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Cytonuclear Discordance and Historical Demography of Two Brown Frogs, Rana tagoi

2 and R. sakuraii (Amphibia: Ranidae)

4 KOSHIRO ETO and MASAFUMI MATSUI\*

- 6 Graduate School of Human and Environmental Studies, Kyoto University, Yoshida
- 7 Nihonmatsu-cho, Sakyoku, Kyoto 606-8501, Japan

- 9 \*Corresponding author. Phone: +81-75-753-6846;
- 10 FAX: +81-75-753-6846;
- 11 E-mail: fumi@zoo.zool.kyoto-u.ac.jp

#### Abstract

Prior studies of mitochondrial genomic variation reveal that the Japanese brown frog *Rana tagoi* comprises a complex of cryptic species lineages, and that *R. sakuraii* arose from within this complex. Neither species forms a monophyletic group on the mitochondrial haplotype tree, precluding a simple explanation for the evolutionary origins of *R. sakuraii*. We present a more complete sampling of mitochondrial haplotypic variation (from the *ND1* and *16S* genes) plus DNA sequence variation for five nuclear loci (from the genes encoding *NCX1*, *NFIA*, *POMC*, *SLC8A3*, and *TYR*) to resolve the evolutionary histories of these species. We test hypotheses of population assignment (STRUCTURE) and isolation-with-migration (IM) using the more slowly evolving nuclear markers. These demographic analyses of nuclear genetic variation confirm species-level distinctness and integrity of *R. sakuraii* despite its apparent polyphyly on the mitochondrial haplotype tree. Divergence-time estimates from both the mitochondrial haplotypes and nuclear genomic markers suggest that *R. sakuraii* originated approximately one million years ago, and that incomplete sorting of mitochondrial haplotype lineages best explains

27 non-monophyly of *R. sakuraii* mitochondrial haplotypes. Cytonuclear discordance elsewhere in

R. tagoi reveals a case of mitochondrial introgression between two species lineages on Honshu.

The earliest phylogenetic divergence within this species group occurred approximately four

million years ago, followed by cladogenetic events in the Pliocene and early Pleistocene

yielding 10–13 extant species lineages, including *R. sakuraii* as one of the youngest.

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Key words: species complex; incomplete lineage sorting; introgression; isolation with migration

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#### 1. Introduction

Japanese brown frogs Rana tagoi and R. sakuraii are known to show a complicated genealogical relationship (Tanaka et al., 1996; Eto et al., 2012, 2013). Rana tagoi occurs widely on the main and peripheral islands of the Japanese archipelago except for Hokkaido and the Ryukyus. While most brown frogs breed in open, still waters, R. tagoi breeds in subterranean streams where the larvae can metamorphose without feeding (Matsui and Matsui, 1990; Maeda and Matsui, 1999). These distinctive traits might be the product of adaptation to the mountainous environments of the Japanese archipelago. Conversely, R. sakuraii, occurring only on Honshu sympatric with R. tagoi, breeds under rocks in open streams, and adult frogs have several characters suitable for a lotic environment (e.g., they possess fully developed toe webs, which are less well developed in R. tagoi), although its eggs and larvae share traits with those of R. tagoi. From these facts, Matsui and Matsui (1990) postulated that R. sakuraii speciated from a R. tagoi-like ancestor when it adapted to stream environments. This hypothesis is supported by phylogenetic analyses of mitochondrial haplotypes, in which R. sakuraii is embedded in R. tagoi lineages (Tanaka et al., 1996; Eto et al., 2012). However, neither of the species is monophyletic on the mitochondrial haplotype tree (Eto et al., 2012). Mitochondrial haplotype variation reveals that R. tagoi is divided into numerous species lineages, and some of these lineages are reproductively isolated from each other (Eto et al., 2012, 2013). As is clear from

these studies, *R. tagoi* contains multiple cryptic species, one of which is the sister taxon to *R. sakuraii*. Two hypotheses potentially explain polyphyly of *R. sakuraii* haplotypes on the mitochondrial haplotype phylogeny. Incomplete lineage sorting (ILS), retention of disparate haplotype lineages from an *R. tagoi*-like ancestor, is the simplest explanation if *R. sakuraii* originated very recently, within the past approximately one million years. Alternatively, introgression of mitochondrial haplotypes resulting from gene flow between *R. sakuraii* and a sympatric lineage of *R. tagoi* could explain the anomalous phylogenetic distribution of *R. sakuraii* mitochondrial haplotypes.

In this study, we analyse sequence data for two mitochondrial and five nuclear loci to test these hypothesis and to estimate divergence times and demographic patterns of these two species. Expanded sampling of mitochondrial haplotype variation relative to earlier studies yields increased precision of the mitochondrial phylogenetic analysis. We test hypotheses of population assignment (Pritchard et al, 2000) and isolation-with-migration (IM; Hey, 2010) using the more slowly evolving nuclear markers to verify inferences made from the mitochondrial haplotype phylogeny.

# 2. Materials and Methods

70 2.1. Sampling strategy

For each species, we chose samples belonging to representative localities/mt-lineages based on previous studies (e.g., Eto et al., 2012). We analysed 107 samples of *R. tagoi* (including three samples each of the subspecies *R. t. yakushimensis* and *R. t. okiensis* from peripheral islands) and 21 of *R. sakuraii* from 81 localities (Fig. 1, Table S1). To the mtDNA phylogenetic analysis, we added GenBank data for *R. kobai* (AB685768), *R. sauteri* (AB685767), *R. tsushimensis* (AB639592, AB639752), and *R. ulma* (AB685780) as outgroup taxa based on known phylogenetic relationships (Tanaka-Ueno et al., 1996, 1998).

#### 2.2. Sequencing of DNA

Total DNA was extracted from frozen or ethanol-preserved tissues using standard phenol-chloroform extraction procedures. Then, we amplified fragments containing the target region (two mitochondrial genes, 16S ribosomal RNA [16S] and NADH dehydrogenase subunit 1 [NDI]; and five nuclear genes, sodium-calcium exchanger 1 [NCXI=SLC8AI], nuclear factor I/A [NFIA], pro-opiomelanocortin [POMC], sodium-calcium exchanger 3 [SLC8A3], and tyrosinase [TYR]) by polymerase chain reaction (PCR). The experimental conditions and PCR techniques were essentially identical to those reported previously (Eto et al., 2012). The amplified PCR products were purified by polyethylene glycol (PEG) precipitation. The cycle sequence reactions were performed out with an ABI PRISM Big Dye Terminator ver. 3.1 Cycle sequencing Kit (Applied Biosystems) and sequenced on an ABI 3130 automated sequencer. We used the primers listed in Table S2 for PCR and sequencing, and all samples/loci were sequenced in both directions.

# 2.3. Alignment of DNA, haplotype determination, and data characteristics

Sequence alignment was conducted using MUSCLE (Edgar, 2004). For heterozygous nuclear genes, we used PHASE ver. 2.1 (Stephens et al., 2001) to determine haplotypes. In this analysis, the threshold of probability was set to small values (0.5–0.6) following Garrick et al. (2010). Before analysing the historical demography, we also used IMgc (Woerner et al., 2007) to detect the largest non-recombining block of nDNA for IM analysis, because IMa2 assumes no intra-locus recombination (Hey and Nielsen, 2004). As data parameters, we calculated the summary statistics of variable sites (vs), number of haplotypes (h), haplotype diversity (Hd), and nucleotide diversity ( $\pi$ ). We also checked the neutrality of the five nuclear loci with Tajima's D (Tajima, 1989). Since none of them showed significant deviation from zero (Table S3), these loci were considered neutral markers. We conducted all of these calculations using DnaSP (Rozas et al., 2003).

2.4. Population assignment based on mtDNA

A phylogenetic analysis was conducted using the two mitochondrial genes. First, we selected the best substitution model for each gene using Kakusan4 (Tanabe, 2011) based on the Akaike information criterion (AIC). Then, phylogenetic trees based on the maximum-likelihood method (ML) and Bayesian inference (BI) were constructed using TREEFINDER ver. Mar. 2011 (Jobb, 2011) and MrBayes ver. 3.2.1 (Ronquist and Huelsenbeck, 2003), respectively. For the ML tree, we conducted non-parametric bootstrap analysis with 1000 replicates, and branches with a bootstrap value (BS) of 70% or greater were regarded as significantly supported. In the BI analysis, two independent runs of four Markov chains were conducted for 10 million generations (sampling frequency one tree per 100 generations); the first three million generations were discarded as burn-in. Convergence of parameters was checked using Tracer ver. 1.5 (Rambaut and Drummond, 2009). We considered a Bayesian posterior probability (BPP) of 0.95 or greater as significant support. From the results of both analyses, we used mitochondrial haplotype clades, levels of haplotype divergence, and geographic distributions to diagnose hypothetical species lineages, which were treated as population units based on mtDNA in the later analyses.

