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# Testing an hypothesis of hybrid zone movement for toads in France

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## Abstract

Hybrid zone movement may result in substantial unidirectional introgression of selectively neutral material from the local to the advancing species, leaving a genetic footprint. This genetic footprint is represented by a trail of asymmetric tails and displaced cline centres in the wake of the moving hybrid zone. A peak of admixture linkage disequilibrium is predicted to exist ahead of the centre of the moving hybrid zone. We test these predictions of the movement hypothesis in a hybrid zone between common (*Bufo bufo*) and spined toads (*B. spinosus*), using 31 nuclear and one mtDNA SNPs along a transect in the northwest of France. Average effective selection in *Bufo* hybrids is low and clines vary in shape and centre. A weak pattern of asymmetric introgression is inferred from cline discordance of seven nuclear markers. The dominant direction of gene flow is from *B. spinosus* to *B. bufo* and is in support of southward movement of the hybrid zone. Conversely, a peak of admixture linkage disequilibrium north of the hybrid zone suggests northward movement. These contrasting results can be explained by reproductive isolation of the *B. spinosus* and *B. bufo* gene pools at the southern (*B. spinosus*) side of the hybrid zone. The joint occurrence of asymmetric introgression and admixture linkage disequilibrium can also be explained by the combination of low dispersal and random genetic drift due to low effective population sizes.

## KEYWORDS

admixture linkage disequilibrium, asymmetric reproductive isolation, *Bufo bufo*, *Bufo spinosus*, cline coupling, hybrid zone movement

## 1 | INTRODUCTION

During allopatric speciation, a taxon's range is split by a physical barrier, and the vicariant populations gradually diverge through processes such as mutation, natural selection, and genetic drift (Lande, 1980; Mayr, 1942; Wu & Ting, 2004). Diverged taxa may later meet in secondary contact and form a hybrid zone, for example when taxa expand their ranges from glacial refugia in the postglacial era (Barton & Hewitt, 1985; Hewitt, 1988, 2011). Upon secondary contact the two populations may have established reproductive isolation,

inhibiting exchange of genetic material (Rice, 1998). Conversely, when the fitness of hybrid populations is equal or increased compared to the fitness of parental populations, gene flow is extensive, and the two populations will merge (Anderson & Stebbins, 1954; Arnold, 2004; Hedrick, 2013; Seehausen, 2004). In between these extremes, a reduced hybrid fitness precludes merging and facilitates limited introgression (Mallet, 2005).

When hybrid fitness is reduced independent of the environment, a narrow tension zone exists between the two species (Barton & Hewitt, 1985; Moore, 1977). In the classical hybrid zone literature

such tension zones are thought to stabilise where population density is low, or at an ecological barrier to dispersal (Barton & Hewitt, 1985; Endler, 1977). Recent studies suggest that hybrid zone movement is more common than previously thought (Arntzen & Wallis, 1991, 1999; Buggs, 2007; Leaché, Grummer, Harris, & Breckheimer, 2017; Roy, O'Connor, & Green, 2012; Ryan et al., 2018; Taylor et al., 2014; Wielstra, Burke, Butlin, & Arntzen, 2017; Wielstra, Burke, Butlin, Avci, et al., 2017). Hybrid zone movement occurs when one parent species has a higher fitness than the other, such as through a competitive advantage, asymmetric hybrid fitness effects, or environmental change (Buggs, 2007). Introgression caused by hybrid zone movement is thought to affect selectively neutral markers distributed randomly across the genome, resulting in genetic traces of the displaced species in populations of the advancing species (Currat, Ruedi, Petit, & Excoffier, 2008; Moran, 1981). In contrast, under adaptive introgression, alleles from one species with a positive effect in the other species may introgress through the hybrid zone, regardless of whether the zone is stable or moving (Hedrick, 2013; Mallet, 2005; Schmickl, Marburger, Bray, & Yant, 2017; Seehausen, 2004). Adaptive introgression affects only the selected marker and markers that are nearby on the chromosome (physical linkage), or markers that interact functionally with the marker under selection (functional linkage; Barton & Hewitt, 1985; Baird, 2015; Sedghifar, Brandvain, & Ralph, 2016). Only a fraction of the genome is involved in adaptive introgression, whereas hybrid zone movement results in genome-wide introgression (Currat et al., 2008; Wielstra, Burke, Butlin, Avci, et al., 2017).

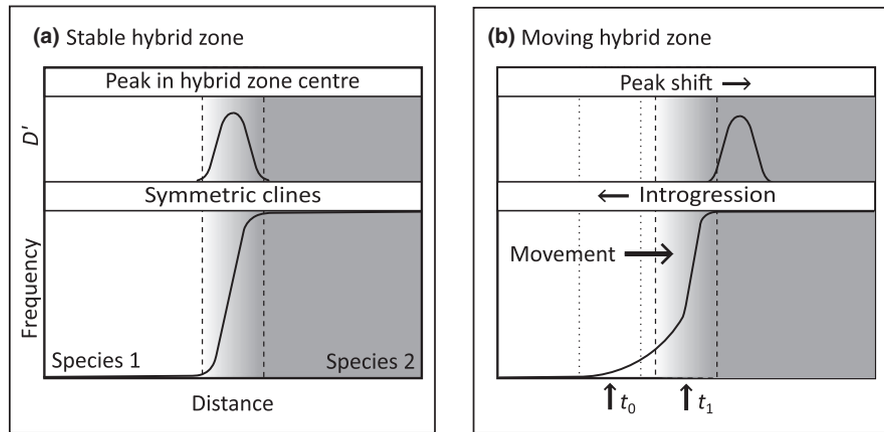
The transition from one species to the other through the hybrid zone can be described by a set of allele frequency gradients, i.e., geographical clines (Schmickl et al., 2017). The positions and shapes of geographical clines are indicative of evolutionary processes involved in hybrid zone formation (Barton & Hewitt, 1985). Two types of cline variation are distinguished based on cline position (coincident or displaced) and cline shape (concordant or discordant). Coincident clines share the same cline centre, whereas displaced clines have a cline centre away from the majority of the clines for other loci. Concordant clines are similar in shape (described by e.g., width and tail shape; Szymura & Barton, 1991), whereas discordant clines have a shape divergent from the majority of the clines. Coincident clines are considered evidence of a hybrid zone in a stable position over time (Abbott et al., 2013). Concordant and narrow clines (steep at the centre and with distinct tails) suggest that the hybrid zone is maintained by a balance between strong selection and dispersal, as opposed to displaced and wide clines indicating that selection against hybrids is low (shallow slope without distinct tails; Barton & Gale, 1993). Displaced and discordant clines exist in many species, both for nuclear and mitochondrial markers, and may be caused by sex-biased gene flow or by adaptive introgression (Bonnet, Leblois, Rousset, & Crochet, 2017; Excoffier, Foll, & Petit, 2009; Mallet, 2005; Sloan, Havird, & Sharbrough, 2016; Toews & Brelsford, 2012; While et al., 2015). However, displaced and discordant clines may also be caused by hybrid zone movement, when a trail of genetic material from the overtaken species remains present in the overtaking species (Arntzen, de Vries, Canestrelli, & Martínez-Solano, 2017; Excoffier et al., 2009;

Gay, Crochet, Bell, & Lenormand, 2008; Rohwer, Bermingham, & Wood, 2001; Wielstra, Burke, Butlin, Avci et al., 2017).

