisgarh [96,97], Jharkhand (Koderma, fide Smith [86]; also

see Husain [98]), Bihar [99], Odisha (Mahanadi valley, fide Wall [99]; also see Boruah et al.

[100]), and northern part of WB [101] to northeastern India, including Arunachal Pradesh

337 [102,103], Assam [99,104,105], Meghalaya [106], MZ [107,108], Tripura [109], Manipur

[110] and Nagaland [111]. A few unverified records are available from Madhya Pradesh
[36], Uttarakhand [35], and southern peninsular India in Tamil Nadu, Karnataka and Kerala
[98].

Here we provide additional distributional records for B. fasciatus sensu stricto based 341 342 on 44 new localities from MZ, and two from WB, India (Supplementary Table S7). The lowest elevation among these new records is 4 m a.s.l. at Chitrasali in Hooghly District, WB 343 and the highest is 1426 m a.s.l. at Champhai Jailveng in Champhai District, MZ. Based on 344 the previous distribution of the species, the elevation range was between 40 and 2300 m a.s.l. 345 [33,34]. Moreover, an estimated distribution range of the species was plotted (Fig. 6) 346 following WHO's range estimation for *B. fasciatus* [112]. 347 348 **Natural history.** Although *B. fasciatus* is a common species, details on the ecology, habitat, population, and breeding are still sparse and further studies are needed. Therefore, here we 349 350 provide some natural history data based on two clutches of eggs encountered from two

351 localities in WB State, India:

(i) On 16th May 2020, at ca. 20:00 h, from Chitrasali village, Hooghly, the snake was 352 353 encountered on the bank of a pond adjacent to a house in the middle of a village. The female was found coiling around a clutch of 19 eggs. The breeding site was located inside a naturally 354 occurring burrow at the base of a dead tree with decayed roots. The burrow was on the bank 355 356 ca. 6 feet from the pond. The pond had a gentle slope and was surrounded by plentiful vegetation. On the day of the egg collection, the recorded ambient temperature at the natural 357 breeding site was 28–38 °C with average humidity of 78%. The eggs were relocated and 358 incubated in a dedicated herpetoculture room at 27.6 °C using 3 cm thick vermiculite 359 bedding in a perforated box. On 10th June at 20:18 h, the first egg slits were observed, and 360 hatching was completed on 18th June at 05:45 h. The fluctuating room temperature and 361 average humidity from the start of hatching until hatching was completed were 26-35 °C and 362 363 81.1%, respectively. Notably, hatchlings crawled out from the pipped eggs on the 12th, 13th, and 14th June. Upon investigation, we found that a total of six eggs failed to hatch, out of 364 which three eggs were unfertilized, two contained partially developed embryos showing 365 366 deformities, and one egg had a fully developed embryo, possibly unable to cut through the eggshell. On 18th June, we recorded the biometric data of the 13 hatchlings (5 females with 367 average SVL 322.2 mm, TaL 32.4 mm, and body weight 21.2 g; 8 males with average SVL 368 318.6 mm, TaL 36.5, body weight 19.9 g), and were subsequently released close to where 369 370 the eggs were collected.

371 (ii) On 05th May 2021 at 12:30 pm, from a construction site at Ankuni village, Hooghly.

A clutch of eight eggs were uncovered under a pile of old bricks at the base of a dead tree 372 with lots of burrows. The breeding site was located on the bank of a pond, and the entire 373 rubble pile was covered in vegetation. However, in this case, the female snake was not found 374 near the eggs, and it is possible that the excavation work might have scared the female away. 375 The eggs were relocated and incubated in the same herpetoculture room using 3 cm thick 376 377 vermiculite bedding in a perforated box. The room temperature recorded on 5th May fluctuated between 24 and 33 °C, with a relative humidity of 65%. Egg slits were seen on 6th 378 June at ca. 22:00 h. On 8th June at ca. 08:00 h, hatching was completed and all of the 379 380 juveniles had emerged from the eggs. From the egg relocation until the completed hatching (6th–8th June), the temperature and humidity fluctuated between 24 and 39 °C and 65–75%, 381 respectively. On 8th June, the biometric data of the eight hatchlings were taken (3 females 382 with average SVL 333.3 mm, TaL 38.7 mm, and body weight 21.3 g; 5 males with average 383 SVL 351.0 mm, TaL 43.2, body weight 21.4 g), and they were also released close to the site 384 385 from which the eggs had been collected.