# 2.5. Population assignment based on nDNA

Rana tagoi and R. sakuraii are so close genetically as to cause difficulty constructing phylogenetic trees using nDNA sequences (Eto et al., 2012, 2013). Therefore, we conducted clustering analysis using STRUCTURE ver. 2.3.3 (Pritchard et al., 2000) to delimit population units based on nDNA. We applied an admixture and allele-frequency-independent model to haplotype data for the nuclear loci, and calculated 500,000 generations following 100,000 generations of burn-in. The number of clusters (K) was set from 1 to 10, and 10 independent iterations were conducted for each K. The most likely K was determined by the likelihood

131 distribution of each iteration and the delta K value (Evanno et al., 2005). We also constructed 132 haplotype networks for each gene based on the median-joining method using Network ver. 4.6 133 (Bandelt et al., 1999) to examine the relationships among nuclear haplotypes. 134 135 2.6. Divergence dating based on mtDNA 136 To estimate the divergence time between mt-lineages, we conducted Bayesian analysis 137 using BEAST ver. 1.7.5 (Drummond et al., 2012). For each calibration, 10 million generations 138 of run (of which the first three million were discarded as burn-in) were conducted under a non-139 autocorrelated log-normal relaxed clock model. Tracer ver. 1.5 (Rambaut and Drummond, 140 2009) was used to check the parameter distributions and effective sample size. We applied the 141 following two different calibrations: 142 Calibration I: The molecular evolutionary rate of 1.38% (0.69% per lineage) per MY was 143 applied. This value was estimated for the ND1 and ND2 regions of Bufo (Macey et al., 1998), 144 and only ND1 data were used in this calculation. The evolutionary rate of this region is similar 145 among a wide range of vertebrates (Macey et al., 2001). We thus used that rate, despite 146 considerable phylogenetic distance between Rana and Bufo. 147 **Calibration II**: Using only 16S data, we applied the evolutionary rate of 0.66% (0.33% per 148 lineage) per MY estimated for 16S of Leiopelma (Fouquet et al., 2009). 149 150 2.7. Estimation of historical demography 151 The historical demography, especially the patterns of gene flow and divergence times 152 among species or genetic groups, was examined using coalescent analysis with the Bayesian IM 153 model. We analysed the nDNA data using the program IMa2 (Hey 2010), and estimated the 154 effective population size,  $N_e$ , population migration rate,  $2N_eM$ , and population divergence time,

T. As the mutation rate of nuclear genes, we applied 0.047% per MY per lineage for NCX1

(reported in the genus *Hydromantes*; Rovito 2010), 0.072% (0.061–0.083%) for *POMC* 

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157 (*Hyperolius*; Lawson, 2010), and 0.047% (0.027–0.067%) for *SLC8A3* (amphibians in general; Roelants et al., 2007). The geometric mean of these values, approximately  $2.71 \times 10^{-7}$  mutations 158 159 per year per locus, was used as the mutation rate  $(\mu)$  to scale each demographic parameter. 160 Based on several test runs, the upper bounds for the parameters were set at  $\theta = 10-20$ , t = 3-5, and m = 10-25, and five million steps (sampling frequency one tree per 50 steps) of 161 162 calculations were performed for 30 heated chains after two million burn-in steps. We conducted 163 three independent runs, and finally combined the results using the L-mode option of IMa2. 164 Since R. tagoi and R. sakuraii typically start to breed at the age of 3 years (Kusano et al., 1995a, 165 b), we applied this value as the generation time of the two species. The trendline plots and effective sample sizes were monitored to ensure good mixing and convergence of parameters. 166 167 The significance of  $2N_eM$  was determined using the log-likelihood ratio (LLR) test of 168 Nielsen and Wakeley (2001). We also used the parameter comparison option (with the -p6 169 command) of IMa2 and output the list of probability, which indicates one parameter to be 170 greater than the other. The relative strength of genetic isolation was evaluated using  $2N_eM$ 171 values (strong  $[2N_eM \le 1]$ , moderate  $[1 < 2N_eM \le 5]$ , and weak  $[5 < 2N_eM \le 25]$ : Wright, 1931; 172 Waples and Gaggiotti, 2006; Reilly et al., 2012).

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#### 3. Results

3.1. Sequence characteristics

We obtained complete mitochondrial *16S* (1612bp) and *ND1* (967bp) sequences for all samples. There were 489 parsimoniously informative sites within the ingroup: 244 for *16S* and 245 for *ND1*. The other statistics are listed in Table S3.

In the sequences of the five nuclear loci for all 128 samples, only *POMC* had in-dels, and these sites were omitted from the subsequent analyses. For haplotype determination using

PHASE, all haplotypes in all samples/loci were determined successfully, except for one sample

for *POMC* and two for *TYR*, which were treated as null alleles in subsequent analyses. The

sequence length and statistics of each locus are listed in Table S3. Overall, each parameter generally indicated great genetic diversity in *R. tagoi* and *R. sakuraii*. Of the five nuclear loci, *TYR* was the most variable ( $H_d = 0.955$  and  $\pi = 0.017$  for all samples) and *NFIA* was the least variable (0.735 and 0.003, respectively).

The best substitution model selected in the ML analysis was the general time reversible

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3.2. Population assignment: Mitochondrial DNA results

190 (GTR; Tavaré, 1986) model with the optimized gamma shape parameter (G) of 0.158 and the 191 proposition of invariable sites (I) of 0.144 for 16S and the J1 (Jobb, 2011) model + G (0.543) + I 192 (0.312) for ND1. For BI, the models were GTR + G (0.082) + I (0.226) and GTR + G (0.892) + 193 I (0.226) for 16S and ND1, respectively. The constructed ML ( $-\ln L = 15500.618$ ) and BI 194 (15863.190) trees were essentially identical in topology, and only the ML tree is shown in Fig. 2. 195 We followed Eto et al. (2012) for the names of each genetic group. 196 The phylogenetic relationships obtained were fundamentally identical to those reported by 197 Eto et al. (2012). The ingroup was divided into two large haplotype clades (A and B), and both 198 of these included subclades judged by their geographic distributions to diagnose separate 199 species lineages (A-1ab to A-9abc and B-1 to B-2ab); Clade B (ML-BS = 82% and 200 BPP = 1.00) contained only haplotypes from R. tagoi, while Clade A (ML-BS = 93% and 201 BPP = 1.00) included both *R. tagoi* and *R. sakuraii* haplotypes. Each clade/lineage was well 202 supported (ML-BS≥70%, BPP≥0.95). The statistical support for nodes was generally better 203 than in the previous study, and more detailed phylogenetic relationships were clarified, 204 particularly those among the lineages in Clade A. In Clade A, the lineages from Honshu Island 205 (A-1ab to A-6) formed a subclade (A' in Fig. 2. ML-BS = 73% and BPP = 0.98) against the 206 Shikoku and Kyushu subclade (A"; ML-BS = 79% and BPP = 0.95). Within Subclade A', three 207 additional lineage groups were recognised: one consisted of Lineages A-1a and A-1b (ML-208 BS = 82% and BPP = 1.00); the second of Lineages A-2 and A-3 (ML-BS = 70% and

BPP = 0.98); and the third Lineages A-4, A-5, and A-6 (ML-BS = 79% and BPP = 1.00). The haplotypes obtained from *R. sakuraii* were included in Lineages A-2 and A-3. Lineage A-2 also contained *R. tagoi* haplotypes, although haplotypes were not shared between the two species.

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3.3. Population assignment: Nuclear DNA results

The results of the clustering analysis using STRUCTURE are shown in Fig. 3. For all samples, K = 2 was supported by the test of delta K, and two clusters (I and II) were recognised. Almost all samples were clearly assigned to each cluster (posterior probabilities  $\geq 80\%$ ), indicating strong genetic isolation between the two nDNA clusters. Although the division of the two nuclear clusters (I and II) did not completely correspond to that of the two mitochondrial clades (A and B), Cluster II was largely concordant with mitochondrial Subclade A', with the exception of Lineage A-1a (Fig. 3). Clusters I and II also were separated on the haplotype networks of some nuclear genes (e.g., NCX1, NFIA, and SLC8A3; Fig. S1). However, in relatively more variable genes like TYR, the haplotype relationships were highly complex and their separation was not clear (Fig. S1). Furthermore, haplotypes were more or less shared between Clusters I and II in all loci, indicating ILS in these nuclear genes. Since the two large clusters seemed to contain several subclusters, we independently reanalysed samples for the two clusters. Within Cluster I, the population assignment with K=2was supported (Fig. 3). In this clustering, the division of subclusters was still roughly correlated with the mt-lineages: the lineages from the main islands (A-1a, A-7, A-9a, and B-2ab) tended to form a subcluster and the lineages from the peripheral islands (A-8, A-9c, and B-1) formed another. One lineage, A-9b, included samples assigned to both of these subclusters. Except for Lineage A-9b, samples of the two subclusters were clearly assigned to either subcluster. In contrast, K = 3 was supported within Cluster II using the delta K test and likelihood distribution. In this division, R. tagoi Lineages A-1b and A-4, R. tagoi A-2, and R. sakuraii (A-3 and part of A-2) each formed a subcluster (Fig. 3). The separation of these subclusters was clear (posterior

probabilities > 80%), with a few exceptional samples in the *R. sakuraii* subcluster. By contrast, many samples of lineages A-5 and A-6 were not clearly assigned to particular subclusters, and showed intermediate genetic structures between *R. tagoi* of A-2 and *R. sakuraii*.

3.4. Divergence times of the mitochondrial lineages

The results of divergence dating for the major nodes on the mitochondrial genealogy are listed in Table 1. Although we applied the evolutionary rates of phylogenetically remote taxa (*Bufo* and *Leiopelma*) the divergence times obtained for the ingroup were similar in the two calibrations. Two major mt-clades (A and B: node 1 in Fig. 2) were estimated to have diverged 4.2–4.0 (95% highest posterior density interval [HPD] of 6.2–2.3) MYA. Then Subclades A' and A" (node 2) split 2.8–2.6 (4.1–1.6) MYA, followed by the separation within Clade B (node 22) 2.7–2.3 (4.3–1.2) MYA. The two lineages including *R. sakuraii* samples, A-2 and A-3, separated from each other 2.1–1.9 (3.1–1.1) MYA (node 7), followed by internal divergence during 1.4–0.9 (2.2–0.4) MYA (nodes 8 and 9). The most recently divergent lineages were B-2a and B-2b (node 23), which split at 1.4 (2.2–0.7) MYA. These estimates indicate that the divergence of each major mitochondrial clade/lineage began in the mid-to-late Pliocene and was approaching completion in the mid Pleistocene.

# 3.5. Historical demography

As shown above, the results of the population assignment were not completely concordant between mt- and n-DNA (Figs. 2 and 3). In estimating demographic parameters, we used only nDNA data because nuclear markers are thought to be more conservative than mitochondrial ones, which are more likely to be affected by introgression than the nuclear markers (Ballard and Whitlock 2004).

