



ELSEVIER

Available online at www.sciencedirect.com

SCIENCE @ DIRECT®

C. R. Biologies 326 (2003) S85–S92



Frog alien species: a way for genetic invasion?

Alain Pagano^{a,b,c,*}, Alain Dubois^d, David Lesbarrères^a, Thierry Lodé^a

^a Laboratoire d'écologie animale, Université d'Angers, Campus Belle Beille, 49045 Angers, France

^b UMR CNRS 5023, Ecologie des hydrosystèmes fluviaux, Université Lyon 1, 43 bvd du 11 novembre 1918, 69622 Villeurbanne cedex, France

^c CEPM, UMR CNRS IRD 9926, Centre de recherches IRD, 911 av. Agropolis, BP 5045, 34032 Montpellier cedex 1, France

^d Laboratoire des Reptiles & Amphibiens, Muséum National d'Histoire Naturelle, 25 rue Cuvier, 75005 Paris, France

Abstract

European water frogs are characterized by anthropic introductions and *Rana ridibunda* may be considered as an invasive species. As such translocations may result in introgression of exotic genes in native populations, i.e. genetic pollution, we studied genetic characteristics (on 11 allozymic loci) of natural versus introduced water frogs. Our study contributed to (1) disclose 3 genetic markers allowing the identification of exotic frogs; (2) quantify the proportion of exotic frogs found in natural populations; and (3) suggest how genetic pollution may arise in these frogs. **To cite this article:** A. Pagano et al., C. R. Biologies 326 (2003).

© 2003 Académie des sciences. Published by Éditions scientifiques et médicales Elsevier SAS. All rights reserved.

Résumé

Les grenouilles introduites : une voie pour l'invasion génétique ? Les grenouilles vertes européennes sont caractérisées par des introductions anthropiques et *Rana ridibunda* peut être considérée comme une espèce invasive. Comme les introductions peuvent causer des introgressions de gènes exotiques dans les populations autochtones, c'est-à-dire de la pollution génétique, les caractéristiques génétiques de grenouilles « naturelles » et introduites ont été étudiées pour 11 locus allozymiques. Notre étude a contribué à (1) identifier 3 marqueurs génétiques permettant l'identification des grenouilles exotiques, (2) quantifier la proportion de grenouilles exotiques présentes dans les populations naturelles et (3) suggérer comment la pollution génétique peut survenir chez ces grenouilles. **Pour citer cet article :** A. Pagano et al., C. R. Biologies 326 (2003).

© 2003 Académie des sciences. Published by Éditions scientifiques et médicales Elsevier SAS. All rights reserved.

Keywords: introgression; introduction; hybridogenesis; conservation; water frogs

Mots-clés : introgression ; introduction ; hybridogénèse ; conservation ; grenouilles vertes

1. Introduction

Introduction of “exotic” (i.e., non native from the area considered) species in ecosystems may have sev-

eral results: (1) ecological consequences such as competitive exclusion, introduction of parasites or diseases, predation, modification of trophic structure, etc. (e.g. [1–3]); or, (2) genetic consequences. Among them, hybridization between exotic and autochthonous species has been documented (e.g. [4–6]). When hybrids are sterile, hybridization represents a cost as parental species “fail” in producing a successful off-

* Corresponding author.

E-mail address: alain.pagano@univ-angers.fr (A. Pagano).

spring [7]. Such a cost is amplified in species close to extinction because the number of effective breeders is low and critical with regard to population persistence. Moreover, in hybrid zones, processes of introgression often occur [8]. In a context of hybridization induced by man's activity, introgression may be viewed as a genetic invasion of exotic genes from one species to another, a process that has been called "genetic pollution" [9–11]. Such a process is a threat for the genetic integrity of native genomes [12] and thus represents a model for conservation genetics studies. We want to underline that genetic pollution does not only affect distinct species that are involved in introgression but also affects, within a species, distinct allopatric populations that are genetically different or locally adapted. Because of anthropic translocation, allopatry is broken and these populations may be involved in gene exchanges.

