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Influence of geology and human activity on the genetic structure and demography of the Oriental fire-bellied toad (Bombina orientalis)*

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ABSTRACT

The Oriental fire-bellied toad (Bombina orientalis) is a commonly used study organism, but knowledge of its evolutionary history is incomplete. We analyze sequence data from four genetic markers (mtDNA genes encoding cytochrome c oxidase subunit I, cytochrome b, and 12S-16S rRNA; nuDNA gene encoding recombination activating gene 2) from 188 individuals across its range in Northeast Asia to elucidate phylogeographic patterns and to identify the historic events that shaped its evolutionary history. Although morphologically similar across its range, B. orientalis exhibits phylogeographic structure, which we infer was shaped by geologic, climatic, and anthropogenic events. Phylogenetic and divergence-dating analyses recover four genetically distinct groups of B. orientalis: Lineage 1-Shandong Province and Beijing (China); Lineage 2-Bukhan Mountain (Korea); Lineage 3-Russia, Northeast China, and northern South Korea; and Lineage 4-South Korea. Lineage 2 was previously unknown. Additionally, we discover an area of secondary contact on the Korean Peninsula, and infer a single dispersal event as the origin of the insular Jeju population. Skyline plots estimate different population histories for the four lineages: Lineages 1 and 2 experienced population decreases, Lineage 3 remained stable, while Lineage 4 experienced a sharp increase during the Holocene. The timing of the population expansion of Lineage 4 coincides with the advent of rice cultivation, which may have facilitated the increase in population size by providing additional breeding habitat.

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1. Introduction

The Oriental fire-bellied toad (Bombina orientalis) is easy to rear and to manipulate (Kawamura et al., 1972), making it a useful study organism in the sciences. Using B. orientalis, ecological research addressed the impact of maternal investment on offspring (Kaplan, 1992; Kaplan and King, 1997; Kaplan and Phillips, 2006), biochemical studies discovered useful compounds from skin secretions (e.g. bombinin- and bombesin-like peptides, bradykinins; Erspamer, 1988; Gibson et al., 2001; Chen et al., 2002), and ecotoxicological work quantified the negative effect of insecticides and

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other chemicals on frog survival and development (Kang et al., 2008; Park et al., 2010). Yet to be resolved is the evolutionary history of B. orientalis, which may provide additional insight into these previous studies.

We studied *B. orientalis* with two goals—to elucidate the genetic relationships of populations across its range and to identify the events that shaped its evolutionary history. Bombina orientalis is widespread across Northeast Asia, and a previous study showed individuals from Shandong Province and Beijing to be genetically distinct from those in Northeast China, Russia, and Korea (Zheng et al., 2009). Additionally, Beijing samples were genetically verified as being artificially introduced from Shandong populations (Zheng et al., 2009). This previous work included only one sample from the Korean Peninsula, leaving us with an incomplete picture of the evolutionary history of the species. Here we extend the findings

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of Zheng et al. (2009) by including comprehensive sampling in
 Korea.

82 Northeast Asia has undergone complex tectonic activity (Yi 83 et al., 2003; Metcalfe, 2006; Shi, 2006) and dramatic climatic change (An et al., 2001; Lambeck and Chappell, 2001; Chen et al., 84 85 2003). Pleistocene glaciation is accepted as a major factor that influenced the diversification of plants and animals in Northeast 86 87 Asia (Serizawa et al., 2002; Aizawa et al., 2007; Lee et al., 2008; Zhang et al., 2008; Bai et al., 2010; Ding et al., 2011), including B. 88 89 orientalis (Zheng et al., 2009). We evaluate this hypothesis of Pleis-90 tocene diversification of B. orientalis by performing divergence dating and demographic history analyses. 91

By sampling across its range, we uncover the phylogeographic pattern of *B. orientalis*. We identify genetically distinct groups in Korea and elucidate the origin of the Jeju Island population. Also, we estimate the diversification of *B. orientalis* to be during the Miocene, casting doubt on hypotheses of a Pleistocene glaciation effect. We offer new hypotheses on the events that shaped the evolutionary history of *B. orientalis*.

