Hybrid zone genomics supports candidate species in Iberian Alytes obstetricans

Christophe Dufresnes1,∗, Íñigo Martínez-Solano2

Abstract. While estimates of genetic divergence are increasingly used in molecular taxonomy, hybrid zone analyses can provide decisive evidence for evaluating candidate species. Applying a population genomic approach (RAD-sequencing) to a fine-scale transect sampling, we analyzed the transition between two Iberian subspecies of the common midwife toad (Alytes obstetricans almogavarii and A. o. pertinax) in Catalonia (northeastern Spain), which putatively diverged since the Plio-Pleistocene. Their hybrid zone was remarkably narrow, with extensive admixture restricted to a single locality (close to Tarragona), and congruent allele frequency clines for the mitochondrial (13 km wide) and the average nuclear genomes (16 km wide). We also fitted clines independently for 89 taxon-diagnostic SNPs: most of them behave like the nuclear background, but a subset (13%) is completely impermeable to gene flow and might be linked to barrier loci involved in hybrid incompatibilities. Assuming that midwife toads are able to disperse in the area of contact, we conclude that these taxa experience partial reproductive isolation and represent incipient species, i.e. Alytes almogavarii and Alytes obstetricans. Interestingly, their evolutionary age and mitochondrial divergence fall below the thresholds proposed in molecular systematics studies, emphasizing the difficulty of predicting the outcome of secondary contacts between young lineages entering the grey zone of speciation.

Keywords: Alytes almogavarii, Catalan midwife toad, RAD sequencing, reproductive isolation, speciation.

Introduction

The delimitation of cryptic species by phylogeographers is inherently challenged by the continuous and dynamic nature of speciation processes (Avise, 2000; Heithoff, 2018; Struck et al., 2018; Chenuil et al., 2019). In the era of the “molecularisation of taxonomy” (Lee, 2004), evolutionary biologists may never agree on how much reproductive isolation and genetic differentiation is required to consider diverging populations as incipient species (Coyne and Orr, 2004; De Queiroz, 2007; Padial et al., 2010). In particular, hybridizing taxa, as well as allopatric taxa of equivalent divergence, fall within a so-called grey zone of speciation, and thus remain at the heart of taxonomic disputes (Roux et al., 2016).

In the wake of recent attempts to relate the evolutionary divergence of phylogeographic lineages with levels of introgression across hybrid zones (Singhal and Moritz, 2013; Dufresnes et al., 2019a), the concept of “speciation thresholds” might circumvent some practical issues (Fouquet et al., 2007; Vieites et al., 2009). As theoretically and empirically supported, reproductive isolation tends to increase rapidly due to the accumulation of barrier loci after a certain degree of genetic divergence, above which the probability to merge back and reverse the speciation process is considerably reduced (Barton, 1983; Orr, 1995; Gourbiere and Mallet, 2009; Singhal and Moritz, 2013). Hence, if properly characterized and implemented, this checkpoint should facilitate the delimitation of incipient species.

Palearctic amphibians are a good system to assess the performance of such speciation thresholds for integrative systematics, in respect to intrinsic limitations such as the choice of markers (mtDNA, nuclear) and metrics to apply (% of divergence, divergence time). In am-
phibians, while some influential studies recommended 3% differentiation at sequences of the mitochondrial 16S gene (Malone and Fontenot, 2008; Vieites et al., 2009), recent comparative surveys suggested a divergence time of 3 My to underlie incipient speciation among European anurans. This age corresponds to the youngest candidate species in several well-studied radiations, while younger lineages usually merge upon secondary contact (Dufresnes et al., 2019a and references therein). However, molecular clocks are specific to the phylogenetic framework considered, and the speciation clock may tick at different paces depending on lineages. Empirical data from additional species groups are thus needed to fully appreciate the progress of reproductive isolation in space and time, and refine the thresholds used in systematics.