386 **Discussion**

387 Bungarus fasciatus sensu stricto. Evidence from this study, based on morphology and molecular data, defines three distinct clades of *B. fasciatus* with non-overlapping 388 389 distribution clusters. The high genetic divergence among lineages also suggests distinct species-level groups within *B. fasciatus* as currently conceived. Our morphometric data 390 analysis also provides evidence of their morphological distinctiveness between Clade I and 391 II. Moreover, the lineage from east Asia is basal to the other two lineages but, if these clades 392 were to be accepted as full species, the name-bearing lineage is Clade II. Thus, according to 393 our newly presented evidence, and partly according to Russell [84], the distribution range of 394 Bungarus fasciatus sensu stricto (Indo-Myanmar clade) comprises east and northeast India 395 extending towards Myanmar. (Figs. 1, 6). 396

397 Systematic challenges. In this study, we elucidate the presence of three independent lineages within B. fasciatus, which is crucial for future nomenclatural revision. In the CYTB 398 gene, while negligible intra-clade genetic divergence was observed within Clade I (0.4%; 399 400 between two locations in JV) and Clade II (0.0-1.3%; Myanmar, east and northeast India), a wide range of intra-clade genetic divergence (00.0-6.5%) was evident within Clade III 401 402 (China, Vietnam, Thailand). Consequently, we speculate that there might still be cryptic diversity within the east Asian lineage (Clade III). Moreover, for robust delimitation of the B. 403 404 *fasciatus* complex, it is necessary to establish whether these lineages have undergone some

405 degree of extrinsic or intrinsic reproductive isolation to be evolving separately [113]. For

instance, due to the high evolutionary rate of hemipenial traits com- pared to the other
morphological traits [114,115], the organ has commonly been used to provide a picture of
sexual barrier even among cryptic species [116–118].

Although it has been previously stressed that delimiting the taxonomic status of 409 geographically diversified populations of venomous snakes alone cannot necessarily predict 410 patterns of venom variation, it can play a pivotal role in overcoming the consequential 411 variability of venoms [119–121]. Fry et al. [120] further indicated that the medical 412 importance of *B. fasciatus* has been overestimated. Moreover, the possible existence of 413 414 undiscovered cryptic species accompanied by more venom diversity with uncharacterized components had been pointed out [122]. Siqueira-Silva et al. [123] observed that more 415 productive environments favour more complex venom, with more toxins in similar 416 proportions. Based on the verbal autopsy we have conducted so far within MZ, there are 417 three cases of fatal envenomation potentially from the bite of banded krait. Therefore, here we 418 419 highlight the importance of analyzing the venom compositions in different populations in each biogeographically isolated clade. 420

421 **Further work.** The combination of multivariate morphometric analysis and mitochondrial

422 gene-based phylogeography has been applied successfully for species delineation

423 [24,124,125] as well as for testing species boundaries [126]. However, nuclear genes

424 provide an independent test of species boundaries [127] as they are capable of measuring the

425 extent of gene flow, and for this reason, recent work has increasingly used a combination of

426 nuclear and mitochondrial genes for phylogeographic analyses and species delineation

427 [128]. Consequently, we believe that the potentially species-level diversity across different *B*.

428 *fasciatus* populations depicted in this study cannot be overlooked, and a thorough

429 comprehension of *B. fasciatus* systematics is still a fundamental challenge.

430 Data availability

431 The generated partial gene sequences were deposited on the NCBI repository (GenBank

432 accession numbers are given in Supplementary Table S2).