99 2. Materials and methods

100 2.1. Sampling

We collected data from 188 *B. orientalis* specimens from across its range (Fig. 1; Table S1). Of these samples, 161 were newly sequenced in this study and 27 retrieved from GenBank. Tissue samples (liver or toe clips) were preserved in 95% ethanol. *Bombina bombina* and *B. variegata* were chosen as outgroups for phylogenetic analyses based on the results of a previous study (Pabijan et al., 2013).

108 2.2. Laboratory protocols

Genomic DNA was extracted using a standard salt extractionprotocol (Sambrook and Russell, 2001). We built upon a previous

B. orientalis dataset (Zheng et al., 2009), sequencing three mito-111 chondrial DNA (mtDNA) fragments (encoding cytochrome c oxi-112 dase subunit I [COI], cytochrome b [cytb], and 12S rRNA and 16S 113 rRNA [12S-16S]) and one nuclear gene (encoding recombination 114 activating gene 2 [RAG2]). The primers COI-1/COI-2 were used to 115 amplify COI (Zheng et al., 2009), cytb-1/cytb-2 to amplify cytb 116 (Zheng et al., 2009), 12S-1/16S-2 to amplify 12S-16S (Kocher 117 et al., 1989; Palumbi, 1996), and rag2-1/rag2-4 to amplify RAG2 118 (Zheng et al., 2009). PCR products were purified and sequenced 119 in both directions using the PCR primers at the National Instru-120 ment Center for Environmental Management (Seoul National 121 University) using an ABI3730XL machine (Applied Biosystems, Fos-122 ter City, CA, USA). Sequences were assembled and edited using 123 Geneious v.5.3.6 (Biomatters, Auckland, New Zealand). Multiple 124 sequence alignments were performed using MUSCLE (Edgar, 125 2004) and manually checked. Additionally, COI, cytb, and RAG2 126 alignments were checked for erroneous stop codons by translating 127 nucleotides to amino acid. Haplotypes of RAG2 were reconstructed 128 using PHASE (Stephens et al., 2001), with the aid of SeqPhase (Flot, 129 2010). 130

2.3. Phylogenetic analyses

Phylogenetic analyses were performed on two datasets: con-132 catenated mtDNA and RAG2. All phylogenetic analyses performed 133 on the mtDNA dataset were partitioned by gene. Maximum Likeli-134 hood (ML) analyses were performed in RAxML v.8.0.2 (Stamatakis, 135 2014), while Bayesian Inference (BI) analyses were performed in 136 MrBayes v.3.2 (Ronquist et al., 2012). ML analyses were performed 137 using the combined rapid bootstrap and search for the best-scoring 138 ML tree analysis. The GTR + G model of nucleotide substitution was 139 used for the ML tree search and 1000 bootstrap replicates. For BI 140 analyses, models of nucleotide substitution were chosen within 141 MrBayes using the reversible jumping model choice (nst = mixed) 142 with both rate variation and invariable sites (rates = invgamma). 143 Two BI searches with random starting trees were run and 144

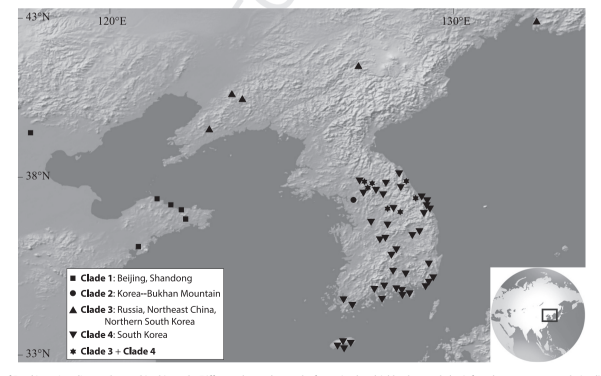


Fig. 1. Map of *Bombina orientalis* samples used in this study. Different shapes denote the four mitochondrial haplotype clades inferred to represent population lineages. The star shape indicates localities where individuals from Clade 3 and Clade 4 were found living syntopically.