#### 3.5.1. Historical demography between Clusters I and II

First, we conducted a coalescent analysis using IMa2 for the two large nuclear clusters: I and II. Each parameter showed single peaks in their probability density distributions (Fig. S2). The parameter values obtained are listed in Table 2. The estimated population migration rate  $(2N_eM)$  for I to II (I  $\rightarrow$  II) was 0.52 (0.24–1.12). In the opposite direction; i.e. II  $\rightarrow$  I, the parameter value tended to be larger, with  $2N_eM_{II \rightarrow I}$  being 1.23 (0.70–2.14). The LLR test showed that all of these values were significantly larger than zero (p < 0.01), suggesting that clusters I and II have maintained a degree of gene flow after their divergence. However, strong to moderate genetic isolation would exist between the two clusters because the 2N<sub>e</sub>M values obtained were relatively small (ca. 1 or smaller: Wright, 1931; Waples and Gaggiotti, 2006; Reilly et al., 2012). The effective population size estimated for I, II, and their ancestor was 2.2 (1.7–2.9), 1.7 (1.3–2.3), and 0.4 (0.2–0.8) million individuals, respectively. The ancestral population size was smaller than those at present, as supported by parameter comparison of  $\theta$ (the posterior probabilities were 1.00 for each comparison). The population size of II tended to be smaller than that of I, but the tendency was not supported statistically (BPP < 0.95). The population divergence time (T) of I and II was estimated as 2.7 (4.4–2.2) MYA. Although its 95% HPD was relatively wide, this estimate was younger than the divergence time of the two major mt-clades (A/B; ca. 4.2–4.0 MYA), but almost equal to those of A'/A" (ca. 2.8–2.6 MYA) and B-1/B-2 (ca. 2.7–2.3 MYA) (Table 1).

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# 3.5.2. Historical demography between R. tagoi and R. sakuraii

Then, we compared demographic parameters between *R. tagoi* and *R. sakuraii*. As *R. tagoi* (*Rt*), we chose mt-Lineages A-2, 5, and 6, which were genetically close to *R. sakuraii* (*Rs*) in the mitochondrial and nuclear DNA analyses, as shown above (see Figs. 2 and 3). Since our dataset was not sufficiently informative to analyse a four-populations model, we combined Lineages A-5 and A-6 as a single group; these showed close genetic relationships in both

286 mitochondrial and nuclear analyses (Figs. 2, 3). We conducted two separate analyses under 287 different population schemes: (1) three-populations model, in which R. sakuraii (Rs) and R. 288 tagoi (Rt) Lineage A-2 were assumed to be mutually close compared to A-5 and 6, based on the 289 mtDNA genealogy, and (2) two-populations model based on the current classification (Rs vs. Rt 290 A-2, 5, and 6). 291 In the three-populations model, significant gene flow (p < 0.05 in the LLR test) was 292 detected only in R. tagoi A-5+6  $\rightarrow$  A-2 (2N<sub>e</sub>M<sub>Rt A-5+6</sub>  $\rightarrow$  Rt A-2 was 3.79 [0.75–9.50]; Fig. S3 and 293 Table 2) and A-5+6  $\rightarrow$  R. sakuraii (2N<sub>e</sub>M<sub>Rt A-5+6</sub>  $\rightarrow$ Rs was 0.40 [0.04–2.00]), and no significant 294 gene flow was recognized between R. sakuraii and R. tagoi A-2 (p > 0.05). These results 295 indicated that the genetic isolation between R. tagoi A-2 and A-5+6 was moderate ( $1 < 2N_eM \le$ 296 5), but the gene flow was strongly biased to one direction (from A-5+6 to A-2). Although gene 297 flow existed between the two species, the direction was limited (R. tagoi A-5+6  $\rightarrow$  R. sakuraii), 298 and the population migration rate obtained was small ( $2N_eM \le 1$ ), indicating strong genetic 299 isolation between R. sakuraii and R. tagoi lineages. The estimated effective population size (a 300 million individuals) was similar between R. tagoi A-2 (0.80 [0.34–2.06]) and A-5+6 (0.79 301 [0.38–1.76]), but was smaller in *R. sakuraii* (0.16 [0.07–0.32]). This tendency was supported in 302 the statistical test, in which  $N_e$  for R. sakuraii was significantly smaller than those for R. tagoi 303 lineages (BPP > 0.95). 304 We could not obtain a sufficient estimate for gene flow between the ancestral populations 305 because no obvious peaks of probability for the parameter  $2N_eM$  were recognised (Table 2). The 306 estimated ancestral population size  $(N_e)$  was 0.21 (0.01–4.06) for R. sakuraii + R. tagoi A-2, and 307 was 0.43 (0.23–0.77) for the common ancestor of R. sakuraii, R. tagoi A-2 and A-5+6. The 308 estimated  $N_e$  for the ancestors tended to be smaller than the present  $N_e$  for R. tagoi (A-2, A-5+6) 309 and larger than that for R. sakuraii, but the tendencies were not supported statistically (BPP < 310 0.95). The time of population divergence estimated for *R. sakuraii/R. tagoi* A-2 (1.1 [2.3–0.6]

MYA) was much younger than that for the ancestors (2.15 [6.11–1.31] MYA), although the credibility intervals largely overlapped.

In the two-populations model, significant gene flow from *R. tagoi* to *R. sakuraii* was again detected ( $2N_eM_{Rt \to Rs}$  was 0.51 [0.14–1.17]: Fig. S3 and Table 2), but such trend was not recognized in the opposite direction (Fig. S3 and Table 2). These results indicate strong to medium isolation between the two species, although small and unidirectional gene flow exists. The  $2N_eM$  value for *R. tagoi*  $\to$  *R. sakuraii* in this model was similar to the value for *R. tagoi* A-5+6  $\to$  *R. sakuraii* in the three-populations model shown above (Table 2).

The estimated  $N_e$  showed values and tendencies similar to those obtained in the three-populations model;  $N_e$  for R. sakuraii (0.17 [0.09–0.34]) was significantly smaller (BPP > 0.95) than that of R. tagoi (1.61 [0.99–2.65]). The estimates for ancestral  $N_e$  (0.37 [0.10–0.68] in the two-populations model) also are similar between the models. The divergence time estimated for the two species, 1.2 (2.9–0.6) MYA, was slightly older than that estimated by the three-population model (ca. 1.1 MYA).

# 4. Discussion

4.1. Discordance between the classification and patterns of genetic variation using different markers

Our new data and analyses confirmed the major patterns of mitochondrial genomic variation reported previously (Eto et al., 2012). Mitochondrial haplotypes obtained from *R. sakuraii* were genealogically embedded in those from *R. tagoi*, and neither species was monophyletic on the haplotype tree. The mitochondrial and nuclear data considered together indicate that *R. sakuraii* constitutes a single species lineage. *Rana sakuraii* corresponds largely to Lineage A-3 on the mitochondrial haplotype tree (Fig. 2), with its sister lineage being *R. tagoi* populations bearing mitochondrial haplotypes of Lineage A-2. Nonetheless, Lineage A-2 includes some *R. sakuraii* 

mitochondrial haplotypes. We examine the hypotheses of incomplete lineage sorting and gene flow as possible explanations for this pattern. The following three scenarios could explain the phylogenetic pattern of mitochondrial haplotypes of lineages A-2 and A-3 (Fig. 2): (1) recent speciation of R. sakuraii from R. tagoi Lineage A-2, which led to ILS of mtDNA at the species level; (2a) past mitochondrial introgression from R. tagoi A-2 to R. sakuraii; and (2b) introgression in the opposite direction (Fig. 4). If recent separation of R. sakuraii from R. tagoi A-2 was the case, the ILS hypothesis (1) would be the simplest explanation. However, if the speciation was shown to be old, especially much older than the divergence time within mt-Lineage A-2, this hypothesis would be rejected. Conversely, the past-introgression hypotheses (2) would be applicable if the speciation of the two species coincided with the split between Lineages A-2 and A-3 (2a), or the separation of these two lineages from the others (2b). Detection of historical gene flow between R. sakuraii and R. tagoi A-2 for the nuclear markers also would support the past introgression hypotheses. The genetic relationship based on the STRUCTURE analysis using nDNA was discordant with the mitochondrial genealogy, and R. sakuraii and R. tagoi A-2 tended to be separated in different subclusters (Fig. 3). This result likely reflects their heterospecific status. The demographic analysis using IMa2 showed that the separation of R. sakuraii from R. tagoi lineages (ca. 1.1 MYA and 1.2 MYA in three- and two-populations models, respectively; Table 2) was younger than the separation of mt-Lineages A-2 and A-3 (ca. 2.1–1.9 MYA; Table 1), and was similar to the divergence within these lineages (ca. 1.4–0.9 MYA). The date of speciation would correspond to, or be younger than, the population divergence time estimated by IMa in this case. So these results favour the ILS hypothesis, although the credibility intervals of these estimates overlapped. Based on the genealogy obtained (Fig. 2), mitochondrial introgression between R. tagoi A-2 and R. sakuraii happened several times if the hypotheses 2 were the case (for example, two independent introgression events should be presumed in the hypothesis 2a). Thus the

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introgression hypotheses assume rampant hybridization of R. tagoi A-2 and R. sakuraii in the past. The IM analyses based on two different models showed gene flow from R. tagoi to R. sakuraii. However, this unidirectional gene flow seems to depend largely on the flow from R. tagoi A-5+6 to R. sakuraii, because no significant flow between R. tagoi A-2 and R. sakuraii was detected (Table 2). These results do not support rampant hybridization of R. tagoi A-2 and R. sakuraii, even though inter-specific gene flow did exist. From these considerations, the ILS hypothesis would be more plausible than the introgression hypothesis to explain the relationships of the two species on the mitochondrial genealogy. The estimated time of the split of R. sakuraii and R. tagoi Lineage A-2 (ca. 1.2–1.1 MYA) is younger than those of other Japanese frogs (e.g., ca. 2.3 MYA between *Odorrana ishikawae/O*. splendida and ca. 1.7 MYA between O. amamiensis/O. narina [Matsui et al., 2005]; and around 5.7–4.0 MYA among *Bufo torrenticola* and two subspecies of *B. japonicus* [Igawa et al., 2006]), and seems to have occurred after the rough formation of the Japanese archipelago (see the next section). Although the ILS of mtDNA at the species level is relatively rare because of its small effective number of gene copies, it occurs occasionally in some situations, such as speciation within the past millions years. It could be applicable in the case of R. sakuraii and R. tagoi, because their speciation is estimated to be only about one million year ago. Rana sakuraii has several traits adaptive to stream breeding in contrast to the subterranean breeding R. tagoi, although they share many other characters (Matsui and Matsui, 1990). It suggests that the speciation of R. sakuraii was triggered by adaptation to a new breeding habitat, which is a process that often promotes rapid speciation (Coyne and Orr, 2004). 4.2. Evolutionary history of the two species Rana tagoi and R. sakuraii are endemic to the Japanese archipelago and no close relatives are known from the continent, although R. sauteri, a lotic breeding brown frog from Taiwan, is thought to be their sister lineage (Tanaka-Ueno et al., 1998). Our data do not contradict with this

idea (Fig. 1). Since the continental allies of R. sauteri are also unknown, the dispersal route of