Whether or not markers reflect past or ongoing demographic processes, including hybrid zone movement and adaptive introgression, is dependent on factors such as the number and genomic distribution of barrier loci, mutation rate, and recombination (reviewed in Ravinet et al., 2017). Barrier loci are genes which inhibit the merging of two diverged populations (Abbott et al., 2013; Barton, 2013). A stronger overall barrier to gene flow is established when multiple barrier effects become coincident by processes summarised as coupling (reviewed in Butlin & Smadja, 2017). If genome-wide coupling of barrier effects occurs, the geographic clines for markers are expected to be coincident even if they are not physically or functionally linked to a barrier gene (Barton, 1983; Barton & Gale, 1993; Bierne, Welch, Loire, Bonhomme, & David, 2011; Vines et al., 2016). To overcome the coupling effect of multiple barrier genes and so to cause gene flow across the hybrid zone, the positive selection on a single gene will have to be disproportionately strong (e.g., Vines et al., 2016).

In addition to physical and functional linkage, associations between markers in hybrid zones can be caused by common descent. This is reflected by an increase in admixture linkage disequilibrium in the hybrid zone as alleles originating from the same parent species tend to be found together within the genomes of early generations of hybrid offspring. Recombination during subsequent backcrossing breaks down admixture linkage disequilibrium (Baird, 2015; Barton & Gale, 1993). However, when the hybrid zone is moving, the peak of admixture linkage disequilibrium is predicted to be positioned just ahead of the hybrid zone centre, where individuals with little history of recombination are involved in reproduction (Gay et al., 2008; Wang et al., 2011). This allows us to make predictions based on the hypothesis of hybrid zone movement: there will be a tail of introgression in the wake of the moving zone combined with a shift of the peak in admixture linkage disequilibrium ahead of the movement. This contrasts with the predictions for a stable situation, where clines are symmetric and the peak of admixture linkage disequilibrium appears in the centre of the hybrid zone (Figure 1).

We test for a genetic footprint of movement in a hybrid zone in northwest France between two morphologically similar but genetically distinct species of toad, the common, *Bufo bufo* (Linnaeus, 1758) and the spined toad *B. spinosus* Daudin, 1803 (Arntzen, Recuero, Canestrelli, & Martínez-Solano, 2013; Recuero et al., 2012; Trujillo, Gutiérrez-Rodríguez, Arntzen, & Martínez-Solano, 2017). While *B. bufo* survived the last glacial maximum in Italy and the Balkans, *B. spinosus* survived in southern France and Spain (Arntzen et al., 2017). Based on four nuclear gene coding markers, twelve microsatellites, and two mitochondrial SNP markers, the width of the hybrid zone (30 km) compared to the dispersal distance (1.3 km/generation) suggested that selection against hybrids restricts gene flow through the hybrid zone (Arntzen et al., 2016). In the southeast of France, the *Bufo* hybrid zone was hypothesised to move southwards based on traces of introgression of one nuclear gene coding marker (of four tested), and discordance between nuclear, mtDNA, and morphological clines (Arntzen et al., 2017). In the northwest of France, a similar data set including microsatellite



**FIGURE 1** Schematic representation of an hypothesis of hybrid zone movement. In contrast to a stable hybrid zone (a) where the peak of admixture linkage disequilibrium ( $D'$ , y-axis, top plot) is situated in the hybrid zone centre and clines are symmetric (frequency, y-axis bottom plot, x-axis of both plots is distance along a transect), a moving hybrid zone (b) is expected to have a peak of admixture linkage disequilibrium shifted ahead of the hybrid zone and a tail of introgression in the wake of the hybrid zone ( $t_0$  is the previous hybrid zone position,  $t_1$  the current position). Dashed lines indicate the hybrid zone centre [Colour figure can be viewed at [wileyonlinelibrary.com](http://wileyonlinelibrary.com)]

data, showed introgression towards the north in two nuclear markers out of four, but was not analysed from the perspective of hybrid zone movement at the time (Arntzen et al., 2016). This scenario provides the opportunity to test for the hypothesis of hybrid zone movement in the northwest of France. We improve the resolution using 31 gene coding markers and one mtDNA marker to interpret interspecific gene flow in the light of the average effective selection on a locus, and place our findings in the context of results from the literature.

## 2 | MATERIALS AND METHODS

### 2.1 | Sample collection and preparation

All DNA extracts were available from previous studies (Arntzen, McAtear, Butôt, & Martínez-Solano, 2018; Arntzen et al., 2017, 2016; Arntzen, Wilkinson, Butôt, & Martínez-Solano, 2014). Eight individuals from four *B. bufo* reference populations and 12 individuals from seven *B. spinosus* reference populations, all positioned >500 km away from the centre of the hybrid zone (blue and red circles in Figure 2a,b), served to identify species diagnostic single nucleotide polymorphisms (SNPs) in a test run. Another 268 individuals from 29 populations, including transect populations (grey circles in Figure 2a,b) and additional samples from the reference populations were genotyped and included in the data set, to a total of 306 individuals. Distances between transect populations were measured in a straight line, which follows the direction of the transect in Arntzen et al. (2016), using a custom R script. Distances for reference populations were measured with GOOGLE EARTH PRO v.7.3.0 (Table 1 and Supporting Information Table S1).