Here we focused on the midwife toad *Alytes obstetricans*, a small anuran that colonized parts of Western Europe from the Iberian Peninsula. This taxon bears the hallmark of the Plio-Pleistocene climatic fluctuations, and previous phylogeographic analyses identified six deeply-diverged mitochondrial lineages distributed across Central Spain (A), N-Spain/W-Europe (B), Galicia and N-Portugal (C), Central Portugal/Sistema Central mountains in Spain (D), the Spanish Pyrenees (E) and Catalonia (F) (Gonçalves et al., 2015). Most of these clades feature nuclear differentiation, some corresponding to taxonomic subdivisions (Arntzen and García-París, 1995; Martínez-Solano et al., 2004; Gonçalves et al., 2007; Maia-Carvalho et al., 2014, 2018; Gonçalves et al., 2015). The most divergent populations (mtDNA lineages E-F), representing the subspecies *A. o. almogavarii*, are of particular interest. Dated to the Mio-Pliocene (Martínez-Solano et al., 2004) or to the Plio-Pleistocene (Gonçalves et al., 2015), this taxon features the highest allozyme distances to all other *A. obstetricans* subspecies (Nei’s $D = 0.17$; Arntzen and García-París, 1995; see also García-París, 1995). While its placement on the nuclear tree is not well resolved (Gonçalves et al., 2007; Maia-Carvalho et al., 2014), *A. o. almogavarii* forms a strongly differentiated genetic cluster, with little gene flow from parapatric subspecies (based on microsatellites; Maia-Carvalho et al., 2018). The divergence time (2.5 My in Gonçalves et al., 2015) and amount of 16S divergence (0.99-1.38% in Martínez-Solano et al., 2004) are both below the proposed thresholds of speciation, yet the apparent lack of genetic introgression suggests potential reproductive isolation between *A. o. almogavarii* and its congeners, which could indicate an unusually young speciation event.

To elucidate whether *A. o. almogavarii* represents an incipient species, we thoroughly analyzed the transition between *Alytes obstetricans*, lineages F (*almogavarii*) and A (*pertinax*) along the Catalanian coast in northeastern Spain (Gonçalves et al., 2015; Maia-Carvalho et al., 2018), based on transect sampling and genomic resources (RAD-sequencing). Interestingly, this phylogeographic break is also found in a related amphibian, the common parsley frog *Pelodytes punctatus*, where subspecies *P. p. punctatus* and *P. p. hespericus* introgress over hundreds of kilometers (CD and IMS, unpublished data) despite equivalent divergence time (Díaz-Rodríguez et al., 2015, 2017). Assuming similar dispersal capacity in *Alytes* and *Pelodytes* (both share mostly terrestrial habits and are often found in syntopy in Mediterranean habitats), we predict widespread admixture across the *A. o. almogavarii/pertinax* transition if, like *P. p. punctatus/hespericus*, they did not evolve significant reproductive isolation during their short history of divergence. On the contrary, a narrow contact zone with limited introgression is expected if they reached the threshold of divergence above which reproductive isolation might have quickly accumulated.

**Methods**

**Sampling and data generation**

We combined 38 samples from a previous study (Gonçalves et al., 2015) with 73 specimens that were newly-collected during the summer of 2017 along a geographic transect in Catalonia (fig. 1, supplementary table S1). Tissue samples...
Figure 1. Mitochondrial phylogeny of *Alytes obstetricans* (left, adapted from Gonçalves et al., 2015), distribution of the main lineages (inset map, adapted from Gonçalves et al., 2015; grey: unclear), and population genomics (433 SNPs) of the Catalonian hybrid zone (right). On the tree, the attributed subspecies (and species, see discussion) are indicated; note that lineage E segregates in *almogavarii* populations from the Pyrenees, but lacks nuclear differentiation (ghost mtDNA lineage, Maia-Carvalho et al., 2018). The lineages studied here are highlighted in bold. The Catalonian map displays the average nuclear ancestry of each population, based on the STRUCTURE analysis (*K* = 2). The barplots show individual nuclear and mitochondrial assignments. Photo: *Alytes obstetricans* (CD).