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389 the ancestor of the R. tagoi complex to the Japanese mainland is uncertain. The estimated time 390 of separation of R. sauteri and R. tagoi complex varies between the calibrations (22.0–11.6 391 MYA; Table 1), but around the early to middle Miocene. In this period the opening of the Japan 392 Sea began (Iijima and Tada, 1990), although the Japanese and Ryukyu archipelagos, as well as 393 Taiwan were not yet isolated from the Eurasian continent (Chinzei and Machida, 2001). 394 Therefore the common ancestor of R. sauteri and the R. tagoi complex would have been distributed in the continental areas corresponding to the present Japanese archipelago to Taiwan, 395 396 but the ancestral allies would have been extinguished thereafter on the continent and the 397 Ryukyus, leaving relict species in Japan and Taiwan. 398 In any case, the ancestral population of the R. tagoi complex is thought to have diverged 399 into two major clades, A and B (Fig. 2), in the mid Pliocene (ca. 4.2–4.0 MYA). The ancient 400 Japanese archipelago was already roughly formed by the late Miocene (Chinzei and Machida, 401 2001), and the separation of lineages ancestral to the clades is thought to have occurred on the 402 archipelago. The ancestor at this period would have been a R. tagoi-like subterranean breeder 403 because all of the present genetic groups of the two species have a common larval trait (e.g., no 404 need to feed until metamorphosis) thought to be adapted to such an environment. 405 Then, the divergence within Clade A occurred in the late Pliocene (ca. 2.8–2.6 MYA), 406 separating populations on or near Honshu from ones on or near from Kyushu and Shikoku. 407 Cluster II as identified by nuclear markers (Fig. 3) is equivalent to populations diagnosed by 408 mitochondrial haplotype Subclade A' excluding Lineage A-1a. Lineages of Subclade A' occur 409 on Honshu, whereas those of mitochondrial Subclade A" are associated with Kyushu or Shikoku. 410 Because populations of mitochondrial haplotype Clade B also occur on Honshu, Honshu is 411 likely the ancestral source of this species complex, and expansion to the ancestral areas of 412 Kyushu and Shikoku likely produced the major cladogenetic event within mitochondrial Clade 413 A. Approximately 1.8–1.4 MYA (the divergence time estimated for mitochondrial haplotypes 414 of Lineages A-1a and A-1b), introgression of mitochondrial haplotypes from a population in

mitochondrial Subclade A' to one in Clade B produced the anomalous result that Lineage A-1a appears in an incorrect position on the mitochondrial haplotype tree. The best interpretation is that Lineage A-1a is closest phylogenetically to the lineages of mitochondrial Clade B as revealed by the nuclear markers, in contrast to its position on the mitochondrial haplotype tree. Occurrence of Subclade A' and Clade B in geographic proximity on Honshu further supports this interpretation. The divergences within SubcladesA', A", and Clade B started around 2.7–2.3 MYA, and splitting of the major mt-lineages was roughly completed by the middle Pleistocene (around 1.4 MYA). In this period, the populations on peripheral islands were isolated geographically, and some survived and evolved into the extant subspecies; *i.e.*, *R. t. yakushimensis* of Lineage A-8 and *R. t. okiensis* of B-1.

The estimated date of speciation of *R. sakuraii* was younger than the formation of the major population lineages discussed above. *Rana sakuraii* would have originated ca. 1.2–1.1 MYA based on the IM analysis (Table 2), likely in association with the adaptation to a new breeding environment as discussed above. The effective population size of *R. sakuraii* (ca. 0.2 million individuals) is smaller than that of the closest mt-lineage of *R. tagoi* (ca. 0.8 million individuals for Lineage A-2), and suggests that a small ancestral population adapted to stream breeding led to *R. sakuraii*.

# **5. Conclusion**

Our data reveal that *R. tagoi* comprises multiple species lineages, which form a paraphyletic group with respect to *R. sakuraii*. Because *R. sakuraii* arose only about one million years ago, incomplete lineage sorting of mitochondrial haplotypes best explains non-monophyly of *R. sakuraii* on the mitochondrial haplotype tree. Our study illustrates how mitochondrial haplotype phylogenies combined with multilocus demographic analyses of nuclear haplotypes permits precise resolution of species lineages and their genetic interactions.

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564	Supporting information
565	Additional supporting information may be found in the online version of this article.

# 567 **Table Captions** 568 **Table 1** The mean estimated divergence times (MYA) for *R. tagoi*, *R. sakuraii*, and the 569 outgroups. Values in parentheses are the 95% highest posterior density interval. For the node 570 numbers, refer to Fig. 2. 571 572 **Table 2** Demographic parameters estimated in the IM analysis. $N_e$ , effective population size 573 (million individuals); $2N_eM$ , effective population migration rate (number of gene 574 copies/generation), for which $2N_eM_{1\rightarrow 2}$ ( $2N_eM_{2\rightarrow 1}$ ) indicates gene flow from group 1 to 2 (2 to 575 1) forwards in time; T, population divergence time (MYA). Values supported by the highest 576 probability are shown as HiPt, and HPD95 indicates the 95% highest posterior density interval. 577 Parameters in bold indicate the values with statistical support, and characters in italics are those 578 with no significant peak of posterior probability density. 579 580 **Figure Captions** 581 Fig. 1 Map showing the sampling localities of Rana tagoi tagoi (circles), R. t. yakushimensis 582 (double circle), R. t. okiensis (stars), and R. sakuraii (triangles). Each species lineage inferred 583 using mitochondrial haplotypes is represented by different markers. For the locality information, 584 see Table S1. 585 586 Fig. 2 Maximum-likelihood tree based on the complete mitochondrial 16S and ND1 sequences 587 (2579 bp in total) for *Rana tagoi* and *R. sakuraii*. For the locality number, see Fig. 1. 588 Haplotypes in Clades A' and B are sampled from Honshu or the Oki Island (B-1). Haplotypes in 589 Clades A" are from Kyushu, Shikoku, or adjacent small islands. 590 591 Fig. 3 Results of STRUCTURE analyses based on the five nuclear genes. Each species lineage 592 inferred using mitochondrial haplotypes is separated by black vertical lines. (top) The best

593 clustering result (K = 2 clusters) for all 128 samples. (left bottom) Results with K=2 (best) and 3 594 for Cluster I. (right bottom) Results with K=3 (best) and 4 for Cluster II. 595 596 Fig. 4 Hypothesized scenarios for non-monophyly of mitochondrial haplotypes in R. sakuraii: 597 (1) the species-level ILS hypothesis; and (2) the past mitochondrial introgression hypothesis, in 598 which introgression occurred from R. tagoi Lineage A-2 to R. sakuraii (a) or in the opposite 599 direction (b). Solid and broken lines indicate the mitochondrial lineages of R. tagoi and R. 600 sakuraii, respectively. Grey arrows indicate massive mitochondrial introgression. 601 602 **Captions for supplementary materials** 603 **Table S1** The samples used in this study with information on the sampling localities, vouchers, 604 and GenBank accession numbers for each locus. KUHE, Graduate School of Human and 605 Environmental Studies, Kyoto University; TMP, temporary number. 606 607 **Table S2** The primers used to amplify mt- and n-DNA in this study. 608 609 **Table S3** Summary statistics of each locus. Tajima's D values; length of sequence after 610 alignment; variable sites (vs); number of haplotypes (h); haplotype diversity (Hd); and 611 nucleotide diversity  $(\pi)$ . 612 613 Fig. S1 Median-joining networks of five nuclear loci. The size of each circle reflects the relative 614 sample size of each haplotype. The color indicates nuclear clusters and species as follows: red = 615 n-Cluster I of *R. tagoi*; green = n-Cluster II of *R. tagoi*; light green = n-Cluster II of *R. sakuraii*. 616 Black circles and bars indicate median vectors and missing haplotypes, respectively. 617

618	Fig. S2 Posterior probability densities for divergence time (T), effective population size $(N_e)$ ,
619	and population migration rate $(2N_eM)$ of Clusters I and II obtained in the IM analyses. The
620	resultant values and 95% confidence intervals for each estimate are listed in Table 2.
621	
622	Fig. S3 Posterior probability densities for divergence time $(T, left top)$ , effective population size
623	$(N_e$ , left middle and bottom), and population migration rate $(2N_eM$ , right) of $R$ . $tagoi$ $(Rt)$
624	lineage A-2, A-5+6, and R. sakuraii (Rs). Estimates with no statistical support are indicated by
625	ns. The parameters obtained in three- and two-populations models are shown as triangles and
626	circles, respectively. The resultant values and 95% confidence intervals for each estimate are
627	listed in Table 2.
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630	The English in this document has been checked by at least two professional editors, both native
631	speakers of English. For a certificate, please see:
632	http://www.textcheck.com/
633	certificate/wc1m0N

**Table 1** The mean estimated divergence times (MYA) for *R. tagoi*, *R. sakuraii*, and the outgroups. Values in parentheses are the 95% highest posterior density interval. For the node numbers, refer to Fig.