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compared using four Markov Chain Monte Carlo (MCMC) chains,
10 million generations, and sampling over 1000 generations. Stationarity was evaluated by plotting the log-likelihood scores in
Tracer v.1.5 (Rambaut et al., 2013), and data from the first 2 million
generations were discarded before building a consensus tree.

For mtDNA, uncorrected *p*-distances were calculated for the major groups using MEGA v.6.06 (Tamura et al., 2013). Calculations were made with partial deletion of missing data and a site coverage cutoff of 95%.

Approximate unbiased topology tests (Shimodaira, 2002) were 154 used to examine whether the ML tree inferred from mtDNA was 155 statistically significant compared to trees with different relation-156 ships among the four major haplotype clades. Tests compared four 157 topologies: (1) the ML topology with Clade 2 (Bukhan Mountain) 158 159 being the sister taxon to a clade comprising all other samples. (2)160 Clade 2 as the sister group to Clade 1 (China only), (3) Clade 2 as the sister group to Clade 3 (Russia, China, Korea), and (4) Clade 2 161 as the sister group to Clade 4 (Korea only). We ran constrained 162 RAxML analyses and calculated the per-site log likelihoods. These 163 per-site log likelihoods were then input into the program CONSEL 164 165 (Shimodaira and Hasegawa, 2001).

166 *2.4. Divergence dating analyses*

For the mtDNA dataset, divergence dates were estimated using 167 168 BEAST v.1.8.0 (Drummond et al., 2012). An uncorrelated lognormal relaxed clock model was implemented with MCMC chains run for 169 100 million generations and sampled every 10,000 generations. A 170 GTR + I + G substitution model was used with a Yule process tree 171 172 prior. Two independent analyses were run and combined using 173 LogCombiner v.1.8. Stationarity and effective sample sizes were evaluated in Tracer v.1.5 (Rambaut et al., 2013). Tree data were 174 summarized in TreeAnnotator v.1.8.0 after removing data from 175 the first 10 million generations as burn-in. We estimated the rate 176 177 of evolution of each mtDNA marker by unlinking substitution 178 models.

179 We set priors for two calibration points based on geologic and fossil data following a previous study of *Bombing* (Pabijan et al., 180 181 2013; Table S2). Parham et al. (2012) outline the data needed for 182 justifying fossil calibrations, but as all data are not available, we use these analyses as a heuristic. The first calibration point was 183 set for the divergence between Asian (B. orientalis) and European 184 (B. bombina and B. variegata) Bombina based on range of possible 185 186 dates for the dispersal from Asia to Europe: the oldest Bombina fossil from Europe (~13 Ma [mega-annum]; Miynarski et al., 1982; 187 188 Böhme and Ilg, 2003) and the closing of the Turgai Strait 189 (~30 Ma; Sanmartin et al., 2001). For the oldest Bombina fossil 190 from Europe, Pabijan et al. (2013) tested two alternative hypotheses (13 Ma vs. 20 Ma), but we chose 13 Ma since these results of 191 192 Pabijan et al. (2013) were consistent with mid-Miocene climate 193 fragmenting Bombina species. We implemented a normal distribution with a mean of 22.5 and SD of 5.7 (13.12-33.67 Ma). The sec-194 ond calibration point was set for the divergence between B. 195 variegata and B. bombina based on the oldest B. bombina fossil from 196 Poland (~4 Ma; Sanchiz and Miynarski, 1979), B. cf. bombina from 197 Austria (~9 Ma; Tempfer, 2005; Böhme and Ilg, 2003), and B. cf. 198 199 variegata from the Czech Republic (~9 Ma; Hodrova, 1987; Böhme and Ilg, 2003). We implemented a normal distribution with 200 a mean of 7.2 and SD of 2.1 (3.74-10.65 Ma). 201

