

GEOGRAPHIC VARIATION IN ASYMMETRIC COMPETITION: A CASE STUDY WITH TWO LARVAL ANURAN SPECIES

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Abstract. The outcome of interspecific competition may be modulated by a large number of factors, both biotic and abiotic. In this paper, we examined experimentally the effects of geographic variation, population density, and abiotic stress on the outcome of interspecific competition between two species of larval anurans. Three different populations of natterjack toad (*Bufo calamita*) were set to compete with common toad (*Bufo bufo*) in replicated artificial ponds under different combinations of osmotic stress (freshwater or brackish water) and intraspecific density (high and low). Among the *B. calamita* populations used, only one of them is sympatric to *B. bufo* and inhabits a freshwater environment. The two other populations, allopatric to *B. bufo*, occupy freshwater and brackish water environments, respectively. Survival, length of the larval period, and size at metamorphosis were analyzed at both the specific and the populational interaction levels. Intra- and interspecific competition did occur and generally resulted in a decrease in survival rates, an increase in the duration of the larval period, and a decrease in size at metamorphosis. Water salinity also prolonged the larval period and interacted with intra- and interspecific density, intensifying its effects. Interspecific competition was asymmetric, since *B. bufo* was competitively superior to *B. calamita*, causing larger effects on *B. calamita* than those experienced by *B. bufo*. However, the three populations of *B. calamita* showed different competitive effects and responses when they were set to compete with *B. bufo*. The population sympatric to *B. bufo* had the stronger competitive effect and response, causing a strong impact on survival, growth, and developmental rate of *B. bufo*. The allopatric population inhabiting the brackish environment also had a marked effect on *B. bufo* survival, although only at high density. The other allopatric population only had a significant effect on the size at metamorphosis of *B. bufo*. The impact of *B. bufo* on *B. calamita* was greater for the allopatric populations than for the sympatric one. This pattern suggests that the nature and outcome of ecological interactions are a function of environmental conditions and population history.

Key words: abiotic factors; amphibians; Bufo; geographic mosaic model; geographic variation; interspecific competition; intraspecific competition; local adaptation.

INTRODUCTION

Populations of species with a wide geographic distribution are likely to be exposed to a vast array of ecological conditions imposed by varying abiotic conditions and biotic interactions. The geographic mosaic theory states that this heterogeneity conforms to a mosaic of local selection pressures of varying intensities (hot spots and cold spots) that, together with gene flow, shape the spatial distribution of traits for that species (Thompson 1994, 1999, Gomulkiewicz et al. 2000). One of the ways of investigating the geographic mosaic theory is testing whether coevolved traits of an interaction differ among communities (Thompson 1999). However, the identity of the species that integrate a given community, and how close they are to their respective optimal conditions, is constrained by the set of abiotic factors acting upon the system, according to

the species' particular niches and tolerance limits (Hoffmann and Parsons 1991, Warner et al. 1991).

Competition is considered to be one of the most important determinants of the structure and dynamics of biological communities (Connell 1983, Schoener 1983, Underwood 1986). The outcome of competitive interactions is often modulated to a great extent by factors such as the abiotic conditions (Dunson and Travis 1991, 1994, Rowe and Dunson 1995, Newman 1998) or the relative intensity of predation (Morin 1983a, b). Amphibian larval guilds have been a fertile ground for the study of community structuring, interspecific interactions, and effects of environmental conditions on the outcome of interactions (Morin 1983a, Dunson and Travis 1991) because of their sensitiveness to environmental variation (both biotic and abiotic), and their relative easiness to handle experimentally. The combination of high population densities and relatively low niche differentiation makes it likely that interspecific competition takes place in tadpole assemblages (Alford 1999). Interspecific competition may be the main agent

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determining community structure in ponds of intermediate permanence, while in extremely ephemeral and permanent ponds the relative effect of abiotic factors (essentially pond duration) and predation, respectively, may prevail over competition (Faragher and Jaeger 1998). Both intra- and interspecific competition in amphibians are typically density dependent (Wilbur and Collins 1973, Travis 1980, Warner et al. 1993, Tejedo and Reques 1994, Newman 1998). In that sense, the study of interspecific competition not only is improved by the simultaneous study of intraspecific density effects, but it seems unreasonable to decouple the former from the latter, since the similarity of requirements for basic resources among conspecific individuals must be greater or at least as great as between species (Underwood 1986, Parsons 1996).