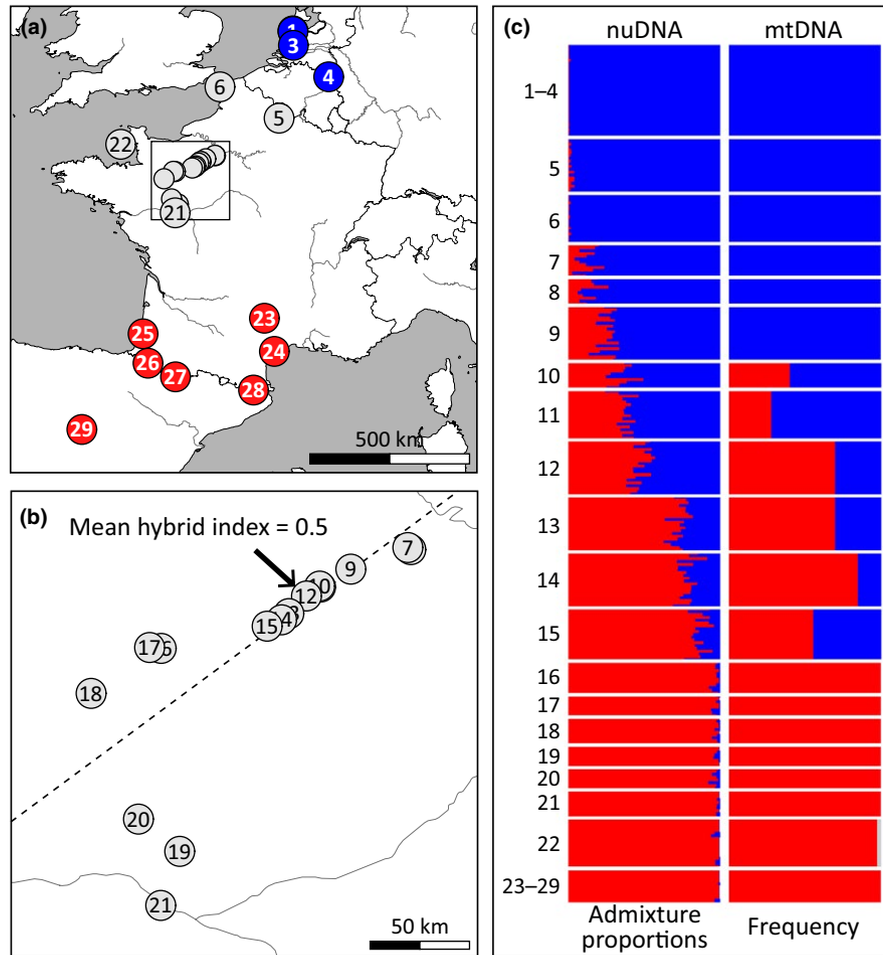
### 2.2 | SNP design based on transcriptome data

Two transcriptomes from liver tissue of a *B. bufo* individual from population 6 (Audresselles, France), and a *B. spinosus* individual from population 18 (Jublains, France) were sequenced commercially by ZF

Screens, Leiden, on the Illumina HiSeq 2000 platform. The data were filtered with TRIMMOMATIC v.0.35 software (Bolger, Lohse, & Usadel, 2014), and assembled with TRINITY v.2.1.1 (Grabherr et al., 2011; Haas et al., 2013). When using exons from transcriptome data for SNP design in expressed genes, paralogs (gene copies) should be excluded to avoid interpreting SNPs between paralogs as SNPs within orthologues (e.g., Ryyänen & Primmer, 2006). Exon boundaries need to be taken into account, as primers designed across exon boundaries are the main cause of genotyping assay failure (Wang et al., 2008). First, a BLAST search (Altschul, Gish, Miller, Myers, & Lipman, 1990) was used to identify paralogs within the *B. bufo* transcriptome assembly. Next, a pipeline for exon boundary identification (Niedzicka, Fijarczyk, Dudek, Stuglik, & Babik, 2016) was used, employing the well-annotated *Xenopus tropicalis* (Gray, 1864) genome (genome version JGI4.2). The *X. tropicalis* genome was also used to annotate markers (after testing for diagnostic differences between the two species, 30 of 31 markers eventually used were annotated, one remained unidentified and was named *exon\_1*). Finally, a second BLAST of the selected exons against the transcriptome of *B. spinosus* was used to exclude potentially undetected paralogs and to identify SNPs. A detailed description of the SNP design from transcriptome data is presented in the Supporting Information Appendix S1.

### 2.3 | SNP validation

Fluorescence-based genotyping (Semagn, Babu, Hearne, & Olsen, 2014) was used in the Kompetitive Allele-Specific PCR (KASP) genotyping system at the SNP genotyping facility of the Institute of Biology, Leiden University. Primer design, PCR setup, and data visualisation followed Arntzen et al. (2016). For 96 exon sequences, primers were designed with the Kraken software (LGC genomics, UK). After a first run, three SNPs were excluded as they failed for all individuals of one species, presumably due to a mutation at a primer binding site. Markers were initially considered diagnostic if they showed species-specific



**FIGURE 2** Map of Western Europe (a) with reference populations to determine diagnostic nature of markers for *Bufo bufo* (blue circles) and *B. spinosus* (red circles), and transect populations (grey circles). Panel (b) details the transect orientation (dashed line) and the position of hybrid zone populations. Panel (c) shows bar plots with individual admixture proportion (Structure Q scores) on the left and mtDNA allele frequency per population on the right. The grey bar refers to one missing mtDNA data point [Colour figure can be viewed at [wileyonlinelibrary.com](http://wileyonlinelibrary.com)]

alleles in the test data set. This resulted in 32 nuclear DNA SNPs (Supporting Information Table S2). Because of limitations due to plate dimensions in the KASP system, the remainder of the samples was analysed with 31 SNPs (by excluding the least well performing marker in the test data set, *scaf4*, for which six out of 20 reference individuals had missing data) and we added one previously published diagnostic mtDNA SNP (16S; Arntzen et al., 2016). We excluded five individuals from the final data set with more than 10 SNPs missing (presumably because of low quality DNA). After assessment of the final data set of 306 individuals, 29 of 31 nuclear SNPs were considered species-diagnostic, defined by minor allele frequencies of <5.0% in reference populations in the final data set (Supporting Information Table S3). Missing data amounted to 5.2% (Supporting Information Table S4).

## 2.4 | Hardy-Weinberg proportions and marker linkage

Because SNPs in genes are more prone to be under direct or indirect selection than noncoding DNA, we were cautious to exclude

any markers showing outlier behaviour. Signals of nonrandom mating or survival were tested for by calculating the Hardy-Weinberg proportions (heterozygote deficit and excess) with the R package *genepop* based on the program *GENEPOP* v.1.0.5 (Rousset, 2008). As the Bonferroni correction (Rice, 1989) can be overly conservative (e.g., Narum, 2006), we chose to account for independence of tests within markers ( $P_c$  for  $N = 31$ ). Deviations from Hardy-Weinberg proportions by heterozygote excess were not significant ( $P_c > 0.05$ ; Supporting Information Table S5). Deviations by heterozygote deficit were significant for the marker *egflam* in five populations ( $P_c < 0.05$ ), hence this marker was excluded in the HZAR cline fitting analysis and admixture linkage disequilibrium calculations (see below).