202 2.5. Estimating demographic history

The demographic history of *B. orientalis* was estimated from the mtDNA dataset using BEAST v.1.8.0 (Drummond et al., 2012). Separate analyses were run for the four groups recovered in the phylogenetic analyses. Priors for the root height of each group were set based on the results from divergence-dating analyses. A GTR + I + G207substitution model was used with a "Coalescent: Bayesian Skyline"208tree prior. MCMC chains were run for 500 million generations,209sampling every 10,000 generations. The amount of burn-in210removed varied between analyses, and was determined using211Tracer v.1.5.212

3. Results

3.1. Phylogenetic analyses

Phylogenetic trees from ML and BI analyses were similar, 215 differing only in relationships of the terminal branches and support 216

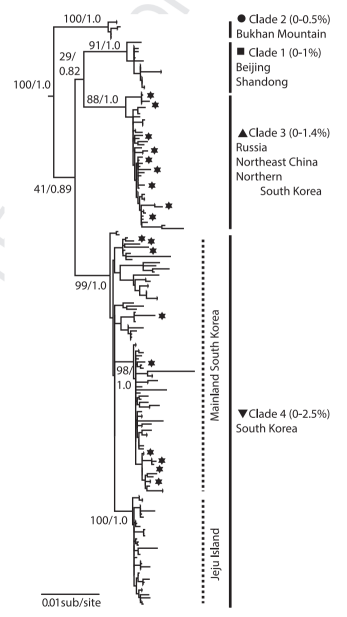


Fig. 2. Maximum likelihood (ML) phylogeny of *Bombina orientalis* inferred from the concatenated mitochondrial DNA dataset. Support values for major clades are labeled on the branches (ML bootstrap/Bayesian posterior probability). Uncorrected *p*-distance values among haplotypes within clades 1–4 are next to the clade labels. *P*-distances between clades tended to be higher (2.4–4.2%). The four major clades are denoted with the same shapes used in Fig. 1. Stars on terminal branches indicate sites where individuals from Clade 3 and Clade 4 were found living syntopically. *Bombina bombina and B. variegata* were used as outgroups for analyses (not shown).

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217 values for clades. Analyses of the mtDNA were well resolved and 218 recovered four major clades, herein designated as Clade 1, Clade 219 2, Clade 3, and Clade 4 (Fig. 2). Each clade was strongly supported 220 by ML and BI analyses (ML bootstrap = 88-100, BI posterior probability = 1.0), but relationships among clades were unclear due to 221 low support values. Clade 1 contains specimens from Shandong 222 Province and Beijing, China (Figs. 1 and 2), while Clade 2 includes 223 224 specimens from a single site in South Korea-Bukhan Mountain. Clade 3 contains samples from the Russian Far East, Northeast 225 China, and northern South Korea. Lastly, Clade 4 covers samples 226 from South Korea. Within Clade 4 is a strongly-supported clade 227 of Jeju Island samples. Seven sites in northern South Korea 228 (37.4-38°N) had individuals from both Clades 3 and 4 (denoted 229 by the star shape in Figs. 1 and 2). RAG2 had low phylogenetic sig-230 231 nal and the phylogenetic trees were poorly resolved, with many 232 polytomies present (Fig. S1). New sequences were submitted to GenBank (KR869168-KR869627: Table S1), and the dataset used 233 for phylogenetic analyses submitted to TreeBASE (http://purl.org/-234 phylo/treebase/phylows/study/TB2:S17697). 235

The maximum uncorrected *p*-distances of mtDNA within the major haplotype clades (Clade 1: 1%, Clade 2: 0.5%, Clade 3: 1.4%, Clade 4: 2.5%) were generally lower than among clades (2.4–4.2%). Uncorrected *p*-distances among haplotypes obtained from Jeju Island were <1%. The mean uncorrected *p*-distances for each mtDNA marker are listed in Table S3.

