RESEARCH ARTICLE

Are invasive marsh frogs (*Pelophylax ridibundus*) replacing the native *P. lessonae/P. esculentus* hybridogenetic complex in Western Europe? Genetic evidence from a field study

Julien Leuenberger · Antoine Gander · Benedikt R. Schmidt · Nicolas Perrin

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Abstract The water-frog L–E system, widespread in Western Europe, comprises the pool frog Pelophylax lessonae and the hybridogenetic edible frog P. esculentus, which originated from hybridization between pool frogs and marsh frogs (P. ridibundus). In P. esculentus, the lessonae (L) genome is eliminated during meiosis and has to be gained anew each generation from a P. lessonae partner, while the *ridibundus* (R') genome is transmitted clonally. It therefore accumulates deleterious mutations, so that R'R' offspring from P. esculentus \times P. esculentus crosses are normally unviable. This system is now threatened by invasive P. ridibundus (RR) imported from Eastern Europe and the Balkans. We investigated the genetic interactions between invasive marsh frogs and native water frogs in a Swiss wetland area, and used genetic data collected in the field to validate several

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J. Leuenberger \cdot N. Perrin (\boxtimes)

Department of Ecology and Evolution, University of Lausanne, 1015 Lausanne, Switzerland e-mail: nicolas.perrin@unil.ch

A. Gander

Association de la Grande Cariçaie, Chemin de la Cariçaie 3, 1400 Cheseaux-Noréaz, Switzerland

B. R. Schmidt

Institute of Evolutionary Biology and Environmental Studies, University of Zurich, Winterthurerstrasse 190, 8057 Zurich, Switzerland

B. R. Schmidt

KARCH, Passage Maximilien-de-Meuron 6, 2000 Neuchâtel, Switzerland

components of a recently postulated mechanism of species replacement. We identified neo-ridibundus individuals derived from crosses between invasive ridibundus and native esculentus, as well as newly arisen hybridogenetic esculentus lineages stemming from crosses between invasive ridibundus (RR) and native lessonae (LL). As their ridibundus genomes are likely to carry less deleterious mutations, such lineages are expected to produce viable ridibundus offspring, contributing to species replacement. However, such crosses with invasive ridibundus only occurred at a limited scale; moreover, RR×LL crosses did not induce any introgression from the ridibundus to the lessonae genome. We did not find any ridibundus stemming from crosses between ancient esculentus lineages. Despite several decades of presence on the site, introduced ridibundus individuals only represent 15 % of sampled frogs, and their spatial distribution seems shaped by specific ecological requirements rather than history of colonization. We therefore expect the three taxa to coexist stably in this area.

Keywords Amphibian · Hybridogenesis · Invasion · Microsatellites · Water frog

Introduction

Accidental or deliberate introductions of exotic species into non-native ranges are now occurring at unprecedented rates, partly due to the intensive development of international trade (Westphal et al. 2008). Some introduced species become invasive, which poses important threats to local biodiversity (Wilcove et al. 1998; Vorburger and Ribi 1999). Potential impacts on native species are both ecological and genetic (Williamson and Fitter 1966; Manchester and Bullock 2000).



Ecological effects are mediated by competition for food or space, predation, parasitism, habitat modification, or spread of new diseases (Mooney and Cleland 2001). Genetic effects result from hybridization with indigenous species, such as documented between wild and domestic cats, or between introduced Mallards and native ducks (Rhymer and Simberloff 1996). If hybrids are fertile, backcrosses with parental species induce genetic introgression that may lead to the progressive elimination of native genomes ("genetic swamping"; Rhymer and Simberloff 1996; Huxel 1999), as documented e.g. between introduced and local trout in the USA, or between Red deer and introduced Sika deer in Scotland (Rhymer and Simberloff 1996).