The 111 samples (18 populations) were barcoded for mitochondrial DNA using custom-designed primers Aly.ND4.F3 (5′-AAACCCCCTGAAGTTTTTCAGG-3′) and Aly.ND4.R3 (5′-TCTTGGTGGGCAAGAAGGTT-3′), which amplify ∼500 bp of the mitochondrial gene ND4. These samples were then included in a RAD genomic library built with enzymes *SbfI* and *MseI*, following the protocol detailed in Brelsford, Dufresnes and Perrin (2016). The library was sequenced on two Illumina lanes and processed bioinformatically using the STACKS pipeline (v. 1.48), with default parameters (Catchen et al., 2013). We then filtered tags present in all populations (∼p 18), in 80% of individuals of each population (∼r 0.8), but with a minimum allele frequency of 0.05 (∼min_maf 0.05) and a maximum observed heterozygosity of 0.75 (∼max_obs_het 0.75) across all individuals. This retained 433 SNPs that were informative across populations, with little missing data (97.4% of matrix completeness), while limiting paralog overmerging.

Genetic analyses

Patterns of genetic structure among Catalanian *A. o. almo-gavarii* and *A. o. pertinax* samples were assessed from the SNP dataset using the clustering algorithm of STRUCTURE (Pritchard et al., 2000). Five replicates from *K* = 1 to 5 were ran, each including 100,000 iterations after a burnin of 10,000. We also performed a Principal Component Analysis (PCA) on individual allele frequencies with the R package adegenet (Jombart, 2008).

Furthermore, we measured the geographic transition between the two gene pools by fitting sigmoid clines on diagnostic allele frequency data from 17 populations along the transect (all but loc. 5), using the R package hzar (Derryberry et al., 2014). Clines were inferred from three genetic datasets: (1) the average population ancestry (*Q*) obtained by STRUCTURE with *K* = 2 (genome-average cline); (2) the mitotype frequencies (mitochondrial cline); (3) the frequencies of 89 SNPs fixed between taxa (all from different RAD tags), i.e. with *Fst* = 1 between loc. 1-3 (*A. o. almo-gavarii*) and loc. 18 (*A. o. pertinax*). We tested clines from two (width *w* and center *c*) to six parameters (accounting for introgression tails), and reported those best fitting the data according to the AIC criterion. Finally, we estimated intrinsic selection against hybrids *s* (∗the fitness difference between populations from the center and the edge of the contact zone) from the cline width *w*, with respect to the dispersal rate per generation *σ* (p 16 in Barton and Gale, 1993). Since little is known about the dispersal behavior in
Figure 2. Cline analyses across the *almogavarii/pertinax* hybrid zone. (A) Cline fitting from a northeast-southwest transect, separately for the genome average (thick black line; 95% CI in red), mtDNA (thick dash line), and 89 species-diagnostic SNPs (thin grey lines). (B) Selective coefficient $s^*$ calculated from the cline widths, for a range of dispersal rates ($\sigma$). Twelve diagnostic SNPs featured clines with $w \sim 0$ km, 11 with $w = 2-4$ km, while the rest span around the genome average.

midwife toads (Tobler, Garner and Schmidt, 2013), here we inferred $s^*$ for a range of $\sigma$ values from 0.1 to 4 km/generation.

Results

Our genomic data (433 SNPs) perfectly differentiated between the gene pools of *A. o. almogavarii* (coded yellow) and *A. o. pertinax* (coded blue) along the Catalonian coast (fig. 1). The best-clustering solution recovered by STRUCTURE involved two clusters ($K = 2$, $\Delta K = 1121.1$; fig. 1). Accordingly, almost half of the total genetic variance was summarized by the first axis of the PCA and discriminates the two taxa (supplementary fig. S1).

Based on our analyses, these two clusters feature highly restricted genetic introgression. First, STRUCTURE ($K = 2$) identified a single locality with clear nuclear admixture (loc. 14, where haplotypes from both mtDNA lineages are present), while the neighboring loc. 13 (inhabited by *A. o. almogavarii*) features two toads with $Q$ scores below 0.95, and two others carrying *pertinax* mtDNA (fig. 1). All other samples received ancestry coefficients above 0.95 (fig. 1). These results are confirmed by the PCA, on which the few introgressed individuals stand out by intermediate values along the first axis (supplementary fig. S1).