Several abiotic factors affect the outcome of intra- or interspecific competition among anuran larvae, including water pH (Warner et al. 1991, 1993, Pehek 1995), pond duration (Tejedo and Reques 1994, Reques and Tejedo 1997), and temperature (Newman 1998). Another stress factor for amphibian tadpoles is water salinity (Balinsky 1981, Shoemaker et al. 1992), but to date this stressor has not been explored as potential modulator of competitive interactions among amphibian species. Inland brackish water environments are relatively frequent in the Mediterranean area and several amphibian species breed in them in spite of the osmotic stress induced. One such species is *Bufo calamita*, long considered a euryhaline toad (Balinsky 1981). Osmotic stress has been experimentally demonstrated to induce high mortality and a strong decrease in developmental rates in *B. calamita* embryos and larvae (Beebee 1985), directly and indirectly reducing fitness through reducing survival and opportunities to reach metamorphosis before pond desiccation (I. Gómez-Mestre and M. Tejedo, unpublished data). Also populational variation in tolerance to osmotic pressure has been observed in *B. calamita* (I. Gómez-Mestre and M. Tejedo, unpublished data). In turn, another bufonid species that largely overlaps in its distribution, *Bufo bufo*, is not found breeding in these kind of environments and thus we predict that the strength of biotic interactions between these two species may be affected by changes in osmotic conditions.

Evidence for geographic variation and local adaptation of life history traits in amphibians, reflecting a geographic mosaic of selection pressures, is accumulating (Berven 1982a, b, Figiel and Semlitsch 1990, Harris et al. 1990, Bernardo 1994, Kurzava and Morin 1994, Fauth 1998, Storfer and Sih 1998, Merilä et al. 2000). If substantial population differentiation occurs, the geographic-mosaic theory of coevolution predicts some degree of geographic variation in the intensity and outcome of interspecific interactions as well. To our knowledge, this aspect has been much less studied in amphibians and information is still scarce (Fauth 1998).

In this paper we examined the outcome of competitive interactions among *Bufo bufo* and three different populations of *Bufo calamita* during the larval phase using an array of artificial mesocosms. Both species range across Western Europe and *B. bufo* has been found to be competitively superior in some British populations (Denton and Beebee 1994, Bardsley and Beebee 1998a). In southern Spain the two species are widespread and sympatric in many places. Reproduction is almost synchronic although *B. bufo* is an earlier breeder, preceding its congeneric by around one month. Both species breed in temporary habitats that differ in durability, those used by *B. bufo* being longer lasting than the ephemeral ponds used by *B. calamita*. However, rainfall is highly variable and affects the availability of breeding ponds so that larvae from both species often co-occur in the same ponds. Thus, there is potential for interspecific interactions during the larval phase. The three populations of *B. calamita* used in the experiments differed in their recent natural history with regard to the factors being analyzed, i.e., water salinity and syntopy with *B. bufo*.

The main hypotheses were (1) that the two bufonid species differed in their competitive ability and osmotic stress tolerance; and (2) different natterjack populations would respond differently in competitive interactions with common toads and in their tolerance to brackish water environments, thus reflecting microgeographic variations in the local selective factors. These hypotheses generated the following predictions: (1) There would be competitive dominance of *B. bufo* on *B. calamita*, but it would be less under brackish water and high-density conditions because *B. bufo* does not occur either in brackish environments (probably as a consequence of a poor tolerance to osmotic stress) or in crowded, ephemeral ponds, like *B. calamita* does; (2) The *B. calamita* population syntopic to *B. bufo* would exhibit a higher competitive response (sensu Goldberg and Landa 1991) to interspecific competition than would allotopic populations; and (3) The *B. calamita* population coming from a brackish environment would show a higher osmotic tolerance than those coming from freshwater ones.

METHODS

Source populations

Three populations of *B. calamita* located in the province of Córdoba, southern Spain, were chosen for the experiment because of their different historical relationship with the factors under study. The first population, Fresh 1, breeds in the same freshwater stream as the chosen *B. bufo* population. The other two populations do not naturally coexist with *B. bufo*. At least during 15 yr of surveys no *B. bufo* were observed breeding in these ponds or in the near surroundings. These two allotopic populations, Fresh 2 and Saline, breed in freshwater and brackish water ponds, respec-

tively. All these populations are separated by distances beyond the dispersal ability of adults. Fresh 2 is ~80 km away from both Saline and Fresh 1 while the two latter are separated by 30 km. However, there are other temporary breeding ponds in between that could allow gene flow among these populations.