Water frogs from the Pelophylax esculentus complex represent a particular situation. The so-called klepto-taxon P. esculentus (edible frog) is a permanent F_1 hybrid that originally emerged from a hybridization between the pool frog P. lessonae and the marsh frog P. ridibundus (Graf and Polls Pelaz 1989). Due to still unknown features of the ancestral ridibundus (R) genome, the lessonae (L) genome is eliminated from the esculentus germ line, so that all gametes transmit the same, non-recombined ridibundus (R') genome (Tunner and Heppich 1981). Mating with a lessonae (LL) partner is thus required to restore esculentus (R'L) offspring—a process called hybridogenesis. Being always expressed in hemizygous state, R' hemiclones are largely protected from purifying selection; in addition, the strong Hill-Robertson interferences resulting from clonal transmission favor the rapid accumulation of deleterious mutations. As a result, crosses between two esculentus bearing the same hemiclone normally produce unviable or unfertile R'R' offspring (Vorburger 2001a), although evidence exists that these newly formed P. ridibundus can occasionally survive and even mature, provided the combining R' have fixed distinct deleterious mutations (Hotz et al. 1985, 1992; Berger et al. 1988; Schmidt 1993; Vorburger 2001b; Guex et al. 2002; Schmeller et al. 2005; Luquet et al. 2011).

The so-called L–E system, consisting of syntopic populations of *P. lessonae* and its obligate genetic parasite *P. esculentus*, is widespread throughout Western and Central Europe, where *P. ridibundus* (the other parental species) is absent (Uzzell and Berger 1975). This system is now threatened by the ongoing invasion of marsh frogs; native to Eastern Europe, the Balkanic peninsula and Western Asia, *P. ridibundus* has been successfully introduced in Central and Western Europe for commercial purpose (Kuzmin et al. 2009). Its expansion has drastic consequences on native amphibian species via competitive exclusion and direct predation. Native pool frogs have already been replaced at several sites (Schmeller et al. 2007; Holsbeek and Jooris 2009). The Bois de Finges, a nature reserve in western Switzerland originally occupied

by the L–E system, was colonized in the 1960s by *P. ridibundus* (Marchesi et al. 1999). The numbers of marsh frogs have increased until in 1999 *P. lessonae* represented only 3 % of water frogs (Marchesi et al. 1999), and has become completely extinct by 2013 (Leuenberger, personal observation).

Replacement of native pool frogs may occur through both ecological and genetic interactions. The latter have been suggested to take place via a mechanism inherent to the hybridogenetic nature of this hybrid complex (Vorburger and Reyer 2003). On one hand, mating between invasive *ridibundus* (RR) and native *esculentus* (R'L) is expected to produce viable *ridibundus* (RR') offspring. On the other hand, mating between *ridibundus* (RR) and *lessonae* (LL) generates *esculentus* (RL) primary hybrids; if the latter are hybridogenetic, mating between RL hybrids should restore pure and viable *ridibundus* (RR) offspring. Thus, the *ridibundus* genome is expected to progressively replace the *lessonae* genome, and eventually eliminate the L–E complex (Vorburger and Reyer 2003).

Several assumptions underlying this genetic mechanism have been tested and validated in the laboratory via experimental crosses (Vorburger and Reyer 2003). These crosses furthermore detected primary hybrid females that were non-hybridogenetic (see also Hotz et al. 1985), producing offspring with recombined ridibundus and lessonae genomes; this suggests an additional (and more classical) mechanism of genetic replacement in which the *ridibundus* genome would replace the lessonae genome by progressive introgression (Rhymer and Simberloff 1996). It is not known, however, whether the different processes mentioned above actually take place in nature, and if they do, how frequently they occur. It is also possible that behavioral or ecological preferences may result in assortative mating at such a high rate that complete genome replacement is prevented and long-term coexistence possible (Plénet et al. 2001; Luquet et al. 2011).

The present study was aimed at investigating the genetic mechanisms of species replacement in a region originally occupied by the L-E system, where invading P. ridibundus, of unknown origin, have been observed for more than three decades (M. Antoniazza pers. comm.; Berthoud and Perret-Gentil 1976). We thus expected the process of replacement to be well advanced, possibly with lessonae and/or esculentus already eliminated locally. We based our investigations on nuclear and mitochondrial markers for several reasons: (i) there is a large overlap between the esculentus phenotype and both parental species, so that species identity cannot be diagnosed with certainty based on morphology alone (Nöllert and Nöllert 2003; Meyer et al. 2009; Holsbeek and Jooris 2009); (ii) genetic markers are expected to provide relevant information on the gene diversity of invading ridibundus and native lessonae, on



Table 1 Sampling sites and sample numbers of *Pelophylax* spp. along the southern shore of Lake Neuchâtel (western Switzerland)