Second, the cline analyses inferred narrow transitions for all markers (fig. 2). Cline models with two parameters (width $w$ and center $c$) were always preferred (lowest AIC scores), and provided remarkably congruent results for mtDNA ($w = 13.5$ km, $c = 86.1$ km), the nuclear genome average ($w = 16.2$ km, $c = 91.1$ km) and the 89 diagnostic nuclear loci ($w = 20.4 \pm 8.5$ km and $c = 90.2 \pm 2.5$ km). For most markers, the transition spanned exclusively between our loc. 13 and 15 (located 30 km apart, fig. 1), near the villages of Valls and Montblanc. While the majority of loci (74%) broadly follow the genome average, two subsets feature narrower clines (13% with $w$ tending towards 0 and 12% with $w = 2-4$ km), as best visualized from the derived $s^*$ values (fig. 2B) and the distribution of cline widths (supplementary fig. S2). Moreover, the hybrid zone may be slightly asymmetric, with the center of the mtDNA cline and of many nuclear SNPs being shifted by a few kilometers on the...
Discussion

The narrow mitochondrial and nuclear transitions suggest that *A. o. pertinax* and *A. o. almogavarii* form a classic tension zone in Catalonia. From the average cline estimates, assuming a dispersal rate of 1.5 km/generation is enough to get $s^* > 0.03$ (fig. 2B), i.e. the values obtained in well-studied hybrid zones involving cryptic anuran species (e.g. *H. arborealorientalis*, Dufresnes et al., 2015; *Pelobates fuscus-vespertinus*, Dufresnes et al., 2019a; *Pelodytes ibericusatlanticus*) (CD and IMS, unpublished data). But it would take up to 3.5 km/generation to reach the strong $s^*$ inferred for deeper-diverged, ecologically differentiated taxa (e.g. $s^* = 0.21$ in *Bombina variegata/bombina*, Barton and Gale, 1993) (fig. 2B). Accounting for a generation time of seven years in *Alytes* (computed from the average age at sexual maturity and lifespan reported by Márquez, Esteban and Castanet, 1997), 1.5-3.5 km/generation thus corresponds to yearly movements of only 0.2-0.5 km.

Such (relatively short) distances seem realistic for *Alytes* in the Catalan landscape. Midwife toads are adapted to xeric environments and accommodate to tiny water points (e.g. droughts), which are visited just a few days per year by the adult males, when they deposit the offspring. Movements up to 1.5 km have been observed in the related *Alytes dickhilleni* endemic to SE-Iberia (Bosch and González Miras, 2012), a region drier than the Catalanian inlands. Note that habitat fragmentation can significantly limit dispersal, even between nearby subpopulations: Tobler, Garner and Schmidt (2013) reported genetic differentiation between sites located <2 km apart in *A. obstetricans* from the heavily impacted Swiss plateau. Here however, a stream network runs along the center of the *almogavarii/pertinax* transition (providing breeding sites, e.g. loc. 14), a landscape element predicted to promote connectivity and persistence among amphibian populations (Grant et al., 2010).

These observations suggest that the limited introgression reflects partial reproductive isolation rather than limited opportunities for dispersal. This is further supported by the heterogeneity of cline widths among our diagnostic markers: the outliers featuring very steep clines (corresponding to strong selective values) could be linked to barrier loci, i.e. involved in hybrid incompatibilities. Theory predicts that most of the genome remains permeable to gene flow in the early stages of incipient speciation, but that post-zygotic isolation arises as barrier loci become recruited and limit introgression across entire genomic regions, due to linkage disequilibrium (Barton, 1983; Orr, 1995; Gourbiere and Mallet, 2009). Surveys across additional *almogavarii/pertinax* contacts would allow testing whether the same clines remain sharp in replicate transects, thus confirming a selective role. Comparative analyses of amphibian hybrid zones shall provide insights on the genomic architecture of introgression at different stages along the speciation continuum, especially to assess how barrier loci accumulate with genetic divergence.