To infer independence of introgression, close physical or functional linkage of markers should be investigated. As some of the 31 nuclear markers are bound to be on the same chromosome (*Bufo* has 22 chromosomes; Olmo, Gargiulo, & Morescalchi, 1970; Olmo, 1973), we used a test of pairwise linkage disequilibrium (LD), indicative of close physical linkage of markers, based on the log likelihood ratio statistic (*G* test) with the R-package *genepop* (Rousset, 2008). One

**TABLE 1** Overview table with the number of individuals sampled per population, the distance in kilometres (km), and the mean hybrid index, pooled for the reference populations (1–4 for *B. bufo* and 23–29 for *B. spinosus*), and for each transect population (5–22)

Population	Individuals sampled	Distance (km)	Hybrid index (HI)
1	10		
2	10		
3	7	0	0.03
4	6		
5	20	174	0.06
6	18	335	0.03
7	12	468	0.16
8	10	469	0.17
9	20	508	0.28
10	10	530	0.38
11	18	530	0.39
12	20	541	0.50
13	20	555	0.74
14	20	561	0.79
15	19	571	0.86
16	12	648	0.96
17	8	644	0.96
18	10	689	0.97
19	8	680	0.98
20	8	695	0.96
21	10	705	0.99
22	18	791	1.00
23	2		
24	2		
25	2		
26	1	1,222	1.00
27	2		
28	1		
29	2		

pair of markers, *exon\_1* and *ttc37*, was found to be in significant pairwise LD in two populations ( $P_c < 0.05$ ; Supporting Information Table S6). As these populations were located in the hybrid zone, where we expect the effect of admixture linkage disequilibrium to increase the number of markers in pairwise LD, we kept the two markers in downstream analyses, but we checked that this marker pair did not distort the general patterns described below. To investigate functional linkage, a protein-interaction network was analysed with STRING v.10.5 (Szklarczyk et al., 2015) on the basis of the *X. tropicalis* genome, and 30 out of 31 annotated nuclear markers. A protein function description was recorded for each annotated marker (Supporting Information Table S7). The markers *brca2* and *rfc1* were found to be functionally linked. As these markers were neither deviating from Hardy-Weinberg proportions nor in pairwise linkage disequilibrium

within populations, they were included in all downstream analyses but, again, we checked that this marker pair did not distort the general patterns described below. Note that it remains possible that individual markers are under direct or indirect selection, and the current and future analyses using these markers need to take this into account.

## 2.5 | Population structure

Allocation of individuals to genetically defined groups was done using all 31 nuclear markers in STRUCTURE v.2.3.4 (Pritchard, Stephens, & Donnelly, 2000) with 10 independent chains of one up to 10 genetic clusters ( $K$ ), a burn in of 10,000, and a chain length of 25,000 under the admixture model following recommendations of Benestan et al. (2016). Other settings were left at default. Credibility intervals (95%) were estimated and convergence was checked comparing log likelihood and similarity of admixture proportion (Structure Q scores) between runs. The results were summarized with CLUMPAK (Kopelman, Mayzel, Jakobsson, Rosenberg, & Mayrose, 2015). The best value of  $K$  was determined using the Evanno method (Evanno, Regnaut, & Goudet, 2005; Supporting Information Table S8), and results were visualised using the R package POPHELPER (Francis, 2017). The admixture proportions are used in the cline analysis (see below) to approach an overall cline shape and position that summarizes all nuclear clines.

## 2.6 | Cline analyses

Classic equilibrium cline models, describing a sigmoid change in allele frequency or phenotype across the hybrid zone, were fitted with the R package HZAR (Derryberry, Derryberry, Maley, & Brumfield, 2014) for 30 nuclear markers, mtDNA, and admixture proportion ( $K = 2$ ), using a set of custom R scripts provided by G. Derryberry. Marker *egflam* was not analysed with HZAR as it was not behaving according to Hardy-Weinberg proportions. First, 30 maximum likelihood estimation searches were performed with random starting parameters, followed by a trace analysis of 60,000 generations on all models with a delta Akaike Information Criterion corrected for small-sample-size ( $\Delta AICc$ ) below 10. Fifteen model variants can be fitted in HZAR, based on all possible combinations of trait interval (allele frequency at the transect ends; three types) and cline shape. The different cline shapes represent either a single sigmoid curve, or a combination of (a) the central sigmoid portion and (b) shallower exponential decay curves at one or both ends (tails) with slopes shallower than expected from the central portion. We refer to these alternatives as “tail types” (five types). Tail types were fitted as an exponential tail to the left (L), exponential tail to the right (R), both tails exponential with independent parameters (B), both tails exponential but with the same parameters mirrored on the cline centre (M), and no exponential tails fitted (N). For cline tails estimated separately (B), we assessed tail slope ( $\tau$ ; shallow tails reflected by low parameter values) and distance from the cline centre where a tail started ( $\delta$ ; short distances reflected by low parameter values). When  $\tau$  and  $\delta$  are both low on one side and both high at the other side of the hybrid zone, introgression is asymmetric. Convergence of the

HZAR analysis was visually assessed in trace plots. Significance of cline model fits was determined by calculating  $\Delta\text{AICc}$  for each best model ( $\Delta\text{AICc} > 2$  compared to the second best model; Supporting Information Tables S9 and S10, Figure S1). Deviation from an equal frequency of left and right tails (as expected in a stable hybrid zone) was tested for all markers with a coincident cline centre with the Chi-square test for equal probabilities.

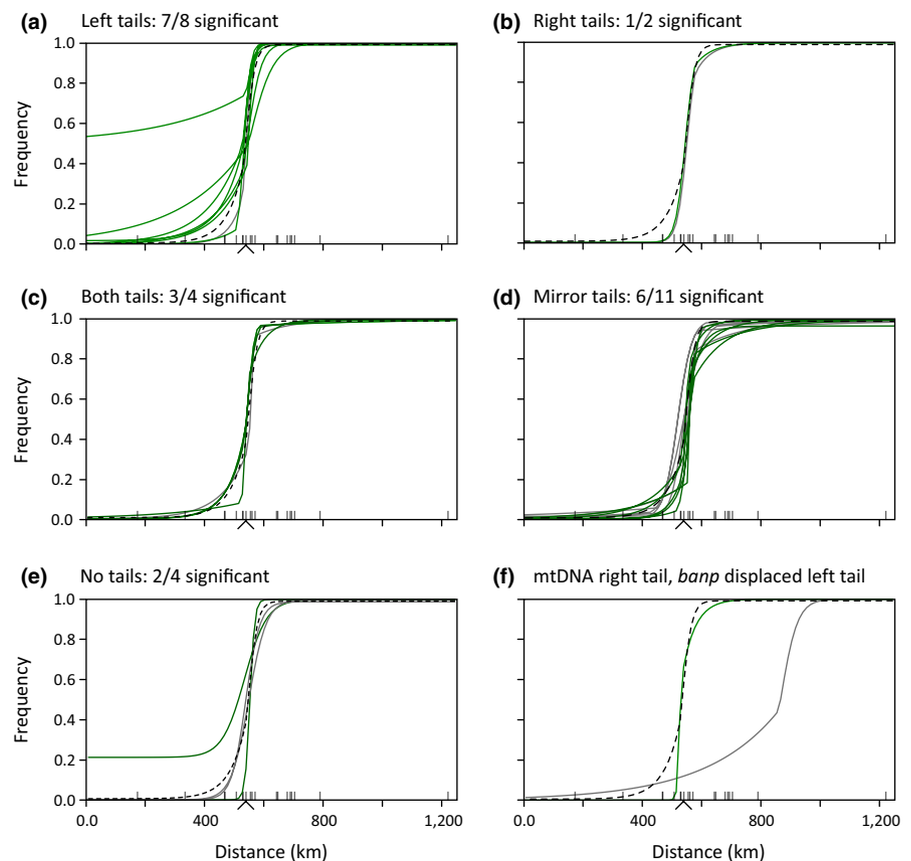
## 2.7 | Admixture linkage disequilibrium