Sampling	Coordinates	P. lessonae			P. esculentus			P. ridibundus		
site		M	F	J	M	F	J	M	F	J
Chevroux	06°54′27″E; 46°53′42″N	0	0	1	0	9	1	4	8	3
Gletterens 1	06°56′14″E; 46°54′32″N	3	2	0	17	4	2	0	0	0
Gletterens 2	06°56′20″E; 46°54′44″N	8	6	0	7	5	0	0	0	0
Motte 1	06°59′06″E; 46°56′30″N	2	5	0	5	13	0	1	0	0
Motte 2	06°58′50″E; 46°56′23″N	5	5	1	11	5	8	0	0	0

M male, F female, J juvenile

their possible introgression, as well as on the diversity of R' hemiclones in ancient esculentus lineages; (iii) nuclear markers offer the potential to differentiate the native hemiclonal R' genomes from the invading R genomes, thereby allowing identification of esculentus primary hybrids resulting from ridibundus (RR)×lessonae (LL) crosses, as well as neo-ridibundus resulting from ridibundus (RR)×esculentus (LR') or esculentus (LR')×esculentus (LR') crosses, respectively; (iv) nuclear markers offer the potential to identify possible polyploid esculentus, which would complicate the analysis further; (v) mitochondrial information might contribute to identify primary hybrids and neo-ridibundus individuals, and deliver information on their paternal versus maternal ancestry. It has been suggested (e.g. Luquet et al. 2011) that introgression of lessonae mtDNA into the ridibundus genome (e.g. via mating of female esculentus with male ridibundus; Spolsky and Uzzell 1984) may facilitate the local adaptation of invasive ridibundus.

Materials and methods

Field sampling and laboratory work

Water frogs were collected from five sites (Table 1) in a nature reserve (la Grande Cariçaie) along the southern shore of the Lake of Neuchâtel (Switzerland) during the 2012 breeding season (May to early August). Individuals were sampled for buccal cells (two sterile cotton swabs per individual; Broquet et al. 2007), then released at the place of capture. Swabs were stored dry at -20 °C before analysis. DNA was extracted with the Qiagen DNeasy Blood and Tissue Kit, using the Spin-Column Protocol (manual

extraction) and the BioSprint 96 workstation (Qiagen, Valencia; PA; USA), following the manufacturer's protocols with overnight sample incubation in proteinase K at 56 °C.

DNA samples were genotyped for a set of 18 microsatellite markers selected for their potential to differentiate the two genomes under study (Table 2), five of them considered specific to the R genome, five other to the L genomes, and the last eight markers expected to amplify in both (Zeisset et al. 2000; Garner et al. 2000; Hotz et al. 2001; Arioli 2007; Christiansen 2009; Christiansen and Reyer 2009). Polymerase chain reactions (PCRs) were performed in four multiplex PCR containing primers for 4–5 loci (Table 2). Reactions were performed in 10 μl, containing 3 µl DNA extraction, 3 µl Qiagen Multiplex PCR Master mix, and from 2.2 to 3 µl of primer mixes with labeled forward and reverse primers (Table 2). The thermal profile of PCR amplification included an initial denaturation step of 15 min at 95 °C followed by 35 cycles of 30 s denaturation at 94 °C, 90 s annealing at T_a (Table 2), 60 s extension at 72 °C, and a final extension of 30 min at 60 °C. PCR products derived from primer mix 1 were diluted five times and from all other primer mixes twice. Of the diluted PCR product, 3 µl were run on an ABI 3130xl Genetic Analyzer (Applied Biosystems, Life Technologies Corporation, Carlsbad, CA) with an internal size standard (GeneScan-350 ROX). Alleles were scored with GENEM-APPER v4.0 (Applied Biosystems).