Invasive species, mainly resulting from introductions by man, are often implied in perturbations of ecosystems [2]. As an example, with their range distribution increasing, hybridization and introgression with localized species become possible. Thus, genetic pollution may be important for invasive species. European water frogs provide a good model to study such processes. Such frogs are subject to multiple and recurrent introductions [13–16]. It is especially the case in France because of accidental introductions of the species *R. ridibunda* that are imported mainly from Egypt, Turkey and Balkanic countries for culinary purposes (ca. 700 t of frogs; [14,17]). As a consequence, *Rana ridibunda* that was previously located to a narrow part of France (Fig. 1) may be considered as an invasive species because its range is now extending to many other parts of France [16]. Several other European water frog species that occur naturally in the Balkanic region (*Rana kurtmuelleri*), the Near East (*Rana bedriagae*) or the northern part of Africa (*Rana saharica*) may also have been introduced in France, although they have not yet been identified [18]. Thus these frogs may be affected by genetic pollution because of translocations.

Moreover, these frogs exhibit a peculiar situation regarding animal hybridisation. They constitute "kleptons" [19,20], i.e. unusual biological "species" which have a peculiar mode of reproduction such as gynogenesis or "hybridogenesis" [21–23]. Especially interesting in this respect are "zygokleptons" [19], i.e.

forms of hybrid origin that are fertile, discard in germinal cells one parental genome and produce clonal (non recombining) gametes allowing the persistence of hybrid lineages through "sexual parasitism" of the parental species to re-establish at each generation the genome lost through meiosis (so-called "hybridogenesis" or creditogenesis; see [24]). Usually, such forms are expected to be prevented from genetic pollution because recombination is not expected to occur. However, occasional recombination has been suspected in natural populations of hybridogenetic European water frogs [25,26]. Thus, the integrity of genomes needs to be specified in the taxa implied in such hybridogenetic complexes.

In this context, our study aims at (1) identifying genetic markers that allow one to evidence introductions in *R. ridibunda* populations; (2) quantifying the proportion of introduced specimens in these natural populations; (3) surveying the "integrity" of their genomes; (4) identifying places where allopatry is broken and co-occurrence (assemblages) arise between different water frog species studied in France; and (5) check if introgression may occur. Moreover, we would like to propose a theoretical scenario for the "genetic pollution hypothesis" and discuss its biological consequences.

2. Materials and methods

2.1. *Rana ridibunda* survey

European water frogs ($n = 254$) were collected in 11 distinct populations from eastern France in 1997 (along the French Rhone river from North to South; see [16]). As introductions mainly originated from frog importation for culinary purposes, two additional samples were studied, coming from a fish store that imports such frog for restaurants. These frogs are known to originate from Turkey and Egypt, the main exporting countries for these living frogs [17].

Genetic variability was studied on 11 presumptive structural gene loci by starch gel electrophoresis, using continuous buffer systems (Tris-citrate pH 6, Tris-citrate pH 8 and Tris-EDTA-borate pH 8). Slices were stained following standard protocols [27,28]. Loci successfully resolved were: ACO-2 (4.2.1.3), GDA, α -GDH-1 and α -GDH-2, LDH-1 and LDH-2