Results from the topology tests comparing alternative topologies for the placement of Clade 2 (Bukhan Mountain) were not significant (all *p*-values > 0.1). These results indicate that non-ML topologies were statistically equal to the ML tree and therefore possible.

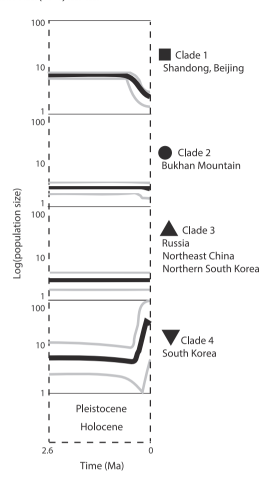


Fig. 3. Skyline plots inferring the demographic history of the four major *Bombina orientalis* lineages recovered in our study. The *y*-axis represents the log-transformed population size (effective population size \times generation time). Graphs have been truncated to only show the Pleistocene and Holocene.

3.3. Demographic history

The Bayesian skyline graphs inferred the demographic histories267of the four *B. orientalis* lineages (Fig. 3). The demographic patterns268of the four lineages differed during the Pleistocene-Holocene:269Lineage 1 and 2 showed population decreases, Lineage 3 showed270a stable population size, and Lineage 4 showed a population271expansion.272

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4. Discussion

4.1. Genetic structure of B. orientalis

Bombina orientalis occurs across Northeast Asia in Northeast 275 China, the Russian Far East, and Korea. A previous phylogeographic 276

Table 1

Divergence date estimates of the major groups recovered in this study. Ma = mega-annum.

Group	Description	Mean (95% confidence interval)
All	B. orientalis, B. bombina, B. variegata	17.17 Ma (8.56-26.15 Ma)
B. orientalis	Age of B. orientalis	14.80 Ma (6.50-23.52 Ma)
Clade 1	Beijing, Shandong	5.01 Ma (1.72-9.29 Ma)
Clade 2	Bukhan Mountain	3.12 Ma (0.61-6.58 Ma)
Clade 3	Russia, Northeast China, Northern South Korea	6.31 Ma (2.29-10.79 Ma)
Clade 4	South Korea	9.20 Ma (3.77–15.33 Ma)
Jeju	Jeju Island population	4.32 Ma (1.64-7.61 Ma)

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247 3.2. Divergence dating analyses

After removing the burn-in, the majority divergence dating 248 249 analysis parameters had effective sample sizes greater than 200, indicating the MCMC chain reached stationarity. All divergence-250 date estimates had broad confidence intervals. The same four 251 major haplotype clades were recovered as with phylogenetic anal-252 253 yses, but the relationship between these groups differed. Compar-254 ison of the divergence-date estimates to previous studies is in 255 Table S4. The estimate for the origin of B. orientalis was 14.8 Ma 256 (6.5-23.52 Ma). For the four major haplotype clades, the diver-257 gence dates varied (Table 1). The estimate for the divergence 258 among the haplotypes within the Jeju clade was 4.32 Ma (1.64-7.61 Ma) (Table 1). Groups corresponding to the four major haplo-259 type clades are considered distinct evolutionary lineages and called 260 261 Lineages 1–4 in the discussion.

Inferred rates of evolution (substitutions/site/Ma) of the three
mtDNA partitions are as follows: COI–0.00632, cytb–0.00415,
12S-16S–0.00113. Comparisons of rates to previous studies
(Evans et al., 2003; Mueller, 2006) are found in Table S5.

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study of *B. orientalis* uncovered two genetically distinct groups: (1)
Shandong Province and Beijing, China and (2) Northeast China,
Russia, and Korea (Zheng et al., 2009). In our study, we increased
sampling throughout Korea and uncovered additional genetic
structure in *B. orientalis*. Since these results are based largely on
the mtDNA dataset (low variation and genetic structure in *RAG2*),
these results should be considered preliminary.