DNA samples were also analyzed for the mitochondrial cytb gene, using the primers F1: 5'CGA AGC CTA GAA GAT CTT TG3' and L1: 5'CTC CTG GGA GTC TGC CTA AT3' (Dubey et al. in prep.). PCRs were performed in a total volume of 25 μl with 2.5 μl of 10× buffer, 0.2 mM of each primer, 0.05 mM of dNTPs, 8 mM of MgCl2, 1 U of Taq DNA polymerase (Qiagen, Valencia, PA, USA) and 2 μl of extracted DNA, under the following conditions: initial denaturation step of 5 min at 94 °C, followed by 35 cycles of 30 s denaturation at 94 °C, 40 s annealing at 50 °C, 90 s extension at 72 °C and a final extension of 10 min at 72 °C. PCR fragments were then purified and sequenced at GATC sequencing service (Cologne, D; http://www.gatc-biotech.com). Sequences were aligned by ClustalW (Higgins et al. 1994) implemented in MEGA v5.2 (Tamura et al. 2011), together with the published sequences of two P. lessonae from central and southern Italy (GenBank accession numbers EU047775 and AB029942), two P. ridibundus from Ukraine and Greece (AB029945 and DQ474163), and one P. nigromaculatus as an outgroup (AB043889). We used MrAIC v1.4.4 (Nylander 2004) to select the best model of DNA substitution for Maximum likelihood (ML) analyses. The HKY+G model best fitted our dataset according to AIC, AICc and BIC criteria. ML analyses were performed with PhyML v3.0.1 (Guindon and



Table 2 Amplification
conditions for the 18
microsatellite markers used,
distributed in four primers
mixes

Provided are for each marker the genome specificity (with published reference: *I* Arioli 2007, *2* Christiansen 2009, *3* Garner et al. 2000; *4*

Christiansen and Reyer 2009, 5 Zeisset et al. 2000; 6 Hotz et al. 2001), labels of forward primers, volume of primers used, and annealing temperature * Difference in specificity between our samples and the references (see "Results"

Primers mix	Marker	Genome specificity as published	References	Forward primers labels	Volume (µl) of each primers (10 mM)	Annealing temperature (°C)
Primers mix 1	Ga1a19 redesigned	L+R	1, 2	FAM	0.4	58
	RlCA1b5	L+R	3	ATTO550	0.2	58
	RICA5	L	3	HEX	0.3	58
	RlCA1b6	L+R	1	FAM	0.2	58
	Rrid064A	R	4	Dyomics630	0.4	58
Primers mix 2	Re2Caga3	R	1	ATTO550	0.1	55
	Res16	L+R	5	FAM	0.2	55
	Res20	L	5	HEX	0.5	55
	RlCA2a34	L+R	4	Dyomics630	0.5	55
Primers mix 3	ReGA1a23	L*	4	ATTO550	0.2	58
	Rrid169A	R	4	HEX	0.4	58
	Res22	R	5	FAM	0.2	58
	Rrid013A	L+R	6	FAM	0.1	58
	Rrid059A	L+R	6	ATTO550	0.2	58
Primers mix 4	Re1Caga10	L+R	1	HEX	0.4	58
	RlCA1a27	L	4	Dyomics630	0.4	58
	Rrid135A	R*	4	ATTO550	0.3	58
	RICA18	L	3	FAM	0.2	58

Gascuel 2003) implemented in SeaView v4.0 (Gouy et al. 2010).

Statistical analyses

section)

Genotypes were clustered using a Bayesian algorithm (Structure v.2.3.3; Pritchard et al. 2000). The most likely number of clusters (K) was calculated from the Δ K method (Evanno et al. 2005) implemented in Structure Harvester v0.6.93 (Earl and vonHoldt 2012). Analyses were carried out on the 18 nuclear loci using an admixture model without prior species information, with 10,000 burn-in steps and 100,000 iterations. All runs were replicated 10 times with K ranging from 1 to 8.

In parallel, we performed principal component analyses (PCAs) with PCAGEN v2.0 (Goudet 2005), which extracts the factors (linear component of the 18 initial variables) that maximize overall differentiation (F_{ST}). A first PCA was applied on individual genotypes, and a second after dissociation of the L and R' genomes in *esculentus* individuals (see Results); each *esculentus* individual was thus represented by two entries in the matrix, one expressing its *lessonae* genome and the other its *ridibundus* genome.

Gene diversity (i.e., expected heterozygosity under HW assumptions, given allelic frequencies) was calculated with FSTAT v2.9.3 (Goudet 1995), based on the 10 markers that amplified products from both genomes (see Results), and independently so in the R' and L genomes of *esculentus* individuals. Differences in gene diversity between genomes

were tested with a Mann–Whitney U test in R (R Core Team 2012).