Our results thus call for a taxonomic revision of *Alytes obstetricans*. We hereby consider the Catalanian Midwife Toad *Alytes almogavarii* Arntzen, García-París, 1995 as a distinct incipient species, which was previously considered as a subspecies of the Common Midwife Toad *Alytes obstetricans* (Laurenti, 1768). The biodiversity of the Spanish and French herpetofauna should thus be re-evaluated to account for this regional endemic, especially for conservation planning and red lists. In France, *A. almogavarii* appears restricted to the eastern Pyrenees (genetically confirmed in the departments of Aude and Pyrénées-Orientales, and perhaps extending to Ariège; Gonçalves et al., 2015; Maia-Carvalho et al., 2018), so it is extremely localized in the country.
Are *Alytes obstetricans* and *A. almogavarii* cryptic species? In the original description, Arntzen and García-París (1995) compared external characters and found high morphological similarity between all Iberian *Alytes*, with the exception of the early-diverged *A. cisternasii* and the habitat-specialist *A. muletensis* (see also Sanchiz, 1984). Based on skull osteology however, *A. almogavarii* shares some character states with species in subgenus *Baleaphryne* (including *A. muletensis*, *A. maurus*, and *A. dickhilleni*) rather than with the other *A. obstetricans* subspecies (Martínez-Solano et al., 2004). Furthermore, bioacoustic investigations suggested inter-population variation in breeding calls between Iberian taxa, including longer notes in *A. almogavarii* compared to *A. obstetricans* subspecies (Márquez and Bosch, 1995).

To our knowledge, the tadpoles of the different subspecies have not been formally compared. We did observe lighter and more contrasted coloration in tadpoles of *A. almogavarii* than in those of *A. o. pertinax* during the fieldwork of this study (IMS & CD, pers. obs.), but this could simply relate to their micro-habitats (streams vs still waters), which are known to influence coloration in *Alytes* (Polo-Cavia et al., 2016). The new taxonomic status of *A. almogavarii* should hopefully motivate comparative assessments, especially to document whether the differing breeding calls may induce some pre-mating isolation with *A. obstetricans*, and thus contribute to the limited hybridization observed.

The tension zone between *A. obstetricans* and *A. almogavarii* sharply contrasts with the wide hybrid zone (>150 km) found for the parsley frogs *Pelodytes punctatus punctatus/hespericus* in Catalonia (Díaz-Rodríguez et al., 2017; CD and IMS, unpublished data), despite putatively similar divergences (~2.5 My for *Alytes*; ~2.8 My for *Pelodytes*). While massive introgression is the rule rather than the exception for Plio-Pleistocene anuran lineages (e.g. Dufresnes et al., 2019a), the opposite patterns between *Alytes* and *Pelodytes* emphasize that the outcomes of secondary contacts are hard to predict for pairs of taxa falling within the grey zone of speciation. First, admixture across hybrid zones depends on a myriad of extrinsic factors, and may thus drastically vary between replicate contacts (e.g. Croucher et al., 2007). For instance, asymmetric introgression across the genome may arise due to hybrid zone movement, a situation increasingly reported in the wild (e.g. van Riemsdijk et al., 2019). Here, some clines appear slightly shifted on the *almogavarii* side (supplementary fig. S2), but it could also be a methodological bias due to our ~20 km sampling gap on the *pertinax* side of the contact (between loc. 14 and 15). Second, the random accumulation of genetic incompatibilities can lead to different levels of postzygotic isolation between independent pairs of lineages, even if they share equivalent divergences (e.g. Arntzen et al., 2017). The latter could be directly assessed in *Alytes* by investigating the transition between *A. almogavarii* and *A. o. obstetricans* (the sister clade of *A. o. pertinax*), most likely extending along the French Pyrenees (Gonçalves et al., 2015). More generally, these aspects are crucial to consider when applying thresholds of speciation: here *A. almogavarii* falls below both the sequence and time thresholds set up by previous studies (see Introduction), which might explain why it was not previously considered a distinct species. Although thresholds are useful for species delimitation, especially for completely allopatric taxa (e.g. insular species), direct evidences from targeted studies of secondary contacts thus appear essential to make the best informed taxonomic revisions.