Four major mtDNA lineages were recovered: (1) Shandong Pro-284 285 vince and Beijing (China), (2) Bukhan Mountain (South Korea), (3) Russia, Northeast China, and northern South Korea, and (4) South 286 Korea (Figs. 1 and 2). The first group from Zheng et al. (2009) in 287 288 Shandong and Beijing corresponds to Lineage 1 of our study, while their second group is divided into two genetically distinct lineages 289 in our study (Lineages 3 and 4). We discovered another genetically 290 291 distinct lineage (Lineage 2) restricted to a single region in South 292 Korea (Bukhan Mountain). These four lineages are all strongly sup-293 ported (ML bootstrap = 88–100, BI posterior probability = 1.0), but 294 based on support values (Fig. 2) and topology tests, the relation-295 ship among them is unclear. Morphologically, there were no obvious differences among the lineages. As our study provided 296 297 increased sampling and new insight in Korea, we focus our discus-298 sion on the three lineages in Korea.

299 4.2. Lineage 2–Bukhan Mountain

300 We uncovered a genetically distinct group with a distribution restricted to Bukhan Mountain (Figs. 1 and 2). Seven individuals 301 were collected across 2012 and 2013 field seasons from two local-302 ities ~3 km apart. Genetically, all samples were highly similar 303 304 (p-distance < 0.5%) and formed a strongly supported haplotype 305 clade distinct from other populations (ML bootstrap = 100, BI pos-306 terior probability = 1.0). The closest collecting locality was \sim 45 km 307 to the northwest in Myoungji Mountain (data retrieved from Gen-308 Bank); this individual was a member of Lineage 4 (Fig. 2). Based on 309 ML and BI analyses, Lineage 2 was inferred to be the sister group to 310 a clade comprising all other *B. orientalis*, but this relationship was 311 not strongly supported, and topology tests could not exclude alter-312 native positions in the phylogeny. To determine the extent of Line-313 age 2's distribution and whether there are areas of sympatry with 314 other lineages, areas around Bukhan Mountain and Seoul should be 315 surveyed further.

316 4.3. Lineages 3 and 4–Russia, China, Korea

317 Excluding Lineage 2, two major groups of B. orientalis occur 318 across the Korean Peninsula. Lineage 3 is widespread in Northeast 319 China, Russia Far East, and the Korean Peninsula, while Lineage 4 is 320 restricted to Korea (Fig. 1). The uncorrected *p*-distances between 321 individuals in these two lineages ranged between 2.4% and 3.8%. 322 At seven locations across northern South Korea, where more than 323 five samples were collected (Pyunggang Botanical Garden, Gapyeong, Chuncheon, Chiak Mountain, Taebaek Mountain, 324 Gangneung, Yeoncheon), individuals from Lineages 3 and 4 lived 325 syntopically (indicated by stars in Figs. 1 and 2). RAG2 had low 326 327 variability, and these data did not provide evidence on whether the two groups hybridized-shared alleles could be a result of 328 329 either hybridization or shared ancestral polymorphisms.

330 The genetic distinctness and geographic distributions of the two 331 lineages lead us to hypothesize that the range overlap resulted 332 from allopatry, divergence, and then secondary contact. The area 333 of sympatry spans approximately 67 km between 37.4 and 38°N. 334 However, since we do not have samples from North Korea, we do not know the northern extent of the sympatric area. Bombina orien-335 336 talis in North Korea probably belongs to Lineage 3, as individuals in 337 this lineage surround North Korea, but the range of Lineage 4 pos-338 sibly extends northward. Sequencing additional nuclear markers will help to clarify the relationship between Lineages 3 and 4,339and whether they hybridize in the region of sympatry.340

4.4. Jeju Island colonization

Lineage 4 consists of a group from Jeju Island, nested in a paraphyletic mainland South Korea group (Fig. 2). This pattern is typical of a single colonization event from continent to island (Bellemain and Ricklefs, 2008). Based on our heuristic, we estimated the age of the Jeju population to be 1.64–7.61 Ma (95% confidence interval). The sister group to the Jeju group has members spread throughout the Korean Peninsula, so we could not determine the source population.