Results

Species identification

A total of 141 water frogs (adults and juveniles) were captured over 27 nights of sampling (Table 1). All 141 individuals could readily be assigned to three genetic clusters based on the patterns of microsatellite amplification; 16 individuals (five males, eight females, three nonsexed juveniles) yielded no amplicons for four markers (RlCA5, Res20, RlCA1a27 and RlCa18) considered specific to the lessonae genome and therefore were assigned to P. ridibundus. Another 38 individuals (18 males, 18 females, two non-assigned individuals) showed no amplification for four other markers (Rrid064A, Re2Caga3, *Rrid*169A and *Res*22) considered specific to the *ridibundus* genome and therefore were assigned to P. lessonae. The remaining 87 individuals (39 males, 36 females, 12 nonassigned) amplified for all 18 markers, and were hemizygous for the eight markers mentioned above; these were therefore assigned to P. esculentus. Contrasting to the results by Christiansen and Reyer (2009), ReGA1a23 and Rrid135A were not strictly specific to the lessonae and ridibundus genomes respectively: the former yielded amplicons in many ridibundus genomes, and the latter in a



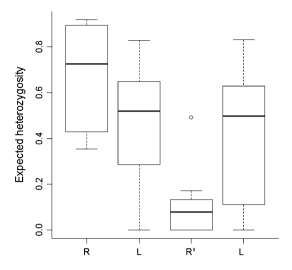


Fig. 1 Gene diversity of the R and L genomes from the two parental species (left; R = P. ridibundus, L = P. lessonae) and the R' and L components of the P. esculentus genomes (right). Boxes represent interquartile ranges (50 % of the data) with the median given by the $bold\ line$. Whiskers identify values that do not exceed 1.5 times the interquartile range; the outlier ($empty\ dot$) is Re1Caga10

few *lessonae* genomes. Despite overall high levels of gene diversity, no individual displayed more than two alleles at any locus, pointing to all-diploid populations.

Gene diversity

The hybridogenetic nature of these 87 *P. esculentus* individuals was corroborated by allelic patterns: for the 10 loci in which two alleles could be amplified, one copy matched

the size range and frequency distribution of *lessonae* alleles, while the other copy matched the *ridibundus* size range and was mostly monomorphic throughout. These patterns allowed straightforward identification of the L and R' genomes in all *esculentus* individuals, and therefore independent estimation of gene diversity on both components (Fig. 1). Expected heterozygosity was highest in the R genomes of *P. ridibundus*, and lowest in the R' genomes of *P. esculentus* (Mann–Whitney U test, U = 97, p < 0.001). The L genomes from *P. lessonae* and *P. esculentus* did not differ from each other (U = 51, p = 0.970), being slightly less diverse than the *P. ridibundus* R genomes (U = 28, p = 0.104).

Mitochondrial DNA

Species assignments were largely corroborated by mtDNA analyzes: all 38 individuals assigned to *P. lessonae* clustered in the same *cytb* clade, together with *lessonae* sequences from Italy, while all 16 *P. ridibundus* with one exception (3_EN_13) clustered in a distinct clade, together with *P. ridibundus* Genbank sequences (Fig. 2). Among the 87 *P. esculentus* individuals identified, 84 displayed a *lessonae* haplotype, while the remaining three (1_EN_40, 3_EN_02 and 2_EN_19) had a *ridibundus* haplotype.

Clustering of individuals

Structure analyses identified K=3 as the most likely number of clusters. All individuals were assigned with high

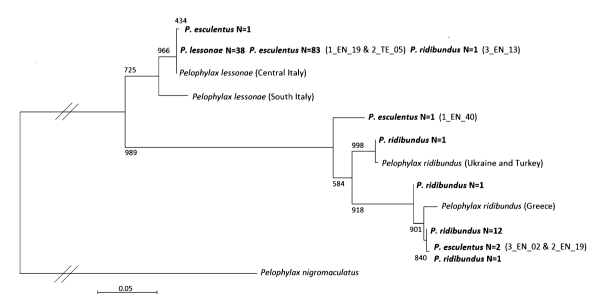


Fig. 2 *cyt-b* maximum-likelihood phylogeny of water frogs from our sampling site, with one outgroup (*P. nigromaculatus*, GenBank accession number AB043889) and four reference genomes (two *P. ridibundus* and two *P. lessonae*, respectively AB029945, DQ474163, EU047775 and AB029942). The names in *bold* correspond to the

samples from our study, identified from the patterns of microsatellites amplification. Provided are bootstrap support values for the major branches (out of 1,000 replicates), as well as localization of the six outliers (one *ridibundus*, five *esculentus*)