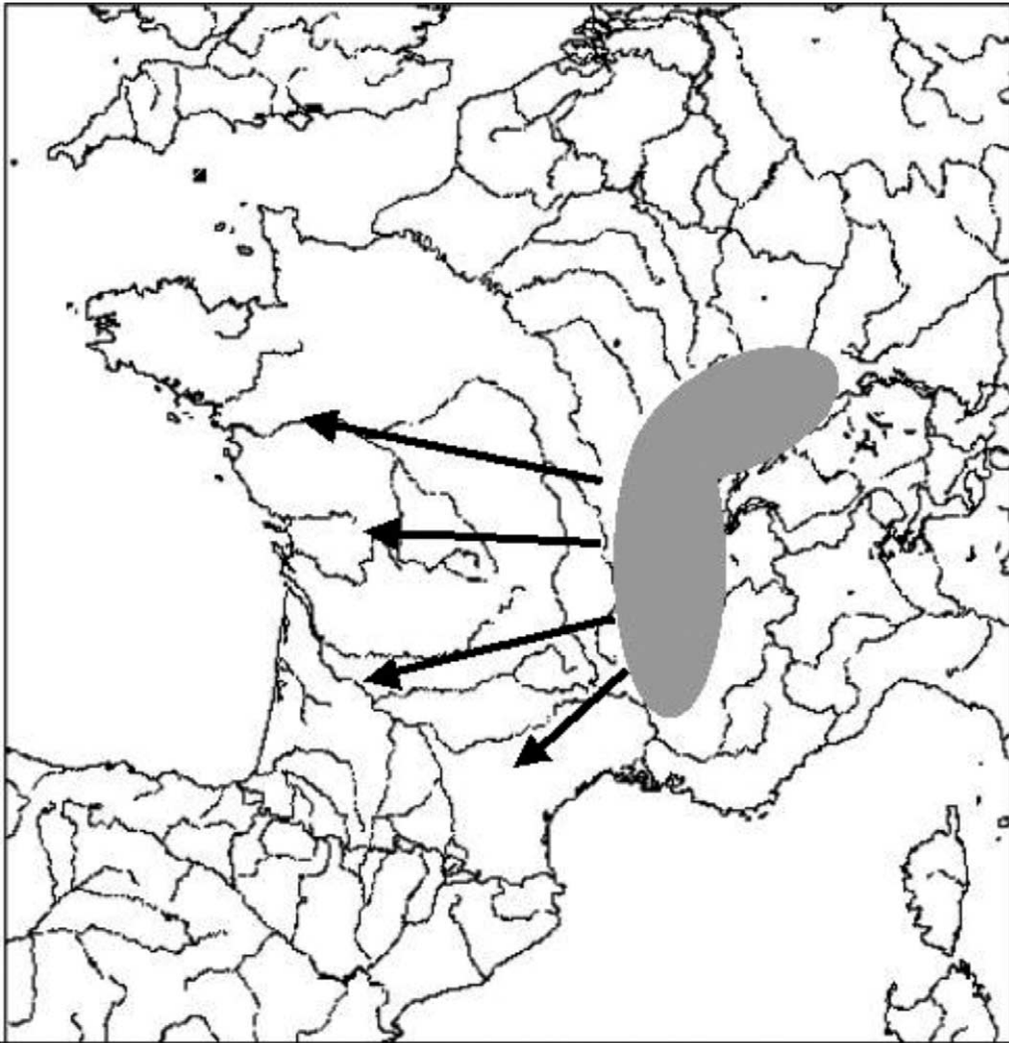


Fig. 1. Range distribution area of *R. ridibunda* in France. The grey area corresponds to the distribution mapped by Graf & Polls Pelaz [23]. The arrows represent range expansion such as that recorded recently (see [16]).

(1.1.1.27), MPI (5.3.1.8), 6PGDH (1.1.1.44), PGM-1 and PGM-2 (2.7.5.1), SOD (1.15.1.1).

A multivariate analysis (FCA) was performed on the 11 loci in order to see if autochthonous and exotic frogs could be distinguished.

2.2. General water frog survey

Altogether 608 frogs were randomly sampled in 32 aquatic sites. In order to determine the taxonomic composition at each site, each frog was identified

using a combination of specific allozymic markers, specifically LDH-B, α GDH (E.C. 1.1.1.8), AHH (E.C. 3.3.1.1), MPI, and PGM-2.

3. Results

3.1. *R. ridibunda* survey

Among the 11 loci, all were polymorphs except the SOD (data not shown). As revealed by allelic frequencies (Table 1), three loci appeared as genetic

Table 1
Allelic frequencies at 3 loci. Underlined numbers represent the (suspected) markers of allochthonous origin

Locus	Allele	France	Turkey	Egypt
		<i>N</i> = 254	<i>N</i> = 9	<i>N</i> = 10
GDA	<u>100</u>	<u>0.0147</u>	–	<u>0.2000</u>
	110	0.2990	0.6111	0.8000
	120	0.6765	0.3889	–
	<u>130</u>	<u>0.0098</u>	–	–
ACO-2	100	0.3088	0.1111	0.0556
	105	0.0069	–	–
	110	0.6567	0.5556	0.3333
	<u>120</u>	<u>0.0276</u>	<u>0.3333</u>	<u>0.6111</u>
MPI	90	0.0042	–	–
	100	0.4937	0.2000	–
	119	0.4958	0.8000	0.80
	<u>135</u>	<u>0.0063</u>	–	<u>0.20</u>

markers of introduction from Turkey and Egypt: GDA, ACO-2 and MPI. The allele GDA-100 was common in the Egyptian frogs analysed (20%) but very rare in autochthonous frogs (1.4%; 4 frogs among which 2 were homozygote 100/100). The allele 120 of ACO-2 reached 33.33% in the Turkish sample and 61.11% in the Egyptian one. Although this allele was scarce in French populations (2.76%), all 6 frogs bearing this allele were homozygote 120-120. The MPI-135 allele was rare in France (0.63%) and more common in the Egyptian sample (12.5%).