Using variance in hybrid index (HI, see also Table 1), admixture linkage disequilibrium ( $D'$ ) can be calculated, and subsequently lifetime dispersal distance ( $\sigma$ ) weighted for individuals in aquatic (pre-) and terrestrial (post-metamorphosis) life stages, expected cline width under neutrality, and average effective selection on a locus ( $s^*$ ) can be calculated following Barton and Gale (1993). This was done using 29 nuclear markers, excluding marker *egflam*, which was not within Hardy-Weinberg proportions and marker *banp*, for which the cline centre was displaced compared to other clines (Supporting Information Figure S2). Average effective selection is the selection pressure on a locus at the zone centre due to direct selection or association with other loci under selection. A few input parameters were needed for these calculations (Supporting Information Table S11). A mean recombination rate between marker pairs of 0.4997 was calculated following formula (6) from Macholán et al. (2007), using the number of chiasmata per bivalent for *Bufo bufo* (1.95; Wickbom, 1945), and the number of chromosomes for *Bufo bufo*

( $N = 22$ ). A generation time of 2.5 years for *Bufo* toads at the latitude of the hybrid zone (mean of three years in females and two years in males; Hemelaar, 1988), and initial secondary contact at 8,000 years ago following Arntzen et al. (2016) were used as input parameters. The width of the hybrid zone was derived from a general sigmoid cline model following the “no-tails” formula in HZAR (Derryberry et al., 2014) fitted to the HI. At the cline centre,  $D'$  was estimated from its regression on  $p^*(1-p)$ , where  $p$  is the average over loci of the frequency of the *B. bufo* allele at a sample location. Means and 95% confidence intervals (CI) for cline width,  $D'$  at the centre, dispersal distance, expected cline width under neutrality, and effective selection were based on 1,000 bootstrap replicates of the original genotype data set (with replacement, maintaining original sample size; Supporting Information Table S11), using custom R scripts. The amplitude and position of the peak in  $D'$  were estimated by fitting a Gaussian curve following Gay et al. (2008).

## 2.8 | Hybrid index analysis

To visualise potential unequal contribution of both species to the genomic composition of admixed individuals, we used the R package Hlest (Fitzpatrick, 2012). This analysis scales the number of markers derived from each parental species ( $S$ ) to heterozygosity, which is calculated as the fraction of heterozygous markers with variants inherited from both parent species, within an individual ( $H_i$ ). We used 29 markers with an allele frequency difference between species more than 0.8 (Larson, Andrés, Bogdanowicz, & Harrison, 2013). To



**FIGURE 3** Best fitting clines for 30 nuclear markers and the mtDNA marker determined with HZAR. Frequencies of 0 and 1 represent pure *Bufo bufo* and *B. spinosus* genotypes. Left tail clines are in panel (a), right tail clines are in (b), both tail clines in (c), mirror tail clines in (d), and no tail clines are in (e). Panel (f) shows the mtDNA cline with a significant right tail and the nuclear marker *banp* with a significantly displaced cline centre and a non-significant left tail. Cline models that fit significantly better than the next best model ( $\Delta\text{AICc} > 2$ ) are shown by green lines and the others by grey lines. Sample localities on the x axis by inside ticks and the arrow (^) indicates the position of the hybrid zone centre based on the admixture proportion cline (dashed line) [Colour figure can be viewed at [wileyonlinelibrary.com](http://wileyonlinelibrary.com)]

assess unequal inheritance of mtDNA in hybrid offspring, we coloured the data points by mtDNA type.

### 3 | RESULTS

The number of genetic clusters ( $K$ ) supported by Structure was two, concordant with the two toad species (Supporting Information Table S8). Species admixture was recorded in populations 7–15. Individuals of populations 7–9 north of the hybrid zone centre were hybrid populations with a relatively high frequency of *B. spinosus* nuclear alleles when compared to the number of *B. spinosus* mtDNA haplotypes (Figure 2c). Credibility intervals of the admixture proportion estimates are shown in Supporting Information Figure S3.

HZAR clines were partitioned according to tail type (Figure 3, for a plot including all clines see Supporting Information Figure S2, Tables S9 and S10). Eight nuclear marker clines best fitted a left tail of introgression (northwards into *B. bufo*) and for seven markers the fit was significantly better than alternative models (Figure 3a). Two nuclear marker clines had a right tail of introgression (southwards into *B. spinosus*), of which one had a significantly better fit than alternative models (Figure 3b). For four nuclear marker clines, both tails were exponential with independent parameters, of which three had a significantly better fit than alternative models (Figure 3c). For these clines, tail slope ( $\tau$ ) and distance from the cline centre where the tail started ( $\delta$ ) showed an increase in one parameter (e.g.,  $\delta$ ) and a decrease in the other (e.g.,  $\tau$ ) in all cases. Such results are not straightforward to interpret in terms of directional introgression and these clines were not taken into account when assessing unidirectional introgression. For eleven nuclear markers, clines had mirrored tails of which six had a significantly better fit than alternative models (Figure 3d). For three nuclear markers, clines had no tails, and one marker had a significantly better fit than alternative models (Figure 3e). All nuclear markers had cline centres within 30 km of the admixture proportion cline centre (positioned at 539 km) except for *banp*, with a cline centre >300 km to the south, inside the range of *B. spinosus* (Supporting Information Figure S2). This cline had a nonsignificant left tail into *B. bufo* (Figure 3f). The mitochondrial marker had a significant right tail into *B. spinosus* (Figure 3f). The functionally linked markers *brca2* and *rfc1* had similar cline models with mirrored tails and overlapping 95% credible cline regions for centre and width, and showed no pattern of introgression deviating from the admixture proportion cline. In summary, seven nuclear markers showed significant introgression of *B. spinosus* into *B. bufo* by cline discordance, whereas two nuclear markers showed significant introgression the other way around, one by cline discordance and one by displacement. The inequality of introgression was nonsignificant (Chi-square test for equal probabilities,  $\chi^2 = 2.78$ ,  $df = 1$ ,  $p = 0.096$ ).