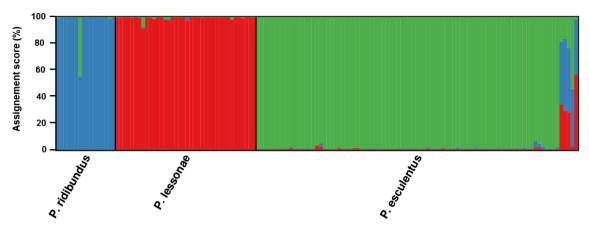


Fig. 3 Bayesian clustering analysis (STRUCTURE) for the most likely number of clusters (K=3), with no prior information on taxonomic identity. All 141 individuals are assigned with high

likelihood to either the *blue* (*ridibundus*), *red* (*lessonae*) or *green* (*esculentus*) cluster, with however six exceptions (one *ridibundus* and five *esculentus*)

likelihoods to one of these three clusters each, fully consistent with the patterns of marker amplification, however with six exceptions (Fig. 3): the *ridibundus* individual with a cytonuclear mismatch (3_EN_13) was assigned to a mix of *ridibundus* and *esculentus* with approximately equal probability, and five *esculentus* (from left to right in Fig. 3: 1_EN_40, 3_EN_02, 2_EN_19, 2_TE_05, and 1_EN_29) were assigned to a mix of *ridibundus* and *lessonae* with high probabilities. This subset includes the three *esculentus* individuals with *ridibundus* mtDNA haplotypes.

The PCAGEN analysis performed on individual genotypes identified the same three clusters and six outliers comprising five *P. esculentus* and one *P. ridibundus* (Fig. 4). The first factor (accounting for 31.22 % of the variance) differentiates *P. lessonae* from *P. ridibundus*, with *P. esculentus* in between. The second factor (9.26 % of the variance) differentiates *P. ridibundus* from *P. esculentus*. The *P. ridibundus* individual with a cytonuclear mismatch (3_EN_13), which was partly assigned to *esculentus* by Structure, takes an intermediate position between these two groups on the 2nd axis, while the five *P. esculentus* individuals assigned to a mix of *lessonae* and *ridibundus* (some of which with *P. ridibundus* mtDNA) also take intermediate positions between these clusters on both axes.

Clustering of genomes

As already pointed out, the assignment of *esculentus* genomes to either *lessonae* (L) or *ridibundus* (R') origin was straightforward: four markers had a *ridibundus*-specific expression, four a *lessonae*-specific expression, and the ten others had parental-specific allelic sizes and/or frequency distributions. The PCAGEN analysis performed after discrimination of the *lessonae* (L) and *ridibundus* (R')

components of P. esculentus genomes also provides clearcut results (Fig. 5). The first factor (accounting for 37.78 % of the variance) mostly differentiates the lessonae (L) from the *ridibundus* (R and R') genomes, while the second axis (5.27 % of variance) discriminates between the R and the R' genomes. All esculentus L genomes perfectly co-localize with the lessonae (LL) genomes (with however a larger variance in the distribution due to their haploid state), while their R' genomes cluster in two groups. On one hand, 82 R' genomes cluster into 11 haplotypes that are markedly diverged from the RR genomes and poorly diversified: 59 individuals share the exact same haplotype H1 (i.e., the same alleles at all of the 14 loci), while the 23 others have 1 of 10 other haplotypes that each differ from H1 by only 1 or at most 3 allelic substitutions. On the other hand, the 5 remaining R' genomes cluster into 4 haplotypes partly overlapping the RR cluster, and diverging from H1 in at least 7 of the 14 loci. These diverged haplotypes characterize the five esculentus individuals that, according to STRUCTURE, share genetic features with the ridibundus and lessonae genomes (Fig. 3). The RR cluster, finally, is quite homogeneous, with the exception of the ridibundus individual with a cytonuclear mismatch (3 EN 13) that takes an intermediate position between the RR and R' genomes.