The multivariate analysis performed on the whole set of loci showed that (1) there is a genetic difference between the autochthonous and exotic frogs (genetic distance $d = 0.33$ for Egyptian frogs, $d = 0.115$ for Turkish ones; [29]); (2) this difference is mainly due to ACO, GDA-100 and GDA-130 (axis 1, 3, and 2 of the analysis, respectively); and (3) the “French” frogs that bear alleles GDA-100 and ACO(2)-120 are very close to exotic populations and different from the other French populations, suggesting that such frogs are, in fact, exotic ones and originated from introductions (Fig. 2). The frogs bearing the GDA-130 allele were also close to exotic frogs because they also show the ACO(2)-120 allele.

Based on the genotype and the factorial map, we can estimate the introduced individuals to at least 10 frogs (3.94%) from 6 distinct populations. Thus, 54.5% of the populations were affected by such introductions.

If considering that “exotic-like” frogs are to be discarded from the French populations to evaluate allelic frequencies, thus GDA-130 and ACO-120 become absent in France i.e. represent specific markers between French populations and allochthonous ones (Table 2). GDA-100 and MPI-135 remain present although scarce because few frogs bear these alleles at an heterozygous state. It suggests cases of genetic pollution i.e. invasion of exotic alleles in the genetic pool of Western populations.

3.2. General water frog survey

The 608 individuals belong to *Rana ridibunda* ($n = 355$ frogs; 58.38%); *R. kl. esculenta* ($n = 178$; 29.28%); and *R. lessonae* ($n = 75$; 12.33%).

Among the 32 sites, the taxonomic composition recorded was:

1. Pure *R. ridibunda* population in 14 ponds (43.75%).
2. L-E assemblage (*R. lessonae* + *R. kl. esculenta*) in six ponds (18.75%).
3. Pure *R. kl. esculenta* population in five ponds (15.63%).
4. Assemblage involving *R. ridibunda* + another unexpected water frog species in three ponds (9.38%).
5. R-E assemblage (*R. ridibunda* + *R. kl. esculenta*) in two ponds (6.25%).
6. Pure *R. lessonae* population in one pond (2.27%).

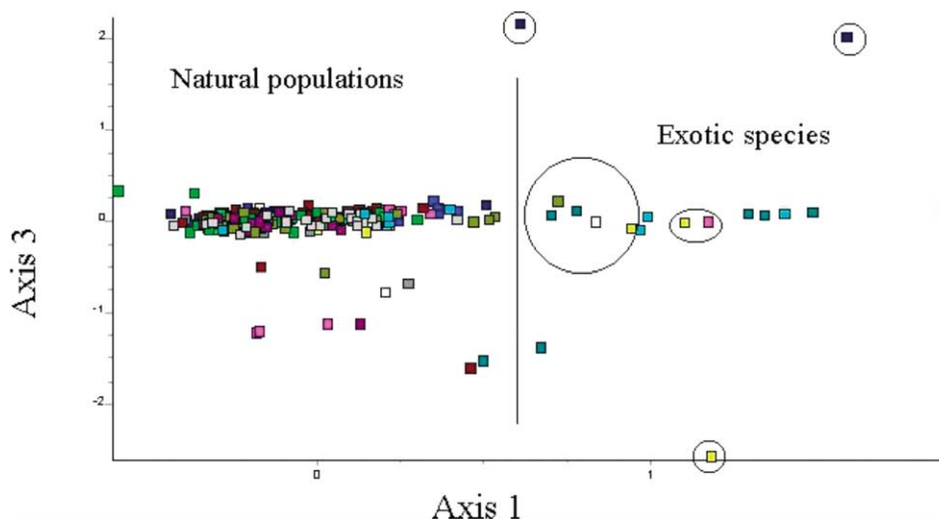


Fig. 2. Factorial map performed on the multilocus genotypes of 11 populations of *R. ridibunda* from France and 2 samples from Egypt and Turkey. Individuals on the right part of the map are from samples obtained from a fish store (blue-sky and blue-green are respectively from Turkey and Egypt). The other individuals surrounded by a circle were sampled in French natural populations and are suspected to represent exotic frogs i.e. introductions.