	interval. For the flode if	
Node	Calibration I	Calibration II
1	4.00 (5.96–2.33)	4.16 (6.16–2.44)
2	2.58 (3.82–1.60)	2.82 (4.07–1.69)
3	2.31 (3.38–1.42)	2.46 (3.54–1.45)
4	1.84 (2.69–1.05)	1.73 (2.60–0.99)
5	1.16 (1.87–0.60)	1.04 (1.66-0.50)
6	1.32 (2.00-0.69)	1.35 (2.06–0.70)
7	1.87 (2.78–1.05)	2.08 (3.07–1.21)
8	0.95 (1.60-0.44)	1.13 (1.78–0.58)
9	0.88 (1.40-0.41)	1.39 (2.16-0.72)
10	1.92 (2.85–1.09)	2.12 (3.14–1.24)
11	1.75 (2.59-0.96)	1.69 (2.50-0.90)
12	0.50 (0.95-0.14)	0.42 (0.81-0.11)
13	1.15 (1.85–0.56)	1.50 (2.30-0.82)
14	0.36 (0.64-0.13)	0.53 (0.90-0.21)
15	2.31 (3.36–1.34)	2.54 (3.70-1.53)
16	0.79 (1.30-0.34)	0.85 (1.38-0.37)
17	0.20 (0.43-0.03)	0.17 (0.37-0.01)
18	1.68 (2.53-0.99)	1.98 (2.91-1.19)
19	0.53 (0.90-0.21)	0.59 (1.01-0.24)
20	1.04 (1.67–0.46)	1.40 (2.11-0.73)
21	1.46 (2.22–0.76)	1.54 (2.34–0.83)
22	2.71 (4.34–1.40)	2.29 (3.50-1.24)
23	1.40 (2.15–0.71)	1.35 (2.07-0.71)
24	0.80 (1.28-0.36)	0.82 (1.32-0.39)
25	0.90 (1.46-0.42)	1.08 (1.74–0.52)
26	0.04 (0.13-0.03)	0.26 (0.53-0.05)
O-1	22.02 (35.97–11.08)	11.59 (18.29–6.42)

**Table 2** Demographic parameters estimated in the IM analysis.  $N_e$ , effective population size (million individuals);  $2N_eM$ , effective population migration rate (number of gene copies/generation), for which  $2N_eM_{1\rightarrow 2}$  ( $2N_eM_{2\rightarrow 1}$ ) indicates gene flow from group 1 to 2 (2 to 1) forwards in time; T, population divergence time (MYA). Values supported by the highest probability are shown as HiPt, and HPD95 indicates the 95% highest posterior density interval. Parameters in bold indicate the values with statistical support and characters in italics are those with no significant peak of posterior

probability density.

<u>probability</u>	z density.					
	$N_{I}$	$N_2$	$N_{\it ancestor}$	$2N_eM_{1\rightarrow 2}$	$2N_eM_{2\rightarrow 1}$	T
(1) Cluster	I vs. (2) Cluste	er II				
HiPt	2.18	1.73	0.40	0.52	1.23	2.72
HPD95	(1.70-2.87)	(1.34-2.30)	(0.17-0.79)	(0.24-1.12)	(0.70-2.14)	(2.10-4.29)
Three-pops	. model: (1) R.	sakuraii vs. (	(2) R. tagoi lir	1		
HiPt	0.16	0.80	0.21	0.01	0.00	1.05
HPD95	(0.07-0.32)	(0.34-2.06)	(0.01-4.06)	(0.00-2.46)	(0.00-0.84)	(0.63-2.26)
Three-pops	. model: (1) <i>R</i> .			neage A-5, 6		
HiPt	0.16	0.79		0.46	0.40	-
HPD95	(0.07-0.31)	(0.38-1.76)	-	(0.00-2.52)	(0.04-2.00)	-
	. model: (1) <i>R</i> .	• • • • • • • • • • • • • • • • • • • •				
HiPt	0.80	0.79		0.17		-
HPD95	(0.34-2.06)	(0.38-1.76)	-	(0.00-3.74)	(0.75 - 9.50)	-
		2.5			(2) 5	
				•	(2) R. tagoi A	
HiPt	0.21		0.43		0.04	2.15
HPD95	(0.01-4.06)	(0.38-1.76)	(0.23-0.77)	(0.00-81.15)	(0.00-34.07)	(1.31-6.11)
<b>T</b>	1 1 (1) D	1 (2		\ ~ ~ <		
	model: (1) R. s	·			0.74	1.01
HiPt				0.01		1.21
HPD95	(0.09-0.34)	(0.99-2.65)	(0.10-0.68)	(0.00-3.19)	(0.14-1.17)	(0.56-2.85)

**Table S1** The samples used in this study with information on the sampling localities, vouchers, and GenBank accession numbers for each locus. KUHE, Graduate School of Human and Environmental Studies, Kyoto University; TMP, temporary number.

loc.		voucher		<u> </u>	GenF	Bank accession	on no.		
nos.	locality	(KUHE)	mt-lineage	mtDNA (16S, ND1)	NCX1	NFIA	POMC	SLC8A3	TYR
	tagoi tagoi								
	Mutsu City, Aomori Pref.	44827	A-1a	AB639413, AB639593					
	Noshiro City, Akita Pref.	46598	A-1a	AB968306			AB969021		
	Ichinoseki City, Iwate Pref.	36699 45622	A-1a	AB639413, AB639598					
	Sendai City, Miyagi Pref. Yamagata City, Yanagata Pref.	45622 37543	A-1a A-1a	AB968302 AB639417, AB639601			AB969017		
	Nihonmatsu City, Fukushima Pref.	29595	A-1a A-1a	AB639417, AB639604 AB639419, AB639604					
	Nihonmatsu City, Fukushima Pref.	36330	A-1a A-2	AB639474, AB639643					
	Daigo town, Ibaraki Pref.	42344	A-1a	AB639420, AB639605					
O	Duigo town, Iburuki 1101.	43723	A-1a	AB968270			AB968980		
		43886	A-2	AB639421, AB639646					
		TMP_081122-1	A-2	AB968251			AB969029		
9	Tsukuba City, Ibaraki Pref.	_ 42747	A-2	AB639479, AB639648	AB968709	AB968838	AB968964	AB969093	AB96922
	Ichihara City, Chiba Pref.	28409	A-2	AB639482, AB639652	AB968673	AB968803	AB968929	AB969058	AB96918
		46172	A-2	AB968305	AB968764	AB968893	AB969020	AB969149	AB96927
	Kanuma City, Tochigi Pref.	40166	A-1a	AB639422, AB639609					
	Uonuma City, Nigata Pref.	36896	A-1a	AB639429, AB639612					
13	Nakanojo Town, Gunma Pref.	44810	A-1a	AB968281			AB968994		
		44811	A-1a	AB968282			AB968995		
		22930	A-4	AB639487, AB639657					
		22936	A-4	AB639487, AB639657					
1.4		44797	A-4	AB968280			AB968993		
	Saku City, Nagano Pref.	43980	A-2	AB968274			AB968987		
15	Akiruno City, Tokyo Pref.	42452 42453	A-2	,			AB968960		
16	Fujikawaguchiko Town, Yamanashi Pref.	42453 45558	A-2 A-2	AB968263 AB968300			AB968961 AB969015		
10	rujikawaguchiko 10wh, Tamahashi Piel.	43480					AB968971		
17	Minobu Town, Yamanashi Pref.	45480 45552	A-6 A-2	AB639493, AB639663 AB968299			AB968971 AB969014		
1 /	Willioud Town, Tamanasin Frei.	45549	A-2 A-6	AB968298			AB969013		
18	Izu City, Shizuoka Pref.	43468	A-2	AB639485, AB639655					
	Hokuto City, Yamanashi Pref.	43483	A-5	AB639489, AB639659					
	Nagano City, Nagano Pref.	18005	A-5	AB639488, AB639658					
	Kurobe City, Toyama Pref.	45102	A-1a	AB968287			AB969002		
		45103	A-1a	AB968288			AB969003		
		45014	A-5	AB968283			AB968998		
		45099	A-5	AB968286	AB968745	AB968875	AB969001	AB969130	AB96925
25	Takayama City, Gifu Pref.	42048	A-1a	AB968261	AB968700	AB968829	AB968955	AB969084	AB96921
		43018	A-1a	AB639434, AB639617	AB968711	AB968840	AB968966	AB969095	AB96922
26	Gujo City, Gifu Pref.	14228	A-5	AB639490, AB639660	AB968652	AB968780	AB969027	AB969036	AB96916
27	Fujieda City, Shizuoka Pref.	17955	A-6	AB639498, AB639668					
28	Neba Village, Nagano Pref.	27335	A-6	AB639500, AB639670					
		27337	A-6	AB968254			AB968921		
	Shinjo City, Aichi Pref.	45913	A-6	AB968304			AB969019		
	Okazaki City, Aichi Pref.	45910	A-6	AB968303			AB969018		
	Ise City, Mie Pref.	42829	A-6	AB639502, AB639672					
	Ibigawa Town, Gifu Pref.	27388	A-1a	AB639436, AB639619					
34	Takashima City, Shiga Pref.	43925	A-1a	AB968273			AB968986		
25	Taga Tayun Shiga Draf	43924 43512	A-1b B-2a	AB968272 AB968266			AB968985 AB968973		
	Taga Town, Shiga Pref. Matsuzaka City, Mie Pref.	41484	В-2a В-2a	AB639551, AB639716					
	Joyo City, Kyoto Pref.	41554	B-2a B-2a	AB639549, AB639714					
	Odai Town, Mie Pref.	40190	B-2a B-2a	AB639553, AB639718					
50	Can Town, who I let.	45047	В-2а В-2а	AB968284			AB968999		
39	Gobo City, Wakayama Pref.	41229	B-2a B-2a				AB968948		
	Kyoto City, Kyoto Pref.	42342	A-1b	AB968262			AB968957		
.0	y	44828	A-1b	AB968307			AB968997		
		42319	B-2a	AB639464, AB639712					
41	Nantan City, Kyoto Pref.	41408	A-1b	AB639452, AB639630					
		41405	B-2a	AB968259	AB968694	AB968823	AB968949	AB969078	AB96920
		41430	B-2a	AB968260	AB968697	AB968826	AB968952	AB969081	AB96920
42	Sasayama City, Hyogo Pref.	10307	A-1b	AB639469, AB639639	AB968647	AB968776	AB968903	AB969031	AB96916
43	Kobe City, Hyogo Pref.	45392	A-1b	AB968297	AB968756	AB968885	AB969012	AB969141	AB96926
	Taka Town, Hyogo Pref.	10330	B-2a	AB639564, AB639729					
	Kyotango City, Kyoto Pref.	14171	A-1b	AB968253			AB968907		
46	Toyooka City, Hyogo Pref.	42711	A-1b	AB639466, AB639637					
		42714	B-2a	AB639467, AB639729					
	Wakasa Town, Tottori Pref.	34743	A-1b	AB639473, AB639642					
	Mimasaka City, Okayama Pref.	27659	B-2a	AB639464, AB639730					
	Misasa Town, Tottori Pref.	24574	B-2b	AB639465, AB639731					
	Kagamino Town, Okayama Pref.	29739	B-2b	AB968256			AB968932		
	Shobara City, Hiroshima Pref.	36040	B-2b	AB639469, AB639734					
	Izumo City, Shimane Pref.	18877	B-2b	AB639467, AB639734					
	Higashihiroshima City, Hiroshima Pref.	30262	B-2b	AB639472, AB639737					
58	Hatsukaichi City, Hiroshima Pref.	unnumbered	B-2b	AB639571, AB639736					
	at 11 at 22	43167	B-2b	AB968265			AB968968		
60	Shimonoseki City, Yamaguchi Pref.	34516	B-2b	AB639575, AB639740					
	Minamiawaji City, Hyogo Pref.	43885	A-7	AB639504, AB639673					
		THE T	. –	AD (00000 - 1					
62	Manno Town, Kagawa Pref. Miyoshi City, Tokushima Pref.	TMP_T2882 TMP_T3498	A-7 A-7	AB639505, AB639674 AB968308			AB969024 AB969025		