Ieiu is a volcanic island that sits atop the 100 m-deep continental shelf of the Korean Peninsula. Jeju formed primarily through volcanic activity (Sohn and Park, 2004). The geological history of Ieiu is still under debate. Surface rock lavers of Ieiu are primarily lavas, which overlay fossiliferous, volcaniclastic sediment (Koh, 1997). The age estimates of Jeju based on different methods are similar: 0.5-1.88 Ma based on ⁴⁰Ar/³⁹Ar dating (Koh et al., 2013), 0.85-1.72 Ma based on nannofossils (Yi et al., 1998), and 0.78-1.95 Ma based on paleomagnetic and biostratigraphic data (Oh et al., 2000). Our estimate for the age of the Jeju B. orientalis population (1.64-7.61 Ma) has a broad confidence interval that overlaps geologic estimates of the age of Jeju Island in the 2 Ma range. The broad confidence interval of our estimate is likely due to the uncertainty in our calibration points, but it raises the possibility that Jeju Island is older than previously believed. Regardless of the timing, these data imply that the *B. orientalis* population on Jeju Island is the result of a single colonization event with no detectable gene flow with the Korean Peninsula, despite intervals of connectedness during lower sea levels of the Quaternary (Butenko et al., 1985; Voris, 2000).

4.5. Miocene diversification

The phylogeographic patterns and diversification of many plants and animals of Northeast Asia can be explained by glacial cycles and refugia during the Quaternary (Serizawa et al., 2002; Aizawa et al., 2007; Lee et al., 2008; Zhang et al., 2008; Bai et al., 2010; Ding et al., 2011). However, our analyses estimate that B. orientalis diverged during the Miocene (Table 1), a result that differs from previous studies that estimated a Pleistocene-Pliocene diversification (Zheng et al., 2009; Pabijan et al., 2013). As estimates for age of the tree root and divergence between B. bombina and B. variegata were consistent across studies (Table S4), we believe that past studies erroneously inferred younger estimates for B. orientalis due to sampling and analytical approaches. For sampling, we expected our dates to be older compared to previous studies because we sampled deeper lineages (Zheng et al. did not have samples from Clade 2 and Pabijan et al. did not have samples from Clades 1 and 2). Additionally for analytical approaches, Zheng et al. (2009) set a prior on the rate of evolution (0.011 substitutions/site/ Ma), artificially constraining the analyses. We compared the rates of mtDNA evolution inferred in our study to previous studies and found B. orientalis to be evolving at a slightly slower rate than other amphibians (Table S4). As there are no studies investigating mtDNA evolution rate across frog diversity, it is unclear whether B. orientalis is especially slow, or falls within the normal rate range.

Our results indicate that *Bombina orientalis* diverged in the Miocene rather than the Pleistocene. The question then becomes: what events during the Miocene caused diversification? Although the broad confidence intervals on the divergence divergence-date estimates do not allow us to identify the specific events leading toward diversification, these results help us understand the broad patterns in *B. orientalis* evolution. Climate probably was not a