The three aquatic systems are temporary. The Saline population breeds in a brackish lagoon, the Laguna de los Jarales reserve, of ~2.6 ha with a maximum depth of 1.2 m. This lagoon is moderately mineralized, showing a salinity range over the last 15 yr of between 2.3 and 18 g/L. The Fresh 1 population breeds in a stream running across a grassland plain at 950 m above sea level. The Fresh 2 population breeds in a mosaic of small and shallow ponds in Sierra Morena (see description in Tejedo 1992).

The experiment was conducted in the Reserva Integral de la Laguna Amarga, located near the populations of Saline and Fresh 1. Six recently laid clutches of *B. bufo* eggs were collected between 11 and 17 January 1998. During the night of 2 February, after a period of heavy rain, we collected six clutches of *B. calamita* from Fresh 2 and another six from Saline that were also brought to the laboratory. On 9 February we collected six samples from a number of *B. calamita* clutches in Fresh 1 that were in a similar developmental stage to those from Fresh 2 and Saline. A total of 500 larvae from each of three *B. calamita* populations and another 800 larvae from the *B. bufo* population were removed and reared in plastic trays until they reached developmental stage 25 (Gosner 1960). All *B. calamita* larvae from all three populations were synchronized, while *B. bufo* larvae were 15–20 d older than *B. calamita* by the time the experiment started.

Experimental design

We measured the effects of population density and osmotic stress on survival, size, and time to metamorphosis on assemblages of *B. calamita* and *B. bufo*. The design consisted of a three-factor fractional factorial design with water salinity, intraspecific density, and presence of heterospecifics as main factors. Densities of 15 and 30 tadpoles per enclosure simulated the low and high intraspecific densities for *B. bufo* and each of the *B. calamita* populations. Presence or absence of competitors was modulated following an additive design rather than following a substitutive one (Underwood 1986), implying that heterospecifics were added while density of the former species remained equal to the density set for the absence of competitors (15 + 15 or 30 + 30 tadpoles per enclosure, respectively, depending on the density level). Finally, two levels of water salinity, fresh and brackish, were established. Both types of water were taken from natural breeding ponds and had a total salinity of <0.4 and 5 g/L, respectively. The design implied 28 treatments that were arranged in a randomized block design with four rep-

TABLE 1. Experimental matrix of all possible interactions among density, competition, and salinity stress.

<i>Bufo calamita</i> origin and density (individuals/ enclosure)	<i>Bufo bufo</i> density (individuals/enclosure)		
	0	15	30
A) Freshwater			
Saline			
0	...	7	8
15	1	9	...
30	2	...	10
Fresh 2			
0	...	7	8
15	3	11	...
30	4	...	12
Fresh 1			
0	...	7	8
15	5	13	...
30	6	...	14
B) Brackish water			
Saline			
0	...	21	22
15	15	23	...
30	16	...	24
Fresh 2			
0	...	21	22
15	17	25	...
30	18	...	26
Fresh 1			
0	...	21	22
15	19	27	...
30	20	...	28

Notes: The density columns and rows refer to the intraspecific density (number of tadpoles) set at each treatment. The numbers in the matrix designate the cells (treatments) to be used in the contrast analysis in order to test specific effects. Cells with the same value represent the same experimental treatment. For instance, the cell numbered 8 represents the replicates belonging to the treatment in which *B. bufo* is set at high intraspecific density but with no interspecific tadpoles in freshwater. A contrast for the intraspecific density effect in *B. calamita* contrasted cells 1, 3, 5 and 15, 17, 19 against 2, 4, 6 and 16, 18, 20.

licates and a total of 112 experimental units. Table 1 shows all combinations in the design.

Experimental units consisted of square plastic boxes 0.50 × 0.37 × 0.28 m as artificial ponds located outdoors and were filled with 40 L of water a week before the beginning of the experiment. Temperature, water levels, and salinity were periodically checked and salinity (measured with an electronic conductivity meter Omega CDH-46, Omega, Stamford, Connecticut, USA) was adjusted within and among blocks. No strict salinity was fixed. We allowed it to increase as water evaporated from the boxes and restrained it so that it would not surpass 10–11 g/L, adding a similar amount of freshwater to all boxes. The boxes remained uncovered, and care was taken to eliminate any potential predators that accidentally entered the boxes. Tadpoles were fed 0.15 g of rabbit chow twice a week regardless

of tadpole density. Three censuses were made during the experiment and prior to metamorphosis. Two boxes had to be discarded for analyses due to the attack of a viperine snake (*Natrix maura*) that preyed upon the tadpoles.