Table 2

Allelic frequencies at 3 loci. The third column represents allelic frequencies calculated on the whole sample of frog from French natural populations while the fourth column represents allelic frequencies after discarding from the sample the individuals suspected to be exotic. As underlined numbers represent the (suspected) markers of allochthonous origin., the fifth column may represent the cases of genetic pollution. The allele with * (GDA-130) represents a potential additional marker because it is linked with ACO(2)-120 in our samples

Locus	Allele	All frogs from France	Sample without "exotic-like" frogs	Remaining frogs with exotic allele
		<i>N</i> = 254	<i>N</i> = 244	
GDA	<u>100</u>	<u>0.0147</u>	<u>0.0102</u>	<u>2 genotypes GDA 100–110</u>
	110	0.2990	0.2944	
	120	0.6765	0.6954	
	<u>130*</u>	<u>0.0098</u>	<u>0*</u>	
ACO-2	100	0.3088	0.3190	
	105	0.0069	0.0071	
	110	0.6567	0.6738	
	<u>120</u>	<u>0.0276</u>	<u>0</u>	
MPI	90	0.0042	0.0043	
	100	0.4937	0.4935	
	119	0.4958	0.4957	
	<u>135</u>	<u>0.0063</u>	<u>0.0065</u>	<u>3 genotypes MPI 119–135 or 100–135</u>

Table 3

Genotypic frequencies recorded at three loci. In bold, genotypes characteristics of *R. kl. esculenta*; underlined, those characteristics of *R. lessonae* (see e.g. [27]). Asterisks represent suspected introgression cases

Taxon	<i>N</i>	LDH-B				MPI			PGM-2		
		Ae	ce	<u>be</u>	<u>ee</u>	ah	ch	<u>hh</u>	dd	cd	<u>cc</u>
<i>R. kl. esculenta</i>	23	12	11	<u>5</u>	<u>10</u>	13	10	<u>16</u>	6*	16	<u>1*</u>
<i>R. lessonae</i>	<u>16</u>	1*		<u>5</u>	<u>10</u>			<u>16</u>		1*	<u>15</u>

7. Assemblage involving *R. lessonae* + another unexpected water frog species in one pond (2.27%).

Several genotypes recorded within two taxa at one pond suggest occasional recombination (Table 3) and thus gene exchanges from one species to another.

4. Discussion

Our study highlights the difference in genetic structure (allelic composition, genetic distance) between Oriental populations (Egypt and Turkey) and Occidental ones. However, in France, several frogs caught in natural populations are, in fact, genetically close to frogs originating from Egypt and Turkey, the two main sources for frog import (Fig. 2). In addition, such individuals bear alleles, in an homozygote condition, that are rare in France but very common or even dominant in Oriental populations. This strongly suggests that they are in fact exotic frogs resulting from recent introductions.

This hypothesis is confirmed by the fact that the elimination of the ten “exotic-like” individuals from the sample of French frogs modifies the allelic frequencies so that the alleles GDA-130 and ACO(2)-120 appear then absent from French natural populations (i.e. specific markers of allochthonous origin). The ACO(2)-120 allele appears without doubt as a marker of Oriental origin.

The GDA-130 allele was not found in the exotic individuals studied probably because of our limited sampling size but it seems also to be a marker of Oriental populations as it is linked with ACO(2)-120 in our sample. The MPI-135 allele has already been suspected to reveal an exotic origin as it is present in populations from Anatolia [17]. These results point to the need for an extensive genetic study of Oriental frogs in order: (1) to identify with more accuracy the alleles that can be considered reliable markers of exotic origin; (2) to be able to establish more precisely which frogs are introduced and how many populations are affected by introductions; and (3) to begin to measure the biological and genetic (e.g. introgression) effects of such introductions.

Our study strongly suggests that introductions have been multiple as they affect at least 6 populations several hundred kilometres distant (near Camargue,

Southern France and near Leman lake, Central-Eastern France). Indeed, 54.5% of the studied populations, seem to contain, at least, one exotic individual while a sample from a population from Southern France disclosed at least three introduced frogs (16.6% of the sample). These observations highlight the potential importance of introductions on the biology of natural populations.