64 Toyo Town, Kochi Pref.	29464	A-7	AB639510, AB639679	AB968675	AB968804	AB968930	AB969059	AB969188
65 Saijo City, Ehime Pref.	27679	A-7	AB639507, AB639676					
55 2 mg 5 5 5 5 7 7 5 5 5 5 5 5 5 5 5 5 5 5 5	43078	A-7	AB968264			AB968967		
66 Saiyo City, Ehime Pref.	TMP_T2241	A-7	AB639509, AB639678					
67 Kitakyushu City, Fukuoka Pref.	28612	A-9a	AB968255			AB968924		
68 Beppu City, Oita Pref.	43637	A-9a	AB639519, AB639688					
69 Yatsushiro City, Kumamoto Pref.	27562	A-9a	AB639524, AB639691					
70 Amakusa City, Kumamoto Pref.	30342	A-9a	AB639525, AB639692					
71 Kanoya City, Kagishima Pref.	27295	A-9a	AB639530, AB639697					
72 Sasebo City, Nagasaki Pref.	27140	A-9a	AB639518, AB639687					
73 Goto City, Nagasaki Pref.	45359	A-9a	AB968295			AB969010		
, a coto city, i tugusumi i ion	45362	A-9a	AB968296			AB969011		
74 Nobeoka City, Miyazaki Pref.	27121	A-9b	AB639528, AB639695					
75 Nishimera Village, Miyazaki Pref.	26088	A-9b	AB639529, AB639696					
76 Miyakonojo City, Miyazaki Pref.	30907	A-9b	AB639532, AB639699					
77 Kimotsuki City, Kagoshima Pref.	43397	A-9b	AB639533, AB639700					
78 Kinko Town, Kagoshima Pref.	27678	A-9b	AB639536, AB639703					
79 Goto City, Nagasaki Pref.	31539	A-9c	AB639538, AB639705					
79 Shinkamigoto City, Nagasaki Pref.	45149	A-9c	AB968291			AB969006		
75 Similaring oto City, Hagasaki Frei.	TMP_110216-1	A-9c	AB968252			AB968906		
80 Goto City, Nagasaki Pref.	44316	A-9c	AB968278			AB968991		
oo doto city, ragasaki rioi.	44317	A-9c	AB968279			AB968992		
	45355	A-9c	AB968294			AB969009		
R. t. okiensis	43333	11-70	11000274	111111111111111111111111111111111111111	7 <b>ID</b> 700002	71B)0)00)	7110707130	7 <b>ID</b> 707200
50 Okinoshima Town, Shimane Pref.	10818	B-1	AB639576, AB639742	AB968649	AB968778	AB968905	AB969033	AB969162
30 Okinosiima Town, Siimane Trei.	22341	B-1	AB639579, AB639742					
51 Nishinoshima Town, Shimane Pref.	43647	B-1	AB639580, AB639742					
R. t. yakushimensis	43047	<b>D</b> -1	AD037300, AD037742	AB)00724	AD700033	Abbooti	AD 707100	AD707230
81 Yakushima Town, Kagoshima Pref.	10182	A-8	AB639578, AB639741	AB968646	AB968775	AB968902	AB969030	AB969159
or rakusinina rown, Ragosinina rici.	45177	A-8	AB968292			AB969007		
	45182	A-8	AB968293			AB969008		
R. sakuraii	43102	71-0	11000273	71B)00732	711111111111111111111111111111111111111	71B)0)000	1111000137	1111000200
11 Kanuma City, Tochigi Pref.	43633	A-2	AB968268	AB968720	AB968849	AB968975	AB969104	AB969232
	43634	A-2	AB968269			AB968976		
	43635	A-2	AB639423, AB639744					
15 Akiruno City, Tokyo Pref.	42450	A-2	AB639583, AB639744					
20 1 2222 2220	43740	A-2	AB968271			AB968981		
17 Minobu Town, Yamanashi Pref.	45620	A-2	AB968301			AB969016		
20 Shizuoka City, Shizuoka Pref.	unnumbered	A-2				AB969022		
20 Simbuona City, Simbuona 1101	44254	A-3	AB968275			AB968988		
	44286	A-3	AB968277			AB968990		
21 Shizuoka City, Shizuoka Pref.	44256	A-2	AB968276			AB968989		
22 Matsumoto City, Nagano Pref.	22887	A-2	AB639585, AB639746					
24 Kurobe City, Toyama Pref.	45105	A-3	AB968289			AB969004		
24 Kurobe City, Toyama Pref.	45106	A-3	AB968290			AB969005		
32 Katsuyama City, Fukui Pref.	43591	A-3	AB968267			AB968974		
38 Odai Town, Mie Pref.	27647	A-3	AB639554, AB639719					
30 Gdai 10wii, iviic 11ci.	40309	A-3	AB639555, AB639720					
	45049	A-3	AB968285			AB969000		
41 Nantan City, Kyoto Pref.	41412	A-3	AB639455, AB639632					
41 Ivanian City, Ryoto 1101.	unnumbered	A-3	AB639454, AB639631					
48 Wakasa Town, Tottori Pref.	34740	A-3	AB968257			AB968938		
59 Iwakuni City, Yamaguchi Pref.	43893	A-3	AB639590, AB639750					
R. tsushimensis	73073	11 5	11000000, 110000100	. 12/00/2/	. 112700000	. 12700704	.110/0/110	. 11.7072TI
Tsushima City, Nagasaku Pref.	10606		AB639592, AB639752	_	_	_	_	_
R. kobai	10000		71B037372, 71B037732					
Amami City, Kagoshima Pref.	10051		AB685768	_	_	_	_	_
R. ulma	10051		11005/00	_	_	_	_	_
Higashi Village, Okinawa Pref.	10056		AB685780	_	_	_	_	_
R. sauteri	10050		11005/00	_	_	_	_	_
Chiayi County, Taiwan	6894		AB685767	_	_	_	_	_
omaji county, turnun	00 <i>7</i> -т		11000707					

**Table S2** The primers used to amplify mt- and n-DNA in this study.

Target	Name	Sequence	Reference
16S	L1507	TACACACCGCCCGTCACCCTCTT	Shimada et al (2011)
	H1923	AAGTAGCTCGCTTAGTTTCGG	Shimada et al (2011)
	L1879	CGTACCTTTTGCATCATGGTC	Shimada et al (2011)
	H2315	TTCTTGTTACTAGTTCTAGCAT	Shimada et al (2011)
	L2188	AAAGTGGGCCTAAAAGCAGCCA	Matsui et al (2006)
	Wilkinson_6	CCCTCGTGATGCCGTTGATAC	Wilkinson et al (2002)
	16L1	CTGACCGTGCAAAGGTAGCGTAATCACT	Hedges (1994)
	16H1	CTCCGGTCTGAACTCAGATCACGTAGG	Hedges (1994)
ND1	L3032	CGACCTCGATGTTGGATCAGG	Shimada et al (2011)
	ND1_Htago	GRGCRTATTTGGAGTTTGARGCTCA	Eto et al (2012)
	ND1_Ltago	GACCTAAACCTCAGYATYCTATTTAT	Eto et al (2012)
	tMet_H	AGGAAGTACAAAGGGTTTTGATC	Shimada et al (2011)
NCX1	NCX1F	ACAACAGTRAGRATATGGAA	Shimada et al. (2011)
	NCX1R1	GCCATATCTCTCCTCGCTTCTTC	Eto et al (2013)
NFIA	NFIA-005_I	FTTTGTCACATCAGGTGTTTT	This study
	NFIA-005_I	RCTTGCCTTGGCTGCT	This study
POMC	POMC1	GAATGTATYAAAGMMTGCAAGATGGWC	Wiens et al. (2005)
	POMC7	TGGCATTTTTGAAAAGAGTCAT	Smith et al. (2005)
SLC8A3	SCF_2F	CAAACACAGRGSAATTATGAT	Shimada et al (2011)
	SCF_2R	ATAATYCCAACTGARAACTC	Shimada et al (2011)
TYR	Tyr_L1	CCCCAGTGGGYRCCCARTTCCC	Kuraishi et al (2013)
	Tyr_H1	CCACCTTCTGGATTTCCCGTTC	Kuraishi et al (2013)