Populations 9–15 within the hybrid zone had higher values of  $D'$  than populations outside the hybrid zone (Figure 4). This is reflected by an increase in pairwise linkage disequilibrium among loci (Supporting Information Table S6). Populations 7, 8, and 10 north of the hybrid zone

centre, and beyond the area of rapid allele frequency change, had larger  $D'$  with larger positive deviations from zero than other populations. However, it was not possible to fit a Gaussian curve through these data points, probably because of the large variation in  $D'$ , and the position and amplitude of the peak of  $D'$  remain undetermined.

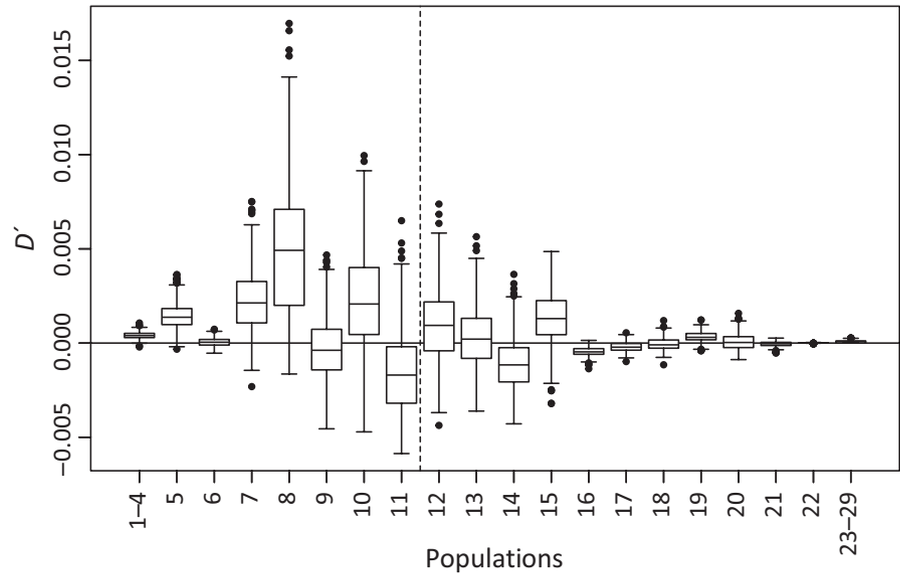
Estimated lifetime dispersal ( $\sigma$ ), based on  $D'$  at the cline centre, was 2.2 km generation<sup>-1/2</sup> (95% CI 0.6–3.7). The observed cline width based on  $HI$  was 114 km (95% CI 103–132). Assuming no selection against hybrids, the expected cline width would be 312 km (95% CI 91–532; Supporting Information Table S11). The average effective selection on a locus ( $s^*$ ) was 0.0017 (95% CI 0.0001–0.0040). Despite variation among clines suggesting weak coupling, the presence of an admixture LD peak supports the use of  $s^*$  as an average value to describe the hybrid zone. Hlest showed a continuum of *B. bufo* to *B. spinosus* genotypes in which hybrids possessing mtDNA haplotypes typical of either *B. bufo* or *B. spinosus* were about equally frequent (Supporting Information Figure S4). The peak of ancestry is at 50% heterozygosity, indicating no asymmetry in the contribution of both species to the hybrid population.

### 4 | DISCUSSION

The features describing the *Bufo* hybrid zone in northwest France in the current study are broadly in line with previous descriptions, but provide more resolution and estimates of key parameters. We found that individual marker cline positions and shapes vary (e.g., Supporting Information Figure S2), widening the admixture proportion cline. We inferred weak effective selection per locus against hybrids ( $s^* = 0.0017$ ), but with low precision of the estimate. Even when effective selection is low and clines are wide, a moving hybrid zone may stagnate at an ecotone, limiting further movement of the zone (Barton & Hewitt, 1985; Buggs, 2007; Endler, 1977; Gompert, Mandeville, & Buerkle, 2017; Moore, 1977). The section of the *Bufo* hybrid zone studied here appears to be located at a weak ecotone in a hilly landscape formed by the “Collines de Normandie”, with *B. bufo* at a lower and *B. spinosus* at a higher altitude (Arntzen et al., 2016). The genetic estimate of lifetime dispersal of 2.2 km generation<sup>-1/2</sup> diverges from the average maximum observed in field studies, but the field estimate is within the CI for the genetic estimate (1.3 km; Smith & Green, 2005; Daversa, Muths, & Bosch, 2012; Trochet et al., 2014). In amphibians, dispersal distance based on genetic data is more often found to be higher compared to field studies, because mark-recapture, seasonal, or field studies covering a small area are at risk of underestimating (rare) long-distance dispersal (Smith & Green, 2005).

The hybrid zone was calculated to be 114 km wide, based on the hybrid index for 30 nuclear markers, and width was 63 km based on the cline of the admixture proportion for 31 nuclear markers. A narrower width (30 km) was earlier recorded based on the first PCA axis of 12 microsatellites, whilst in the same study cline width for four nuclear markers ranged from 27 to 91 km, and morphological cline widths exceeded 200 km (Arntzen et al., 2016). Microsatellites are noncoding, have higher mutation rates, and they are more suitable

**FIGURE 4** Admixture linkage disequilibrium ( $D'$ ) recorded over a transect from the north (*Bufo bufo*, populations 1–4) to the south (*B. spinosus*, populations 23–29). Note that the horizontal axis does not represent linear geographical distance. The dotted vertical line is the position of the hybrid zone centre inferred from the admixture proportion cline. Estimates for  $D'$  are obtained by bootstrap analysis and shown by box-and-whisker plots



for investigating small scale evolutionary processes, therefore they may reflect more recent demographic processes (Ellegren, 2000). This poses the question what is causing a more spatially restricted transition in the microsatellite markers as opposed to a wider transition in nuclear gene markers, as one would expect stronger selection against hybrids and thus narrower clines in markers from coding regions than in microsatellites. Including more markers would cause the hybrid index or admixture proportion clines to get wider, because drift causes the centres of individual marker clines to vary. The larger number of markers in this paper may thus explain the wider zone currently reported compared to the previously reported width based on less markers. Confidence intervals overlap for many of these width estimates and all widths are large relative to dispersal. This indicates the general conclusion of weak selection is robust.

We examined patterns of gene flow and admixture linkage disequilibrium to test the hypothesis of movement of the hybrid zone in two common toad species (*B. bufo* and *B. spinosus*) in northwest France. Introgression from *B. spinosus* into *B. bufo* was more frequent than introgression the other way around, a result that was found before with four nuclear markers (Arntzen et al., 2017, 2016). Disregarding whether cline model fits are significant, eleven markers show introgression from *B. spinosus* into *B. bufo* and two show introgression the other way around, a significant difference (Chi-square test for equal probabilities,  $\chi^2 = 6.23$ ,  $p = 0.013$ ). Cases where introgression is not explicit can also point out methodological difficulties, such as determining unidirectional introgression from cline shape parameters. To better compare the asymmetry of introgression between individual markers, future methods could compare e.g., the area underneath and above the geographic cline on both sides of the cline centre. Further simulations are needed to provide clear expectations for stable and moving hybrid zones.