Our study validates the hypothesis of multiple introductions that was invoked to account for the expansion of the distribution of *R. ridibunda* in most parts of France ([16] and Fig. 1). Thus, *R. ridibunda* may be considered as an invasive species in France. However, the presence of ancient native populations cannot be fully discarded nor the hypothesis of more ancient introductions from other countries such as those recorded in Switzerland [30]. All biological consequences of these introductions are to be studied. Because of their importance in France, one major risk is the extinction of remaining natural populations of *R. ridibunda* if they still exist. Another risk is to modify species distribution, favouring unexpected sympatries and give rise to new hybridisation opportunities [31] that is the basis for processes of genetic pollution. As an example, we recorded 3 sites with unexpected assemblages likely because of translocations of *R. ridibunda* in a region where it was expected previously to be absent.

The scarce presence of exotic (Egyptian) alleles GDA-100 and MPI-135 in the French population even after discarding “exotic-like” individuals suggests an introgression of exotic genes in French populations (i.e. genetic pollution). This would mean a first way for genetic pollution, at the intra-specific level, through matings between Oriental and Western populations of *R. ridibunda* (Fig. 3a). This may have important biological consequences, as it is known that Oriental populations of this species exhibit strong differences in genetic structure and biology: in particular, they are hybridogenetic-resistant [32], and no hybrids are known to be present in these Oriental populations.

A second way for genetic pollution may occur at the inter-specific level through (natural) hybridization between a “polluted” *R. ridibunda* and a *R. lessonae* (Fig. 3b). Such matings would result in “polluted hybrids”. As, in European water frogs, hybrids are characterized by fertility, hybridogenesis (i.e. hemiclonal reproduction through genome exclusion of the

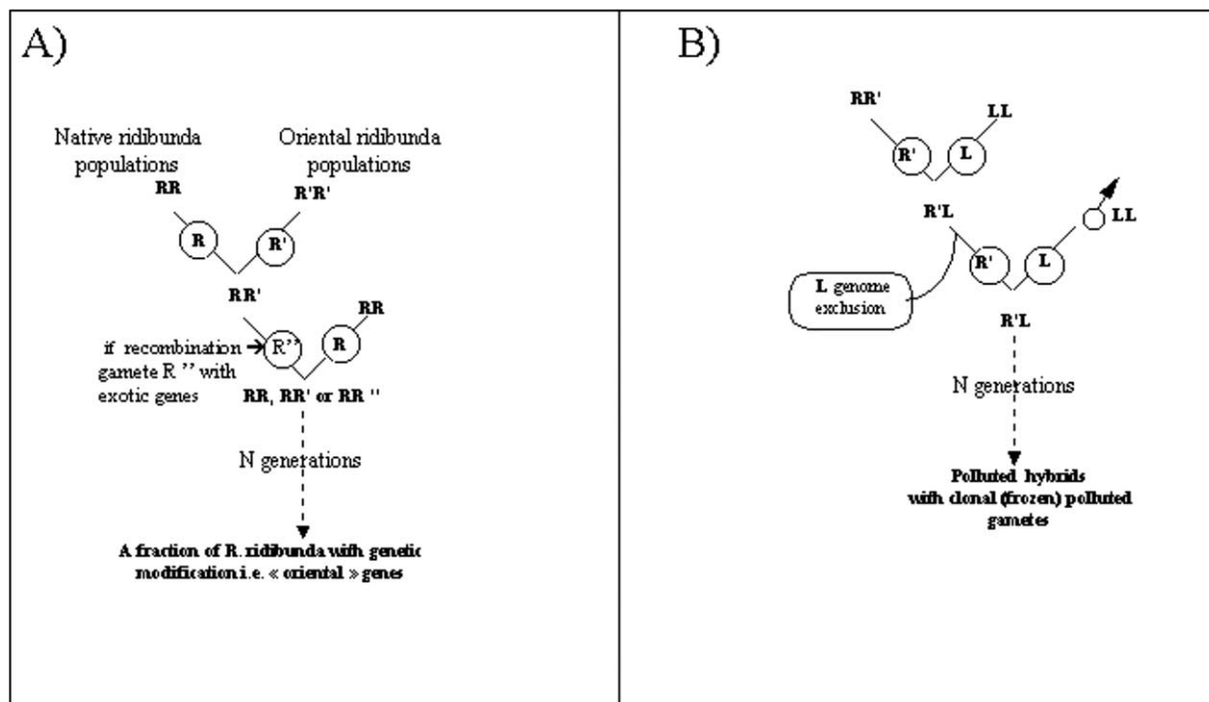


Fig. 3. Theoretical scenario of genetic pollution. (A) Within the *R. ridibunda* species. (B) Transmission of exotic alleles to another taxon (a hybrid taxon; *R. kl. esculenta*) through natural hybridization between a polluted *R. ridibunda* and *R. lessonae*.