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459 Author contributions

Lal Biakzuala: Conceptualization, Data curation, Formal analysis, Investigation, 460 Methodology, Software, Visualization, Writing – original draft. Hmar T. Lalremsanga: 461 Conceptualization, Data curation, Funding acquisition, Investigation, Methodology, Project 462 administration, Validation, Visualization, Writing – original draft, Resources, Supervision. 463 Vishal Santra: Investigation, Resources, Writing – review & editing. A.A. Thasun 464 465 Amarasinghe: Data curation, Formal analysis, Software, Validation, Visualization, Investigation, Writing - review & editing. Arindam Dhara: Investigation, Resources. Molla 466 T. Ahmed: Investigation, Resources. Ziniya B. Mallick: Investigation, Resources. Sourish 467 Kuttalam: Investigation, Software, Writing – review & editing. Anita Malhotra: Formal 468 analysis, Validation, Writing - review & editing. 469

470 Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personalrelationships that could have appeared to influence the work reported in this paper.

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842 Figure Legends

Fig 1. Bayesian inference (BI) phylogenetic tree based on concatenated mitochondrial 843 16S, COI, ND4 and CYTB genes; lineage partitions recovered from CYTB-based 844 PTP analyses are presented besides the BI tree (only the CYTB dataset was utilized 845 for PTP analyses because it contains more representative samples from the three 846 847 clades compared to the other genes). Values at each node represent Bayesian posterior probabilities (PP) and Ultrafast Bootstrap (UFB) values from the Maximum 848 Likelihood (ML) analysis (PP/UFB). Abbreviations of country and state/province 849 850 names are: ID: Indonesia, JW/J: Java; MM: Myanmar, AY: Ayeyarwady; IN: India, WB: West Bengal, MZ: Mizoram, AS: Assam; VN: Vietnam, VC: Vinh Phuc; CN: 851 852 China, GZ: Guizhou, GX: Guangxi, GD: Guangdong, YN: Yunnan; TH: Thailand. Fig 2. Ordination of Bungarus fasciatus populations from Mizoram, West Bengal and 853 Java along the first two principal components based on a PCA of the characters Ve, 854 BB, BT, and NBW. Total variance associated with the PC1 and PC2 are 64% and 855 20%, respectively. 856 Fig 3. Bungarus fasciatus sensu stricto (MZMU1883) from Northeast India: (A) 857 dorsal view of full body, (B) ventral view of full body, (C) dorsal view of head, (D) 858 lateral view of the left side of head, and (E) ventral view of head. 859 Fig 4. Live individuals of Bungarus fasciatus sensu stricto (A) from Keitum village, 860 Mizoram, India (MZMU1421), and (B) a juvenile with creamish dorsum coloration 861 862 from Saikhawthlir village, Mizoram, India. Fig 5. Sulcal (left) and asulcal (right) views of the right hemipenis of Bungarus 863

fasciatus sensu stricto (MZMU2935) from Mizoram, India.

Fig 6. Map showing the distribution range of *Bungarus fasciatus* sensu lato, based on

- the latest species map provided by the World Health Organization (2022); the
- 867 coloration corresponds to the three distinct evolutionary lineages recovered in the
- 868 phylogenetic analyses. The type locality of *Bungarus fasciatus* sensu stricto is
- indicated by a black star. Localities of specimens used in the morphological analyses
- are indicated by black filled diamonds (WB), circles (MZ), and triangles (JV).
- Abbreviations for countries are: IN: India, NP: Nepal, BT: Bhutan, BD: Bangladesh,
- 872 LK: Sri Lanka, CN: China, MM: Myanmar, LA: Laos, TH: Thailand, VN: Vietnam,

- 873 KH: Cambodia, MY: Malaysia, BN: Brunei Darussalam, ID: Indonesia (KA:
- 874 Kalimantan, SM: Sumatra, JW: Java).



878 Figure 1.



Figure 2.



Figure 3.



Figure 4.



894 Figure 5.



Figure 6.

Table 1. Evaluation on the meristic and mensural characters measured for 38 Bungarus fasciatus individuals from Java (JV), Mizoram (MZ), 901 and West Bengal (WB), including mean, standard deviation, minimum and maximum values. Standardized meristic data were utilised for the 902 following tests: Ve of Java and Mizoram was tested for inter-population difference and sexual dimorphism using separate one-way ANOVA 903 with locality and sex as the factors, respectively; Sc of Java and Mizoram was tested using two-way ANCOVA using sex and locality as factors; 904 BB and NBW were tested for inter-population difference (among the three populations) and sexual dimorphism (within JV and MZ) using 905 separate one-way ANOVA with locality and sex as the factors, respectively; since BT violated the assumption of homoscedascity, it was tested 906 using the alternative Brown-Forsythe test and was indicated by octothorp (#). For mensurals, two-way ANCOVA was performed for the log 907 908 transformed TaL, HL, and HW values from JV and MZ by using the log transformed SVL as a covariate, with locality and sex as the factors. The characters with statistically significant variations at the alpha level of 0.05 are shown in boldface. The characters tested for inter-population 909 difference across the three populations are indicated by asterisk (*). Significant values are in bold. 910