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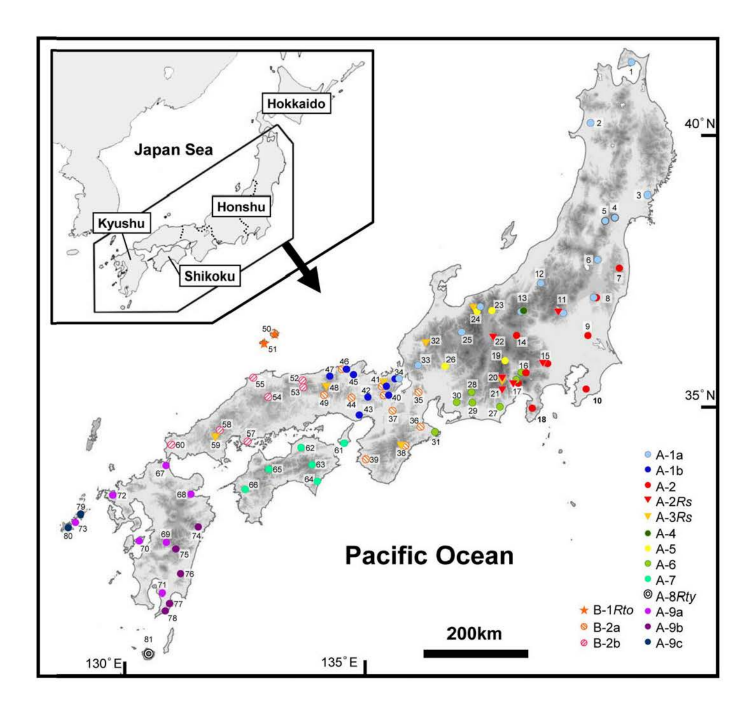
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**Table S3** Summary statistics of each locus. Tajima's D values; length of sequence after alignment; variable sites (vs); number of haplotypes (h); haplotype diversity (Hd); and nucleotide diversity  $(\pi)$ .

( ),	,														
Tajima's $L$ sites	vs h	Hd	$\pi$	VS	h	Hd	$\pi$	VS	h	Hd	$\pi$	VS	h	Hd	$\pi$
,	whole (n =	128)		mt	-lineaş	ge A-1a (1	n = 18)	m	t-linea	ge A-1b (	n = 9	mt-	lineag	e A-2Rt (	n=12)
16S -1.307 1612	285 11	5 0.998	0.024	44	15	0.980	0.006	50	9	1.000	0.010	51	20	0.970	0.010
<i>ND1</i> -0.816 967	287 10	4 0.997	0.047	44	14	0.974	0.010	70	9	1.000	0.026	53	10	0.970	0.016
NCX1 (SLC8A1) -0.648 505	26 37	0.851	0.006	7	5	0.651	0.003	3	4	0.525	0.002	5	4	0.649	0.003
<i>NFIA</i> -1.626 414	18 21	0.735	0.003	3	5	0.548	0.002	3	4	0.700	0.003	2	3	0.177	0.000
<i>POMC</i> -1.538 475	40 48	0.870	0.007	9	6	0.712	0.004	7	6	0.775	0.004	12	10	0.859	0.006
SLC8A3 (NCX3 -1.429 524	21 23		0.003	3	4	0.236	0.001	2	3	0.242	0.000	8	6	0.659	0.002
<i>TYR</i> -1.210 318	50 97		0.017	17	12	0.867	0.013	15	12	0.958	0.018	20	15	0.946	0.018
·															
,	mt-line	age A-2Rs	(n = 9)	m	t-linea	ige A-3 (n	= 12)	m	t-line	age A-4 (1	n=3)	mt-lineage A-5 $(n = 5)$			
16S	26 7	0.964	0.006	43	12	1.000	0.009	7	2	0.667	0.003	41	5	1.000	0.012
ND1	38 7	0.964	0.016	37	8	0.939	0.013	9	2	0.667	0.006	38	4	0.900	0.018
NCX1 (SLC8A1)	2 2	0.125	0.001	2	3	0.163	0.000	3	2	0.533	0.003	3	3	0.607	0.002
NFIA	1 2	0.125	0.000	2	3	0.163	0.000	2	3	0.600	0.002	1	2	0.250	0.001
POMC	6 5	0.556	0.003	9	5	0.652	0.004	3	2	0.533	0.003	4	3	0.607	0.003
SLC8A3 (NCX3)	1 2	0.125	0.000	2	3	0.554	0.001	1	2	0.533	0.001	1	2	0.571	0.001
TYR	11 4	0.442	0.008	9	5	0.493	0.007	1	2	0.533	0.002	12	7	0.964	0.018
	mt-lir	eage A-6 (	n=8)	m	t-line	age A-7 (1	n=7)	m	t-line	age A-8 (1	n=3)	mt	t-linea	ge A-9a (	$\overline{n=8}$
16S	25 8	1.000	0.005	27	7	1.000	0.006	3	2	0.667	0.001	28	7	0.964	0.006
ND1	15 6	0.964	0.005	32	7	1.000	0.011	3	2	0.667	0.002	21	7	0.964	0.006
NCX1 (SLC8A1)	4 5	0.505	0.002	2	3	0.538	0.001	4	3	0.733	0.004	6	6	0.792	0.004
NFIA	2 3	0.425	0.001	2	3	0.275	0.001	2	3	0.600	0.002	2	3	0.433	0.001
POMC	9 7	0.850	0.006	_	1	_	_	4	3	0.600	0.003	2	3	0.242	0.001
SLC8A3 (NCX3)	2 2	0.363	0.001	2	2	0.440	0.002	_	1	-	-	1	2	0.264	0.001
TYR	16 13	0.975	0.016	9	7	0.846	0.009	10	4	0.800	0.013	11	7	0.692	0.009
	mt-lin	eage A-9b	(n=5)	m	t-linea	ige A-9c (	n = 6	m	ıt-line	age B-1 (r	n=3	mt	-lineag	ge B-2a (r	n = 12
16S	44 5		0.014	41	5	0.933	0.014	4	3	1.000	0.002	36	10	0.970	0.007
ND1	36 5	1.000	0.018	39	4	0.800	0.023	-	1	-	-	42	10	0.970	0.013
NCX1 (SLC8A1)	5 5	0.822	0.003	6	7	0.879	0.003	_	1	_	_	2	3	0.636	0.002
NFIA	2 2	0.200	0.001	4	5	0.756	0.004	1	2	0.333	0.001	1	2	0.228	0.001

POMC	6	3	0.378	0.003	7	5	0.742	0.006	8	5	0.933	0.008	9	7	0.851	0.004
SLC8A3 (NCX3)	3	4	0.644	0.001	-	1	-	-	3	4	0.867	0.003	4	5	0.361	0.002
TYR	12	9	0.978	0.013	11	6	0.848	0.014	6	5	0.933	0.009	10	9	0.812	0.010
									·							
	m	t-linea	ige B-2b (	(n=8)		n-clu	ster I (n =	68)		n-clus	ster II (n =	56)				

	mt-lineage $B$ -2 $b$ ( $n = 8$ )					n-cluster I ( $n = 68$ )					n-cluster II ( $n = 56$ )				
16S	24	7	0.964	0.006	204	61	0.996	0.026	176	53	0.997	0.018			
ND1	33	7	0.964	0.014	218	57	0.994	0.049	192	47	0.994	0.037			
NCX1 (SLC8A1 )	7	6	0.833	0.005	19	26	0.875	0.005	12	13	0.483	0.002			
NFIA	1	2	0.125	0.000	11	16	0.493	0.002	6	8	0.486	0.001			
POMC	4	4	0.350	0.001	29	26	0.784	0.006	24	24	0.849	0.007			
SLC8A3 (NCX3)	3	3	0.633	0.002	9	10	0.593	0.002	12	12	0.577	0.002			
TYR	10	13	0.967	0.010	34	58	0.935	0.014	33	43	0.907	0.016			
												•			