Spatial distribution of genomes

The three taxa were not homogeneously distributed across our five sampling sites (Fig. S1): while the hybridogenetic *P. esculentus* occurred widespread at all sites, *P. ridibundus* was mostly limited to the westernmost site (Chevroux). *P. lessonae* was nearly absent from this *ridibundus*-dominated site, but constituted a significant part of the four other sites, and the majority in one of them (Gletterens 2). The *P.*



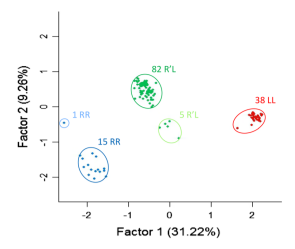


Fig. 4 Biplot of factor scores from the PCA performed on genotypes; RR *P. ridibundus* in *blue* (with the neo-*ridibundus* individual 3_EN_13 in *pale blue*), ancient R'L *P. esculentus* in *dark green*, neo R'L *esculentus* in *pale green*, and *P. lessonae* in *red*

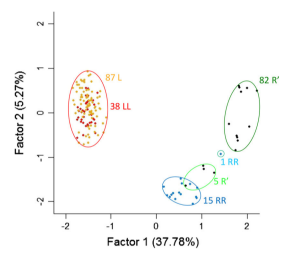


Fig. 5 Biplot of factor scores from the PCA performed on genomes; LL genomes of *lessonae* in *red*, L genomes of *esculentus* in *yellow*, RR genomes of *ridibundus* in *blue* (with the neo-*ridibundus* individual 3_EN_13 in *pale blue*), R' genomes of *esculentus* in *black*

ridibundus with a cytonuclear mismatch (3_EN_13) was found at the site dominated by ridibundus and esculentus genomes (Chevroux), while the five esculentus with diverged hemiclones were spread among four sampling sites (Fig. S1).

Discussion

Our 18 microsatellite markers allowed straightforward species assignment: the patterns of amplification and allelic size ranges were highly specific to the three *Pelophylax* taxa, with no evidence of introgression between the

lessonae and ridibundus genomes. Nuclear identification was corroborated by mtDNA analysis: the cytb tree consisted of two reciprocally monophyletic groups, with all lessonae clustering on one side, and all ridibundus except one on the other side (Fig. 2). Nuclear genetic diversity was high in P. lessonae, and even higher in the invading P. ridibundus (Fig. 1), suggesting multiple introductions (as also supported by the high cytb diversity).

The L genome from P. esculentus individuals was undistinguishable from that of pure P. lessonae (Fig. 5), as expected from ongoing hybridogenesis: the L genome is eliminated at each generation, and has to be obtained anew from a P. lessonae partner. In contrast, the ridibundus (R')genome from most P. esculentus individuals (82 out of 87) differed drastically from the R genomes of invading P. ridibundus (Fig. 5), pointing to anciently diverged hemiclones. The clonal transmission of these frozen R' genomes was confirmed by their very low genetic diversity (Fig. 1) and low number of distinct haplotypes. All these 82 ancient esculentus lineages harbored lessonae mtDNA, as expected from a long history of coexistence. For size and behaviorrelated reasons, L×E crosses often take place between esculentus females and lessonae males (Berger 1970), but reverse crosses also occur at some lower rate (Lengagne et al. 2006). Assuming lessonae females to contribute as few as 10 % of the crosses, the ridibundus mtDNA is expected to decline at a 10 % rate per generation, reaching frequencies below 5 % within 30 generations. Thus, occasional crosses between P. esculentus males and P. lessonae females are expected to progressively eliminate the original ridibundus mtDNA from L-E systems (Spolsky and Uzzell 1986; Plötner et al. 2008).

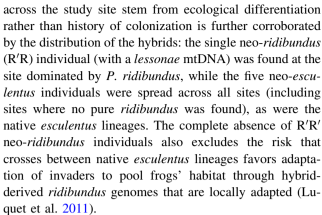
However, five P. esculentus displayed a very distinct pattern. Their R' genomes cluster with those of P. ridibundus (RR), not those of other P. esculentus (Figs. 4, 5), pointing to recent hybridization events between invasive marsh frogs and native pool frogs. Three of them (3_EN_02, 2_EN_19, 1_EN_40) harbored ridibundus mtDNA, revealing their origin as a cross between a male lessonae and a female ridibundus. Interestingly, this includes two females (3 EN 02, 2 EN 19) coming from different localities (respectively Chevroux and Gletterens 1) that share the exact same R' hemiclone and exact same ridibundus cytb haplotype. They therefore represent the 2nd generation (or more) of a new hybridogenetic lineage, created anew from a cross between a ridibundus mother and a lessonae father, with clonal transmission of the nuclear and mitochondrial DNA through R'L mother(s). In addition, one *ridibundus* female (3 EN 13) also displayed a distinct pattern, with a RR' genome and a lessonae cyth, thus clearly resulting from a cross between a P. esculentus (R'L) mother and a P. ridibundus (RR) father. We did not find any R'R' ridibundus, suggesting that, even if some