Alternatively, in this case, it is conceivable that the mitochondrial inferences (Gonçalves et al., 2015) misplaced *A. almogavarii* in the phylogeny and underestimated its evolutionary age. Relying on transformed-branch length estimates, Martínez-Solano et al. (2004) rather advocated for a Mio-Pliocene split for this clade. Moreover, some nuclear gene trees group *A. almogavarii* with the *Baleaphryne* subgenus (Gonçalves et al., 2007; but see Maia-Carvalho et al., 2014), a topology consistent with their os...
teological similarities (Martínez-Solano et al., 2004). Could A. almogavarri be much older than previously assumed, but lost its mtDNA during Plio-Pleistocene episodes of hybridization? Complete mitochondrial replacement appears to be common in Palearctic amphibians (e.g. Zieliński et al., 2013), eventually concealing speciation events (the super-cryptic species concept, Dufresnes et al., 2019b). Phylogenomics of the entire Atyes genus should hopefully clarify this situation and bring complementary insights on the evolutionary history of A. almogavarri. Finally, resolving the complex taxonomy of this group will benefit from genomic analyses targeting additional transitions among the polytypic A. obstetricans, and notably within the genetically-structured A. o. obstetricans (Maia-Carvalho et al., 2018).

Acknowledgements. We thank Generalitat de Catalunya for providing tissue collection permits, Agents de l’Alt Penedès del Cos d’Agents Rurals, P. Albornó, V. Bros, A. Escayol, G. Giner, R. Jiménez, C. Luque, D. Martínez, A. Montori, R. Ruiz and D. Villero for their help in the field, as well as B. Álvarez and I. Rey from the DNA and Tissue collection of the MNCN-CSIC for providing samples under their care. We are grateful to N. Perrin for support. This study was funded by the Swiss National Science Foundation (fellowship P2LAP3_171818 to CD, and grant 31003A_166323 to N. Perrin).

Supplementary material. Supplementary material is available online at: https://doi.org/10.6084/m9.figshare.10075025

References

Arntzen, J.W., García-París, M. (1995): Morphological and allozyme studies of midwife toads (genus Alytes), including the description of two new taxa from Spain. Contrib. Zool. 65: 5-34.

Arntzen, J.W., de Vries, W., Canestrilli, D., Martínez-Solano, I. (2017): Hybrid zone formation and contrasting outcomes of secondary contact over transects in common toads. Mol. Ecol. 26: 5663-5675.

Avise, J. (2000): Phylogeography: the History and Formation of Species. Harvard University Press, Cambridge (MA).

Barton, N., Gale, K.S. (1993): Genetic analysis of hybrid zones. In: Hybrid Zones and the Evolutionary Process, p. 13-45. Harrison, R., Ed., Oxford University Press, New York.

Barton, N.H. (1983): Multilocus clines. Evolution 37: 454-471.

Bosch, J., González-Miras, E. (2012): Seguimiento de Alytes dickhilleni: informe final. In: Monografías SARE, 02. Asociación Herpetológica Española y Ministerio de Agricultura, Alimentación y Medio Ambiente, Madrid.

Brelsford, A., Dufresnes, C., Perrin, N. (2016): High-density sex-specific linkage maps of a European tree frog (Hyla arborea) identify the sex chromosome without information on offspring sex. Heredity 116: 177-181.

Catchen, J., Hohenlohe, P., Bassham, S., Amores, A., Cresko, W. (2013): Stacks: an analysis tool set for population genomics. Mol. Ecol. 22: 3124-3140.

Chenuil, A., Cahiill, A.E., Délemontey, N., Du Salliant, E., Fanton, H. (2019): Problems and questions posed by cryptic species. A framework to guide future studies. In: From Assessing to Conserving Biodiversity. History, Philosophy and Theory of the Life Sciences, vol. 24, p. 77-106. Casetta, E., Marques da Silva, J., Vecchi, D., Eds, Springer, Cham.

Coyne, J.A., Orr, H.A. (2004): Speciation. Sinauer Associates Inc, Sunderland.

Croucher, P.J., Jones, R.R., Searle, J.B., Oxford, G.S. (2007): Contrasting patterns of hybridization in large house spiders (Tegenaria atrica group, Agelenidae). Evolution 61: 1622-1640.

De Queiroz, K. (2007): Species concepts and species delimitation. Syst. Biol. 56: 879-886.

Díaz-Rodríguez, J., Gonçalves, H., Sequeira, F., Sousa-Neves, T., Tejedo, M., Ferrand, N., Martínez-Solano, I. (2015): Molecular evidence for cryptic candidate species in Iberian Pelodytes (Anura, Pelodytidae). Mol. Phylogenet. Evol. 83: 224-241.