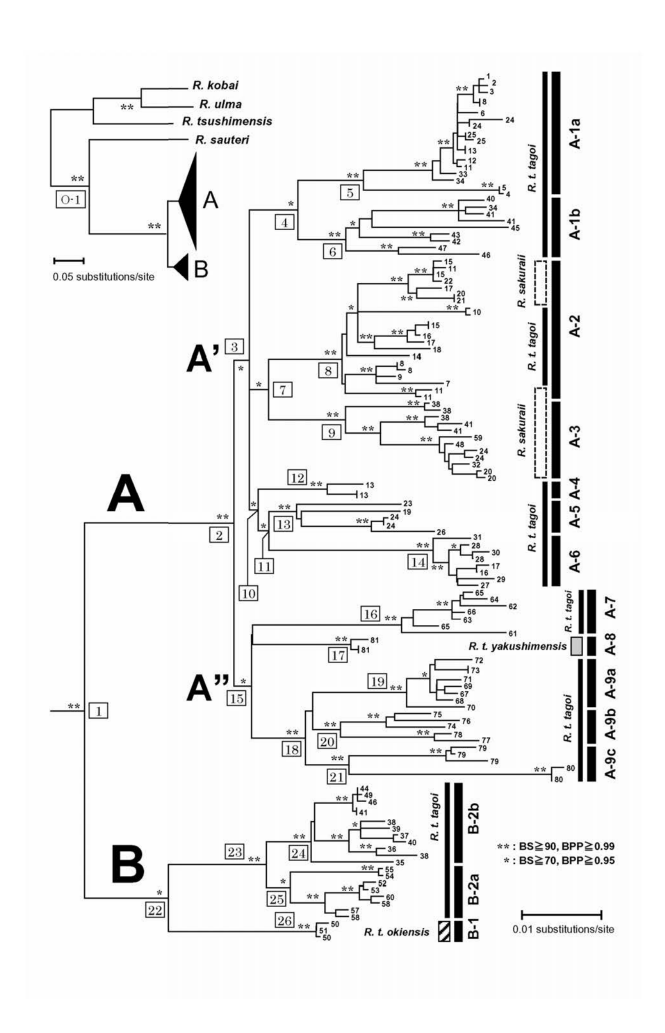


Fig. 2

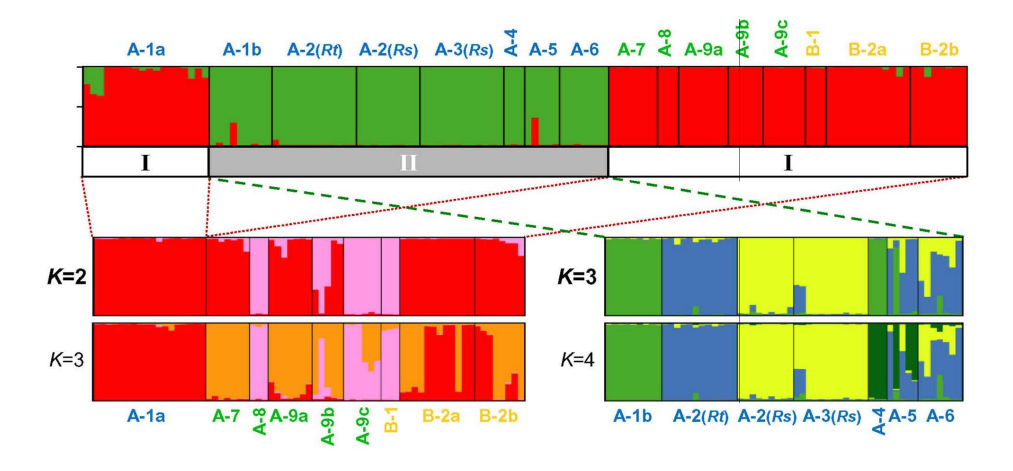


Fig. 3

# 1: ILS 2: introgression b a A'(node 3): ca.2.8-2.6<sub>MYA</sub> other ••••• R. tagoi lineages ←node 7: ca.2.1-1.9 муа A-3 ← nodes 8 and 9 :са.1.4-0.9 мүд R. sakuraii R. sakuraii R. tagoi R. sakuraii R. tagoi R. tagoi

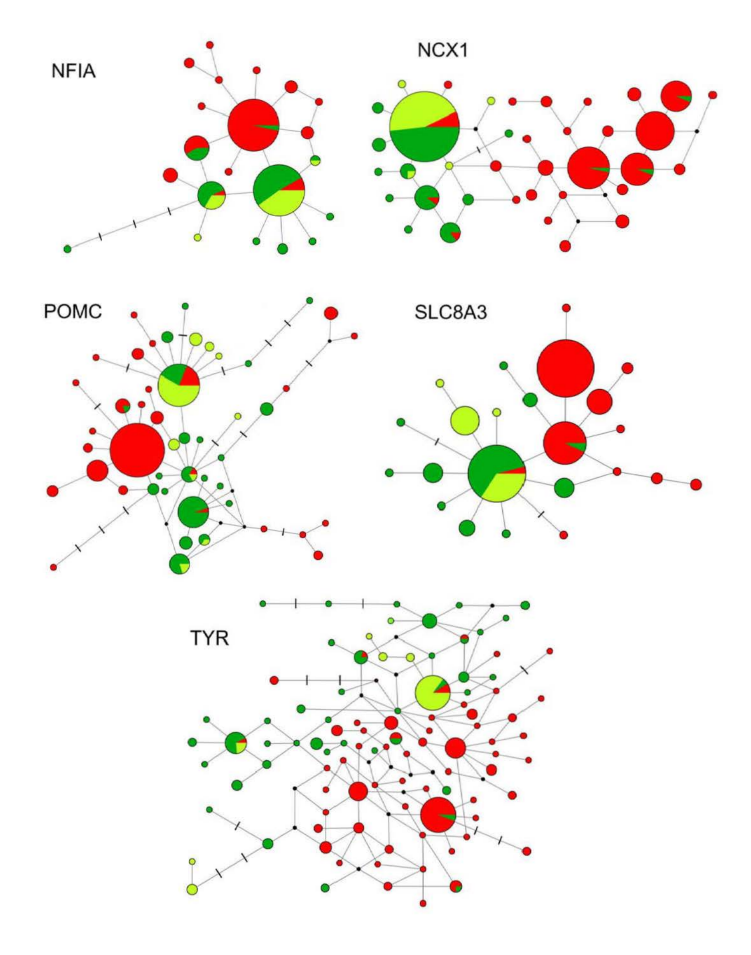


Fig. S1

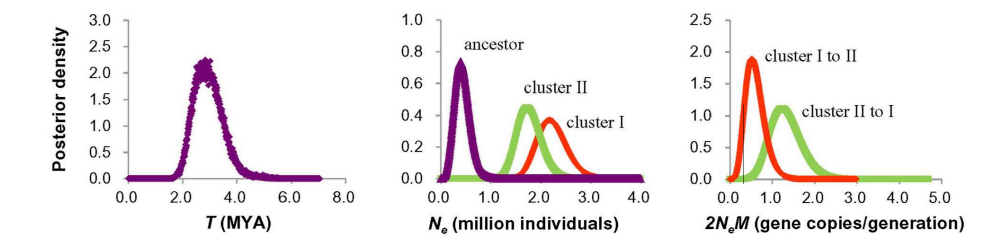


Fig. S2

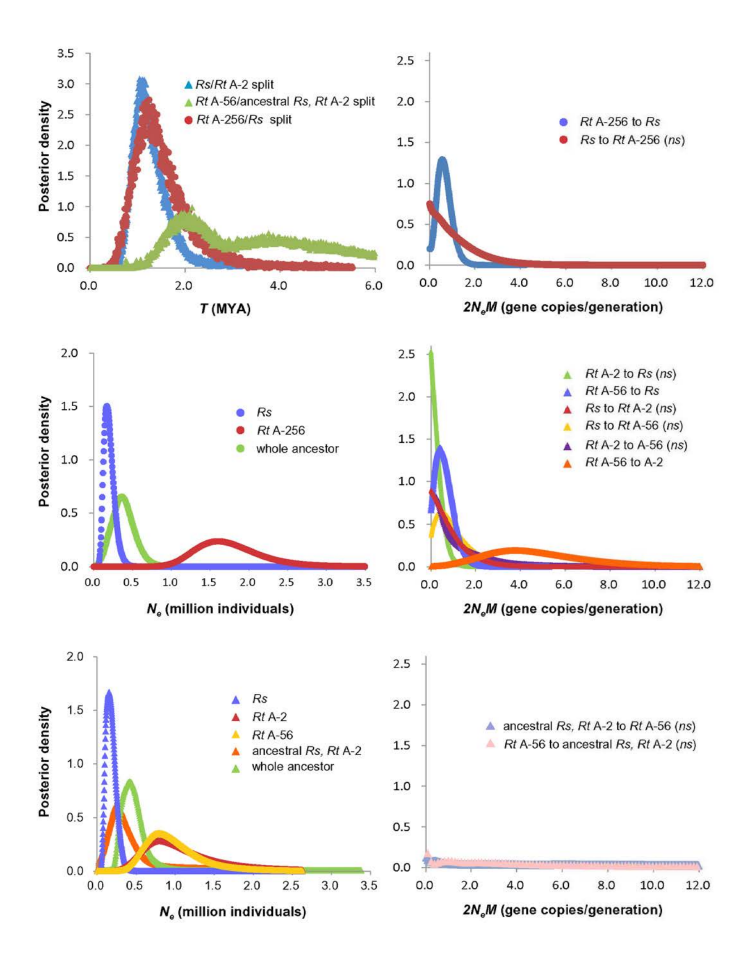
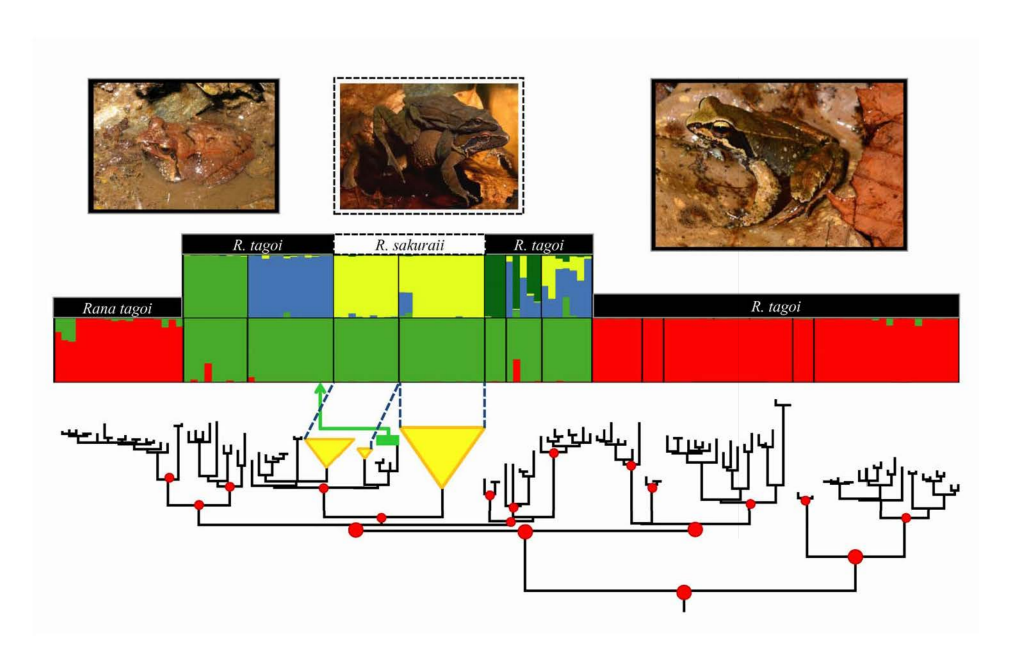


Fig. S3



Graphical abstract