lessonae genome; see e.g. [23]) and long-term persistence through matings at each generation with *R. lessonae*, such hybrid may produce “polluted gametes” bearing exotic genes. In such cases, the polluted genome is persistent in hybrids through generations. A genetic investigation of hybrids using markers of allochthonous origin remains to be performed in order to establish the existence of polluted hybrids. Such a way does not appear purely hypothetical, as our results suggested occasional recombination and introgression within water frogs.

5. Conclusions

Considering water frogs, studies are still needed to test the validity of the genetic pollution models proposed and to establish their biological consequences, because our work is preliminary. However, we aim to underline that the genetic pollution model has a more general value as introgression is often caused by introduction of exotic populations or exotic species. Studies on genetic pollution have been too much neglected

in conservation biology [10,11] because it is not clear if it may represent a direct risk for species survival. Nevertheless, it clearly affects the genetic integrity of natural populations and cannot be without consequences on their fate. Thus, it represents a problem to be taken into account in future conservation genetics studies.

Acknowledgements

We want to thank F. Renaud for assistance in performing protein electrophoresis.

References

- [1] J.A. Drake, H.A. Mooney, F. Di Castri, R. Groves, F. Kruger, M. Rejmanek, M. Williamson, *Biological Invasion. A Global Perspective*, Wiley J. & Sons, 1989.
- [2] D.M. Lodge, *Biological invasions: lessons from ecology*, TREE 8 (1993) 133–137.