Díaz-Rodríguez, J., Gehara, M., Márquez, R., Vences, M., Gonçalves, H., Sequeira, F., Martínez-Solano, I., Tejedo, M. (2017): Integration of molecular, bioacoustical and morphological data reveals two new cryptic species of Pelodytes (Anura, Pelodytidae) from the Iberian Peninsula. Zootaxa 4243: 1-41.

Dufresnes, C., Brelsford, A., Cnrbomija-Isailovici, J., Tzankov, N., Lymbrekisk, P., Perrin, N. (2015): Time-frame of speciation inferred from secondary contact zones in the European tree frog radiation (Hyla arborea group). BMC Evol. Biol. 15: 155.

Dufresnes, C., Strachinis, I., Siriadni, N., Mykhytytnets, G., Cogălniceneanu, D., Székely, P., Vukov, T., Arntzen, J.W., Wielstra, B., Lymbrikisk, P., Geffen, E., Gafny, S., Kumluataş, Y., Ilgaz, Ç., Candan, K., Mizsei, E., Szabolcs, M., Kolenda, K., Smirnov, N., Génieus, P., Lukanov, S., Crochet, P.-A., Dubey, S., Perrin, N., Litvinchuk, S.N., De-noël, M. (2019a): Phylogeography of a cryptic speciation continuum in Eurasian spadefoot toads (Pelobates). Mol. Ecol. 28: 3257-3270.

Dufresnes, C., Mazeppa, G., Jablonski, D, Caliari Oliveira, R., Wenseleers, T., Shabanov, D.A., Auer, M., Ernst, R., Koch, C., Ramírez-Chaves, H.E., Melder, K.P., Simonov, E., Tuitenko, A., Kryvokhyzha, D., Wennekes, P.L., Zinenko, O.I., Korshunov, O.V., Al-Johany, A.M., Peregontsev, E.A., Masroor, R., Betto-Colliard, C., De-noël, M., Borkin, I., Skorinov, D.V., Panynkova, R.A., et al. (2019b): High-density sex-specific linkage maps of a European tree frog (Hyla arborea) identify the sex chromosome without information on offspring sex. Heredity 116: 177-181.
Mazanaeva, L.F., Rosanov, J.M., Dubey, S., Litvinchuk, S.N. (2019b): Fifteen shades of green: the evolution of _Bufo_ toads revisited. Mol. Phylogenet. Evol. 141: 106615.

Fouquet, A., Gilles, A., Vences, M., Marty, C., Blanc, M., Gemmell, N.J. (2007): Underestimation of species richness in Neotropical frogs revealed by mtDNA analyses. PLoS ONE 2: e1109.

García-París, M. (1995): Variabilidad genética y distribución geográfica de _Alytes obstetricans almogavarii_ en España. Rev. Esp. Herp. 9: 133-138.

Gonçalves, H., Martínez-Solano, I., Ferrand, N., García-París, M. (2007): Multilocus phylogeography of the common midwife toad, _Alytes obstetricans_ (Anura, Alytidae): contrasting patterns of lineage diversification and genetic structure in the Iberian refugium. Mol. Phylogenetic. Evol. 93: 363-379.

Gourbiere, S., Mallet, J. (2009): Are species real? The shape of the species boundary with exponential failure, reinforcement and the ‘missing snowball’. Evolution 64: 1-24.

Grant, E.H.C., Nichols, J.D., Lowe, W.H., Fagan, W.F. (2010): Use of multiple dispersal pathways facilitates amphibian persistence in stream networks. Proc. Natl Acad. Sci. USA 107: 6936-6940.

Heethoff, M. (2018): Cryptic species – conceptual or terminological chaos? A response to Struck et al. Trends Ecol. Evol. 33: 310.

Jombart, T. (2008): adegenet: a R package for the multivariate analysis of genetic markers. Bioinformatics 24: 1403-1405.

Lee, M.S.Y. (2004): The molecularisation of taxonomy. Invertebr. Syst. 18: 1-6.

Maia-Carvalho, B., Gonçalves, H., Ferrand, N., Martínez-Solano, I. (2014): Multilocus assessment of phylogenetic relationships in _Alytes_ (Anura, Alytidae). Mol. Phylogenetic. Evol. 79: 270-278.