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401 major factor, as reconstruction of Northeast Asian vegetation suggests stable, warm climate through the Miocene (Kong, 1996, 402 403 2000; Liu and Leopold, 1994). The flora was dominated by ther-404 mophilous, evergreen, broadleaf trees (Castanopsis, Cyclobalnaopsis, 405 Cinnamomum, Ilex, Myrica) extending around the northern Pacific 406 Rim from Northeast China eastward across the Bering land bridge 407 into North America (Kong, 1996, 2000; Liu and Leopold, 1994). Geologically, the Miocene was a time of major events in Northeast 408 409 Asia such as uplift, faulting, and volcanic activity forming the major mountain ranges along the Korean Peninsula (Wang et al., 1989; 410 Park and Son, 2005, 2008; Wan, 2012), opening of the East Sea/ 411 412 Sea of Japan (Otofuji et al., 1985, 2003), and low sea level exposing a dry Yellow Sea basin (Kim, 1987; Park, 2001; Yi et al., 2003). 413 Based on divergence-dating estimates, these events during the 414 415 Miocene coincided with and probably influenced the diversifica-416 tion of *B. orientalis*. Additional analyses clarifying the relationship 417 between the four clades and estimating the timing of diversifica-418 tion events more precisely will allow us to understand the exact 419 geological events driving divergence.

420 4.6. Demographic history and human activity

421 Estimates of demography imply different histories for each of the four lineages (Fig. 3). Lineages 1 and 2 show population 422 decreases during the Pleistocene-Holocene. These are the two lin-423 eages with relatively restricted distributions, with Lineage 1 found 424 425 in Shandong Province, China (Beijing is an introduced population; 426 Zheng et al., 2009) and Lineage 2 found in Bukhan Mountain, Korea 427 (Fig. 1). It is possible that these populations were previously widespread, but experienced a population size and distribution contrac-428 429 tion influenced by the four to five Pleistocene glacial cycles in 430 Northeast Asia (Liu, 1988).

Lineage 3, since its appearance during the late Miocene, main-431 tained a stable population size. Lastly, Lineage 4 had a stable pop-432 433 ulation for most of its history, followed by a sharp population expansion during the Holocene. We suggest that human activity 434 435 contributed to the population expansion of Lineage 4. Previous 436 studies reveal that agricultural ponds provide breeding habitats 437 for amphibians (Beebee, 1997; Knutson et al., 2004), and popula-438 tion sizes may fluctuate with the availability of these man-made 439 water bodies (Beebee, 1997; Curado et al., 2011). Individuals of B. 440 orientalis aggregate and breed in shallow water bodies in the springtime (Kuzman et al., 2004), just as paddies are flooded for 441 rice cultivation. Rice paddies provide stable water bodies in which 442 443 B. orientalis breed, and population densities in rice paddies are often much higher than in natural habitats (pers. obs.). Currently, 444 rice paddies occupy approximately 10% of Korea's total area 445 (934 km²; KOSTAT, 2015). The cultivation of rice (Oryza sativa 446 japonica) in Asia is believed to have begun 8000-13,900 years 447 ago in southern China (Zhao, 1998; Jiang and Liu, 2006), and 448 449 arrived in Korea between 4000 BC (Im, 1990; Park and Lee, 1995) 450 and 300 BC (Ahn, 2010). As the timing of population expansion is 451 similar to development of rice cultivation, we hypothesize that 452 the increased availability of rice paddies during the Holocene con-453 tributed to the population size increase of B. orientalis.

454 **5. Conclusion**

Bombina orientalis is a genetically diverse species with four distinct lineages across Northeast Asia. In the middle of the Korean Peninsula, two lineages overlap, with genetically distinct individuals living syntopically. Individuals from Jeju Island were monophyletic for mitochondrial haplotypes and nested within a paraphyletic mainland grouping, indicating that the population arose from a single colonization event. Divergence-dating analyses

imply that this colonization occurred soon after the formation of 462 Jeju Island. Contrary to previous views that Pleistocene glacial 463 cycles caused population substructure in *B. orientalis*, our analyses 464 suggest a Miocene diversification. As climate was relatively stable 465 in the Miocene, diversification was probably driven by geologic 466 events. However, the Pleistocene was an important time for B. ori-467 entalis; Pleistocene glacial cycles may have caused population 468 decreases in some lineages, while rice cultivation provided stable 469 breeding habitats for a population expansion in another lineage. 470

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Appendix A. Supplementary material

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.ympev.2015.12. 019.

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