Characters	Sex	Java (<i>i</i>	n=15)	Mizoram	(<i>n</i> =15)	West Beng unsex	al (<i>n</i> =8) and	Sexual dimorphism		Inter-population difference	
		Mean±SD	Range	Mean±SD	Range	Mean±SD	Range				
Ve	Male	205.44±3.43	199–210	226±2.10	222–228	217.63±3.12	212-222	$F_{1,28} = 1.35$	p = 0.256	$F_{1,28} = 469.80$	p < 0.001
	Female	206.83±1.94	205-210	229.11±2.15	224–231						
Sc	Male	34.43±0.98	33–36	35.83±0.75	35–37	34.63±1.49	31–36	$F_{1,25} = 2.44$	p = 0.131	$F_{1,25} = 1.30$	p = 0.266
	Female	31.17±1.60	30–34	33.75±1.28	32–36						
BB	Male	22.67±1.12	21–25	24.33 ± 1.97	22–27	28.38±1.73	26–31	$F_{1,28} = 0.44$	p = 0.511	$F_{2,35}=39.78*$	p < 0.001*
	Female	21.83±1.17	20–23	25.00 ± 1.58	23–27						
BT	Male	3.22±0.67	2–4	5.00 ± 0.00	5	5.25 ± 1.09	4–7	$F_{1,21} = 0.12^{\#}$	$p = 0.728^{\#}$	$F_{2,12} = 17.86^{*\#}$	p < 0.001*#
	Female	3.17±0.41	3–4	4.22±0.44	4–5						
NBW	Male	19.00 ± 1.00	18–20	18.20 ± 0.45	18–19	15.63 ± 1.11	14–17	$F_{1,27} = 0.40$	p = 0.533	$F_{2,34}=22.16*$	p < 0.001*
	Female	19.00±0.63	18–20	17.67 ± 1.73	15–20						
TaL	Male	120.74 ± 20.01	90–145	101 ± 38.92	47–133	-	-	$F_{1,24} = 18.96$	p < 0.001	$F_{1,24} = 6.01$	<i>p</i> = 0.022
	Female	107.86 ± 23.43	85-145	97.88±15.56	76–119						
HL	Male	35.06±4.97	27.10-40.90	21.60 ± 5.71	12.80-26.60	-	-	$F_{1,24} = 4.37$	p = 0.047	$F_{1,24} = 79.38$	p < 0.001
	Female	34.81±6.19	25.90-44.50	21.03 ± 5.03	15.74-29.68						
HW	Male	20.88±4.03	13.80-25.70	17.79 ± 5.10	12.18-22.46	-	-	$F_{1,25} = 4.33$	p =0.048	$F_{1,25} = 0.97$	p = 0.334
	Female	20.70±3.13	16.40-26.20	16.12 ± 4.30	10.40-22.76						

Table 2. Some comparative morphological data of *Bungarus fasciatus* sensu lato in each biogeographic region, based on this study and published data.

	Population / clade					
Character	Indo-Myanmar	East Asia	Greater Sunda			
	(<i>n</i> =23)	(<i>n</i> =11)	(<i>n</i> =15)			
Ventrals	200–234	217–237	199–210			
Subcaudals	23–39	33–41	30–36			
Number of dorsal bands on body	22–31	19–21	20–25			
Number of dorsal bands on tail	4–7	?	2–4			
Nuchal band covered by vertebral scales	14–20	14–20 ?				
Background body color	Yellow / cream Yellow		Yellow / cream			
		Yang & Rao				
	Smith [75]	[76];				
Source	This study	Chen et al. [54];	This study			
	This study	Leviton et al.				
		[77]				