Maia-Carvalho, B., Gomes Vale, C., Sequeira, F., Ferrand, N., Martínez-Solano, I., Gonçalves, H. (2018): The roles of allopatric fragmentation and niche divergence in intraspecific lineage diversification in the common midwife toad (_Alytes obstetricans_). J. Biogeogr. 45: 2146-2158.

Malone, J.H., Fontenot, B.E. (2008): Patterns of reproductive isolation in toads. PLoS ONE 3: e3900.

Márquez, R., Bosch, J. (1995): Advertisement calls of the midwife toads _Alytes_ (Anura, Discoglossidae) in continental Spain. J. Zool. Syst. Evol. Res. 33: 185-192.

Márquez, R., Esteban, M., Castanet, J. (1997): Sexual size dimorphism and age in the midwife toads _Alytes obstetricans_ and _A. cisternasi_. J. Herpetol. 1: 52-59.

Martínez-Solano, I., Gonçalves, H.A., Arntzen, J.W., García-París, M. (2004): Phylogenetic relationships and biogeography of midwife toads (Discoglossidae: _Alytes_). J. Biogeogr. 31: 603-618.

Orr, H. (1995): The population genetics of speciation: the evolution of hybrid incompatibilities. Genetics 139: 1805-1813.

Padial, J.M., Miralles, A., de la Riva, I., Vences, M. (2010): The integrative future of taxonomy. Front. Zool. 7: 16.

Polo-Cavia, N., Oliveira, J.M., Redondo Villa, A.J., Márquez, R. (2016): Background colour matching in a wild population of _Alytes obstetricans_. Amphibia-Reptilia 37: 253-260.

Pritchard, J.K., Stephens, M., Donnelly, P. (2000): Inference of population structure using multilocus genotype data. Genetics 155: 945-959.

Roux, C., Fräisse, C., Romiguier, J., Anciaux, Y., Galtier, N., Bierne, N. (2016): Shedding light on the grey zone of speciation along a continuum of genomic divergence. PLoS Biol. 14: e20000234.

Sanchiz, B. (1984): Analisis filogenético de la tribu Alytini (Anura, Alytidae) mediante el estudio de su morfoestructura ósea. In: Historia Biológica del Ferreret (Baleaphryne muletensis), p. 61-108. Hemmer, H., Alcover, J.A., Eds, Ed. Moll., Palma de Mallorca.

Singhal, S., Moritz, C. (2013): Reproductive isolation between phylogeographic lineages scales with divergence. Proc. R. Soc. B. 280: 20132246.

Struck, T.H., Feder, J.L., Bendiksby, M., Birkeland, S., Cerca, J., Gusarov, V.I., Kistenich, S., Larsson, K.H., Liow, L.H., Nowak, M.D., Stedje, B., Bachmann, L., Dimitrov, D. (2018): Finding evolutionary processes hidden in cryptic species. Trends Ecol. Evol. 33: 153-163.

Tobler, U., Garner, T.W.J., Schmidt, B.R. (2013): Genetic attributes of midwife toad (_Alytes obstetricans_) populations do not correlate with degree of species decline. Ecol. Evol. 3: 2806-2819.

van Rijmsdijk, I., BUTLIN, R.K., WIELSTRA, B., ARNTZEN, J.W. (2019): Testing an hypothesis of hybrid zone movement for toads in France. Mol. Ecol. 28: 1070-1083.

Vieites, D.R., Wollenberg, K.C., Andréone, F., Köhler, J., Glaw, F., Vences, M. (2009): Vast underestimation of Madagascar’s biodiversity evidenced by an integrative amphibian inventory. Proc. Natl. Acad. Sci. USA 106: 8267-8272.

Zielinski, P., Nadachowska-Brzyska, K., Wielstra, B., Szotak, R., Covaciuc-Marcov, S.D., Cogălniceanu, D., Babik, W. (2013): No evidence for nuclear introgression despite complete mtDNA replacement in the Carpathian newt (Lissotrition montandoni). Mol. Ecol. 22: 1884-1903.

Submitted: October 7, 2019. Final revision received: October 28, 2019. Accepted: October 28, 2019. Associate Editor: Judit Vörös.