Three populations with elevated admixture linkage disequilibrium ( $D'$ ) suggest a peak of early-generation hybrid offspring north of the hybrid zone. Fitting a Gaussian curve through  $D'$  proved impossible, as there was too much variance within the data, especially

in populations 5, 9 and 11. The Gaussian curve is the shape of the peak expected under a symmetrical cline model (Gay et al., 2008). The absence of a clear peak can be caused by asymmetry in the cline model or insufficient sample size of individuals, populations, or markers, or a combination of both. More data should be generated to be able to determine the shape and position of the peak with higher precision. While the unidirectional introgression is in line with southward movement of the hybrid zone (Buggs, 2007), the northern position of a peak in  $D'$  is in line with movement in the opposite direction (Gay et al., 2008; Wang et al., 2011). Since our results do not fully support the original hypothesis of (southward) hybrid zone movement in *Bufo*, alternative explanations have to be considered, in particular for the position of the peak in admixture linkage disequilibrium, for which we compare our result with other case studies.

Concordant and narrow clines suggest that a hybrid zone is maintained by a balance between strong selection and dispersal, as opposed to wide and displaced clines indicating that selection against hybrids is low (Barton & Gale, 1993). By surveying the hybrid zone literature, we found eight studies with empirical data on effective selection (i.e.,  $s^*$ ) and cline centre and shape in a wide variety of organisms (Supporting Information Table S12; Mallet et al., 1990; Szymura & Barton, 1991; Phillips, Baird, & Moritz, 2004; Macholán et al., 2007; Kawakami, Butlin, Adams, Paull, & Cooper, 2009; Carneiro et al., 2013; Baldassarre, White, Karubian, & Webster, 2014; Hollander, Galindo, & Butlin, 2015; Wielstra, Burke, Butlin, & Arntzen, 2017). Six studies report substantial effective selection (0.19–0.37) with mostly concordant, coincident and steep clines, and three studies report low average effective selection (0.007–0.011) with multiple discordant and displaced clines. Our results for *Bufo*, with low effective selection and substantial cline variability, fall into the latter category.

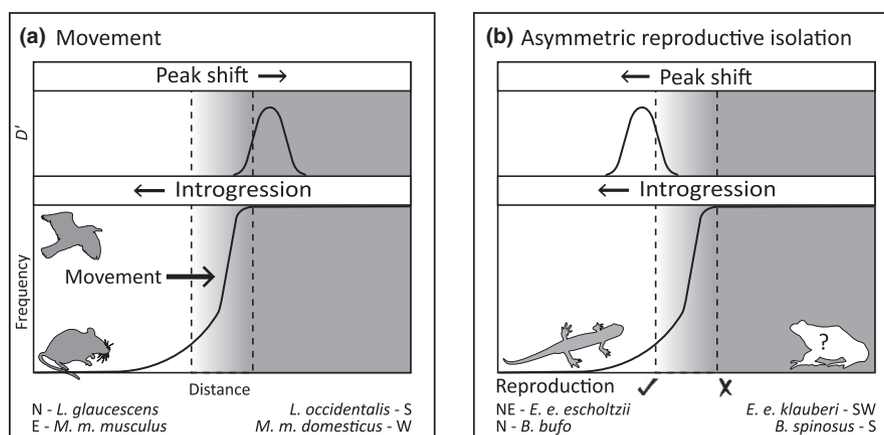
We also consulted the literature on the direction of introgression and the location of elevated levels of admixture linkage disequilibrium in hybrid zones. Two studies document a peak in admixture linkage disequilibrium opposite to where introgression takes place (Figure 5a), and one study documents a peak of admixture linkage disequilibrium

and introgression on the same side (Figure 5b). In the hybrid zone of the gull species *Larus glaucescens* (Naumann, 1840) and *Larus occidentalis* (Audubon, 1839), genotypic and phenotypic clines showed introgression towards the north, and a peak in admixture linkage disequilibrium towards the south, indicating a southward movement of the hybrid zone (Figure 5a; Gay et al., 2008). In the hybrid zone of the house mouse subspecies *Mus musculus musculus* Linnaeus, 1758 and *M. m. domesticus* Schwarz & Schwartz, 1943 introgression to the east and a peak of admixture linkage disequilibrium to the west, supported a westward movement of the hybrid zone (Figure 5a; Wang et al., 2011). Conversely, in the hybrid zone of the salamander subspecies *Ensatina eschscholtzii eschscholtzii* Gray, 1850 and *E. e. klauberi* Dunn, 1929, unidirectional introgression coincides with a peak of admixture linkage disequilibrium (Figure 5b; Devitt, Baird, & Moritz, 2011). This result was explained by asymmetric pre- or postzygotic isolation. On the side of the hybrid zone where reproduction is reduced, introgression and admixture of genetically distinct individuals are absent, whereas at the other side where hybridisation succeeds, introgression and admixture of genetically distinct individuals occur. A deficit of hybrids with *E. e. eschscholtzii* mtDNA confirms that offspring of female *E. e. klauberi* and male *E. e. eschscholtzii* are successfully reproducing, whilst offspring of other parental combinations occur rarely.

When comparing patterns of introgression between hybrid zones of different biological systems, dispersal rate and effective population size need to be taken into account (Canestrelli et al., 2016; Ravinet et al., 2017). Introgression generally increases when dispersal rate is high, because more genetically distant populations interbreed (Nichols & Hewitt, 1994). The effect of a higher dispersal on introgression is greater for a narrow hybrid zone where selection against hybrids is high, than for a wide zone where selection against hybrids is low, because in a narrow zone the distance between genetically distinct individuals is smaller (Barton & Gale, 1993).

Therefore, we chose to express dispersal in terms of the number of generations needed to cross the hybrid zone, as this summarises width in relation to dispersal, and thus also in relation to selection against hybrids. To include the effect of genetic drift in the comparison of hybrid zones, effective population size can be used. When effective population size is low, an increase in the effect of random genetic drift obscures genetic patterns of demographic and evolutionary history, and clines are predicted to be more randomly distributed in the landscape and vary in shape, despite a high selection against hybrids (Polechová & Barton, 2011). The *Larus* hybrid zone would take five generations of unimpeded dispersal to cross (Figure 5a; Gay et al., 2008). Populations consist of 10,000–100,000 individuals (Crochet et al., 2003). The *Mus* hybrid zone would take about 60 generations to cross (Figure 5a; Raufaste et al., 2005; Macholán et al., 2007; Wang et al., 2011). The effective population size of mice is 500 to 5,000 individuals (Pocock, Hauffe, & Searle, 2005). The *Ensatina* hybrid zone would take seven generations to cross (Figure 5b; Staub, Brown, & Wake, 1995; Devitt et al., 2011). In high quality habitat *Ensatina* population density may be 1,300 individuals/ha with effective population size unknown (Rosenberg, Noon, Megahan, & Meslow, 1998; Stebbins, 1954).