- [3] J.M. Kiesecker, A.R. Blaustein, C.L. Miller, Transfer of pathogen from fish to Amphibians, *Conserv. Biol.* 15 (2001) 1064–1070.
- [4] D. Gottelli, C. Sillero-Zubiri, G.D. Appelbaum, M.S. Roy, D.J. Girman, J. Garcia-Moreno, E.A. Ostrander, R.K. Wayne, Molecular genetics of the most endangered canid: the Ethiopian wolf *Canis simensis*, *Mol. Ecol.* 3 (1994) 301–312.
- [5] E. Randi, M. Pierpaoli, M. Beaumont, B. Ragni, A. Sforzi, Genetic identification of wild and domestic cats (*Felis silvestris*) and their hybrids using Bayesian clustering methods, *Mol. Biol. Evol.* 18 (2001) 1679–1693.
- [6] Z. Andersone, V. Lucchini, E. Randi, J. Ozolins, Hybridisation between wolves and dogs in Latvia as documented using mitochondrial and microsatellite DNA markers, *Mammal. Biol.* 67 (2002) 79–90.
- [7] R.G. Harrison, Hybrid zones: windows on evolutionary process, *Oxford Surveys in Evolutionary Biology* 7 (1990) 69–128.
- [8] T.E. Dowling, C.L. Secor, The role of hybridisation and introgression in the diversification of animals, *Ann. Rev. Ecol. Syst.* 28 (1997) 593–619.
- [9] A. Dubois, J.-J. Morère, A propos des introductions d'espèces réalisées par Raymond Rollinat, *Bull. Soc. Herp. Fr.* 9 (1979) 59–61.
- [10] A. Dubois, J.-J. Morère, Pollution génétique et pollution culturelle, *C. R. Soc. Biogéogr.* 56 (1980) 5–22; *Bull. Assoc. Franç. Ingén. Ecologues* 3 (1982) 10–14, reprint.
- [11] A. Dubois, Renforcements de populations et pollution génétique, *C. R. Soc. Biogéogr.* 59 (1983) 285–294.
- [12] J.M. Rhymer, D. Simberloff, Extinction by hybridization and introgression, *Ann. Rev. Ecol. Syst.* 27 (1996) 83–109.
- [13] A. Dubois, Notes sur les grenouilles vertes (groupe de *Rana kl. esculenta* Linné, 1758). I. Introduction, *Alytes* 1 (1982) 42–49.
- [14] A. Dubois, A propos de cuisses de grenouilles. Protection des amphibiens, arrêtés ministériels, projets d'élevage, gestion des populations naturelles, enquêtes de répartition, production, importations et consommation: une équation difficile à résoudre. Les propositions de la Société Batrachologique de France, *Alytes* 2 (1983) 69–111.
- [15] B. Arano, G. Llorente, M. García-Paris, P. Herrero, Species translocation menaces Iberian waterfrogs, *Conserv. Biol.* 9 (1995) 196–198.
- [16] A. Pagano, P.-A. Crochet, J.-D. Graf, P. Joly, T. Lodé, Distribution and habitat use of Water frog hybrid complexes in France, *Global Ecol. & Biogeogr.* 10 (2001) 433–442.
- [17] A. Pagano, P. Joly, H. Hotz, Taxonomic composition and genetic variation of water frogs in the mid-Rhône floodplain, *C. R. Acad. Sci. Paris, Sciences de la vie/Life Sciences* 320 (1997) 759–766.
- [18] A. Dubois, Les Amphibiens et les introductions d'espèces allogènes dans les milieux, in: *Gestion et protection des Amphibiens : de la connaissance à la prise en compte dans les aménagements*, Assoc. Franç. Ingén. Ecologues, Paris, 2002.
- [19] A. Dubois, R. Günther, Klepton and synklepton: two new evolutionary systematics categories in zoology, *Zool. Jb. Syst.* 109 (1982) 290–305.
- [20] A. Dubois, Nomenclature of parthenogenetic, gynogenetic and “hybridogenetic” vertebrate taxons: new proposals, *Alytes* 8 (1991) 61–74.
- [21] A. Dubois, Les problèmes de l'espèce chez les amphibiens anoures, in: C. Bocquet, J. Générmon, M. Lamotte (Eds.), *Mém. Soc. zool. Fr.* 39, in: *Les problèmes de l'espèce dans le règne animal*, Vol. 2, 1977, pp. 161–284.
- [22] R.C. Vrijenhoek, R.M. Dawley, C.J. Cole, J.P. Bogart, A list of known unisexual vertebrates, in: R.M. Dawley, J.P. Bogart (Eds.), *Evolution and Ecology of Unisexual Vertebrates*, Bulletin 466, New York State Museum Publications, Albany, 1989, pp. 24–31.
- [23] J.-D. Graf, M. Polls-Pelaz, Evolutionary genetics of the *Rana esculenta* complex, in: R.M. Dawley, J.P. Bogart (Eds.), *Evolution and Ecology of Unisexual Vertebrates*, New York State Museum Publications, Albany, 1989, pp. 289–302.
- [24] L.J. Borkin, I.S. Darevsky, Reticulate (hybridogenous) speciation in vertebrates, *Zh. Obshch. Biol.* 41 (1980) 485–506.
- [25] A. Pagano, D. Schmeller, Is recombination less negligible than previously described in hybridogenetic water frogs?, in: C. Miaud, R. Guyétant (Eds.), *Current Studies in Herpetology*, Proceedings of the 9th Ordinary General Meeting of the Societas Europaea Herpetologica, Chambéry, France, 1999, pp. 351–356.
- [26] C. Vorburger, Non-hybrid offspring from matings between hemiclinal hybrid waterfrogs suggest occasional recombination between clonal genomes, *Ecol. Lett.* 4 (2001) 628–636.
- [27] H. Hotz, Genetic diversity among water frog genomes inherited with and without recombination. Ph.D. Dissertation, Universität Zürich, Zürich, 1983.
- [28] N. Pasteur, G. Pasteur, F. Bonhomme, J. Catalan, J. Britton-Davidian, *Manuel technique de génétique par électrophorèse des protéines*, Lavoisier, Paris, 1987.
- [29] M. Nei, Genetic distances between populations, *Am. Nat.* 106 (1972) 283–292.
- [30] K. Grossebacher, Atlas de distribution des Amphibiens de Suisse, *Documenta Faunistica Helvetia* 8 (1988) 1–208.
- [31] A. Pagano, T. Lodé, P.-A. Crochet, New contact zone and assemblages among water frog of Southern France, *J. Zool. Syst. Evol. Res.* 39 (2001) 63–67.
- [32] H. Hotz, G. Mancino, S. Bucci-Innocenti, M. Ragghianti, L. Berger, T. Uzzell, *Rana ridibunda* varies geographically in inducing clonal gametogenesis in interspecies hybrids, *J. Exp. Zool.* 236 (1985) 199–210.