crosses between ancient *esculentus* lineages occasionally produce viable tadpoles (Hotz et al. 1992; Vorburger 2001a; Luquet et al. 2011), these offspring perform poorly at post-metamorphic stages.

As the six outliers clearly show, several components of the genetic mechanism of species replacement identified by Vorburger and Reyer (2003) are underway in the study site; invasive P. ridibundus are giving rise to neo-ridibundus lineages (RR') by crossing with native P. esculentus, and to new hybridogenetic esculentus lineages by crossing with native P. lessonae. As these new esculentus lineages harbor ridibundus hemiclones devoid of deleterious mutations, further crosses between esculentus should produce viable ridibundus offspring. In theory, therefore, all ingredients of the genetic mechanism of species replacement envisioned by Vorburger and Reyer (2003) are in place. This raises the question of whether this mechanism will end up in the complete elimination of the native L-E system by invasive marsh frogs, as otherwise documented in several places throughout Western Europe (see "Introduction" section). Fully answering this question requires long-term monitoring of the study site; similar genetic sampling in approximately another decade would certainly provide useful insights on the dynamics of the process. However, the data already in hand suggest some reasons for optimism in this specific case. Although marsh frogs have been recorded for several decades at the site, the L–E system still represents 85 % of all water frogs, with 27 % of pure P. lessonae and 58 % of ancient esculentus lineages. The lessonae genome harbors high gene diversity, testifying to large effective populations sizes. We found no evidence for genetic introgression between the R and L genomes, expected if introduced P. ridibundus do not induce hybridogenesis (Hotz et al. 1985; Vorburger 2001b). This may suggest either that new esculentus are all hybridogenetic or that non-hybridogenetic hybrids are not fertile. Both mechanisms exclude a risk of genetic swamping (Rhymer and Simberloff 1996).

More importantly, the *ridibundus* and L–E complex show strikingly distinct distributions over the study sites (Fig. S1), with one single site heavily colonized by P. *ridibundus*, and four other sites dominated by the L–E system. This heterogeneity likely stems from ecological differences: P. *ridibundus* prefers open and well-oxygenated water bodies such as lakes and large ponds, whereas P. *lessonae* seems more adapted to small water bodies with low oxygen content and dense vegetation cover (Negovetic et al. 2001; Plénet et al. 2001, 2005; Holenweg Peter et al. 2002; Schmeller et al. 2007; Meyer et al. 2009). Accordingly, densely vegetated sites in the eastern part of our study area only harbored P. *lessonae* and P. *esculentus*, P. *ridibundus* being limited to the open site with low vegetation cover of Chevroux. That their distinct distributions



Thus, although our study provides direct evidence for genetic interactions between the invasive P. ridibundus and the native P. lessonae and P. esculentus that might, in theory, lead to the replacement of this native species complex, our data also suggest that ecological differentiation is sufficient to limit the extent of interbreeding, so that the three taxa can be expected to stably coexist in the study area over the coming decades. This rather optimistic statement obviously assumes long-term maintenance of important landscape features, including old ponds with a dense vegetation cover. This, however, may pose a dilemma in terms of wetland management. Due to intrinsic dynamics, any water body is bound to progressively disappear via the processes of alluviation, silting, and natural succession. A usual way to counter these dynamics is through pond restoration by re-digging and re-profiling. Pond restoration is certainly a highly successful option for pioneer species (e.g. Petranka et al. 2007; Rannap et al. 2009), not only among amphibians. However, this may also pose a threat for specialist species like P. lessonae, which require shallow stagnant water bodies with dense herbaceous vegetation. If space and resources allow, wetland management policies should in such cases focus on the creation of new ponds. Alternatively, old ponds should be restored in such a way that they match the ecological requirements of the native species (Plénet et al. 2001, 2005; Holenweg Peter et al. 2002; Schmeller et al. 2007).

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