The *Bufo* hybrid zone would take about 60 generations to cross, and dispersal capacity seems comparable to *Mus*. However, the effective dispersal distance in *Mus* is thought to be underestimated by population structure and migration studies, because effective genetic dispersal in mice is inflated by (human-mediated) long-distance movements and frequent extinction-recolonization effects (Barton & Hewitt, 1985; Macholán et al., 2007; Slatkin, 1985). Compared to *Bufo*, the effective population sizes in *Larus* and *Mus* are one to three orders of magnitude higher. Toads breed in large water bodies, and adult *Bufo* population sizes regularly consist of 2,500–5,000 individuals, but effective population sizes are about two orders of



**FIGURE 5** Schematic representation of inferred hybrid zone movement in (a) the gulls *Larus glaucescens* and *L. occidentalis* (Gay et al., 2008), and the house mice *Mus musculus musculus* and *M. m. domesticus* (Wang et al., 2011), and asymmetric reproductive isolation in (b) the salamanders *Ensatina eschscholtzii eschscholtzii* and *E. e. klauberi* (Devitt et al., 2011), and possibly also in the toads *Bufo bufo* and *B. spinosus* in the present study. In Panel (b), the tick mark shows where reproduction can take place and the cross shows where reproduction cannot take place. The y-axis in the top panel shows admixture linkage disequilibrium ( $D'$ ), the y-axis in the bottom panel shows the frequency, and the x-axis in both panel shows distance along the transect. Cardinal directions are given by N, E, S and W. The question mark in panel b refers to the uncertainty of unidirectional introgression and of the magnitude and position of the peak of  $D'$  in the *Bufo* hybrid zone [Colour figure can be viewed at [wileyonlinelibrary.com](http://wileyonlinelibrary.com)]

magnitude lower (Scribner, Arntzen, & Burke, 1997). Population sizes in *Bufo* may be more comparable to *Ensatina*, considering that effective population sizes are often much smaller than census population sizes in amphibians (Funk, Tallmon, & Allendorf, 1999; Vences & Wake, 2007; Zeisset & Beebee, 2003). A combination of low dispersal and small population sizes may impede introgression and cause patterns of admixture linkage disequilibrium to be less pronounced in *Bufo* than in the other hybrid zones.

Drift is strong in populations with low dispersal and small population sizes, generating random variation in allele frequency in different populations. This variation causes clines to differ in shape and position more than would be predicted based on the width of clines of individual alleles (cline wobbling; Polechová & Barton, 2011). Cline wobbling increases the width of the hybrid index (or admixture proportion) cline. A low number of markers, sampling a small portion of the variation in cline shapes and position, may therefore be insufficient to assess patterns of admixture linkage disequilibrium. Further increasing the number of individuals sampled and the number of markers employed would increase our ability to test the hypothesis of hybrid zone movement. It is not currently known how sensitive the methods employed here are to the numbers of individuals or markers included in the data set. Further investigation through simulations could help to improve our understanding of the limitations of these methods. It will be interesting to see if a higher number of markers generated randomly across the genome, such as RAD sequencing data (Baird et al., 2008), will behave similar or different to these gene-coding markers, as gene-coding markers may be more prone to local selection forces in small populations.

Asymmetric pre- or postzygotic isolation, such as displayed in *Ensatina*, seems most congruent with the co-occurrence of introgression and a peak of admixture linkage disequilibrium seen in *Bufo* (Figure 5b). However, we find only slight evidence for pre- or postzygotic isolation in the *Bufo* hybrid zone. First, we find marginally higher introgression from *B. spinosus* into *B. bufo* than the other way around. Second, hybrids with both parental types of mtDNA are about equally frequent, and the peak of ancestry is symmetric and centred at 0.5 heterozygosity (Supporting Information Figure S4), whereas in *Ensatina* almost all hybrid individuals carried mtDNA of only one parental type. Third, low effective selection within the *Bufo* hybrid zone allows the unimpeded exchange of genes and previous studies showed no indication of asymmetric incompatibility (Arntzen et al., 2016; Trujillo et al., 2017). The only evidence in favour of asymmetric isolation besides the position of the admixture linkage disequilibrium peak, is the relatively high introgression of *B. bufo* mtDNA into *B. spinosus* (e.g., right tail of introgression in mtDNA). Therefore, asymmetric pre- or postzygotic isolation might be weak, with hybridisation inhibited slightly more on the *B. spinosus* side. But this result can also be explained by other processes, such as sex-biased dispersal or adaptive introgression (Currat & Excoffier, 2005; Toews & Brelsford, 2012).

In conclusion, we cannot reject the null hypothesis of a stable *Bufo* hybrid zone in northwest France, and low levels of dispersal and random genetic drift due to small population sizes are

important in shaping the patterns of introgression. Based on earlier studies, this hybrid zone was predicted to move southwards (Arntzen et al., 2016, 2017). Our data set shows similar characteristics to those of previous studies (e.g., northward introgression), but with additional analyses we showed asymmetric reproductive isolation may provide a more important driver of asymmetric introgression than hybrid zone movement. However, multiple processes appear to be at play in shaping the *Bufo* hybrid zone. One might imagine a situation where asymmetric reproductive isolation and hybrid zone movement co-occur where the strongest process may overrule any genetic patterns related to the other process. In addition, small population sizes and other demographic factors such as dispersal distance per generation may cause patterns to become obscured. With 31 nuclear SNPs, we sampled only a tiny portion of the 6 Gbp *B. bufo* genome (Vinogradov, 1998). Further increasing the number of individuals sampled and the number of markers employed can possibly increase our ability to test the hypothesis of hybrid zone movement. It is not currently known how sensitive the methods presented here are to the number of individuals or markers included in the data set. Further investigation through simulations could help improve our understanding of the limitations of these methods. Looking back on past literature on hybrid zone movement in various organisms, it will be interesting to see if the hypotheses previously inferred still stand with new data and new analytical approaches. The *Bufo* hybrid zone exemplifies the complex influences of interspecific hybridization on genomic composition and can be used to test various hypotheses of introgression and speciation.

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## AUTHOR CONTRIBUTIONS

I.v.R., B.W., and J.W.A. designed the study and collected data. I.v.R. and R.B. analysed the data. IvR. wrote the manuscript, with input from all coauthors.

## DATA ACCESSIBILITY

Supplemental text contains details of the SNP design: Appendix S1. Supplemental tables are available in: Tables S1–S12. Supplemental

figures are available in: Figures S1–S4. In- and output of analyses, and custom R scripts are freely available online: Dryad link (sponsored by MEC): <https://doi.org/10.5061/dryad.rg18034>.

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## SUPPORTING INFORMATION

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