Metamorphic toads first began to emerge on 2 April for *B. bufo*, and on April 4th for *B. calamita*. Tadpoles were considered to be in metamorphosis when at least one forelimb protruded, i.e., when they reached Gosner stage 42 (Morin 1983a, Tejedo and Reques 1994). After the occurrence of the first metamorph, the boxes were checked daily and metamorphs removed and kept in the laboratory in individual Petri dishes (9 cm diameter) filled with humidified moss until they completed tail resorption. Towel-dried wet mass of toadlets was then measured to the nearest 0.1 mg using a precision balance Mettler Toledo AG245 (Mettler Toledo, Geifensee, Switzerland), after which all toadlets were returned to their original population site.

Statistical analyses

We analyzed the data using a cell-means approach to a general linear model (Searle 1987) and a contrast analysis through planned comparisons (Dunson and Travis 1991) to test for the existence of interspecific competition. Since our analysis involved not only the mutual competitive effects under varying abiotic conditions, but also the effect of both biotic and abiotic factors on conspecific populations with different natural histories determined by a different ecological background, we further modified the model proposed by Dunson and Travis (1991). First, a one-way ANOVA was carried out on each species and response variable in which all the cells with various treatment combinations where a particular species is present in the design were used as class factors. Where overall ANOVA results were significant, a contrast analysis was carried out to test for a specific biological hypothesis. For our three *B. calamita* populations, seven contrasts are sufficient to test for main effect of salinity, intraspecific, and interspecific effects, and for all possible interactions among the main effects. An additional eighth contrast can be performed to test for the equivalence of intraspecific and interspecific density on the response variables, e.g., that *B. calamita* may be equally affected by the addition of 15 conspecifics or by the addition of 15 *B. bufo*. This test per se is not a contrast for the effects of interspecific competition because it already assumes that competitive effects are occurring (Dunson and Travis 1991), and it should not be done if neither of the previous contrasts for the effect of interspecific competition is significant. With respect to *B. bufo*, the calculations are further complicated because it is independently confronted with three different *B. calamita* populations, in which case there are 14 possible contrasts. An inherent difficulty of this design is the lack of orthogonality among some of the contrasts required, as well as the need for comparison-wise error rate ad-

justment due to the large amount of contrasts performed. Thus we minimized the number of comparisons by discarding those response variables for which the initial one-way ANOVA was nonsignificant, and by avoiding the calculation of equivalence contrasts when any of the interspecific contrasts was nonsignificant. We adjusted the significance level of the tests following the Dunn-Sidak algorithm (Sokal and Rohlf 1995). We used the mean of individual responses for each box as the effect of any given treatment on a particular experimental unit to avoid pseudoreplication of individual measures within boxes (Hurlbert 1984, Tejedo and Reques 1994).

Survival rates were angularly transformed by the arcsine-square-root transformation, and mass at and time to metamorphosis were ln-transformed in order to reduce skewness (Snedocor and Cochran 1989). Survival was introduced as a covariate in the analysis of both length of the larval period and mass at metamorphosis. The initial overall ANOVA yielded a significant block effect on both mass at and time to metamorphosis, and thus further analyses were corrected for block effect. All calculations were made with the PROC GLM from SAS statistical package (SAS Institute 1990).

RESULTS

Species level responses

Bufo calamita.—

1. *Water salinity*.—Being raised in either brackish or freshwater did not significantly affect survival rates of *B. calamita*. Salinity had a marginally significant effect on the duration of the larval period ($F_{1,90} = 7.65$, $P = 0.0069$, $\alpha' = 0.0064$), which was prolonged by 7.6 d under brackish conditions (Fig. 1). The mean mass of *B. calamita* at metamorphosis experienced an increase of ~28% ($F_{1,90} = 13.87$, $P < 0.001$) in saline water treatments with respect to freshwater. A correlation between mean date of and mean mass at metamorphosis for *B. calamita* yielded coefficients of $r = 0.67$ at low density, and $r = 0.03$ at high density, both in the absence of heterospecifics.

The interaction effect between intraspecific density and water salinity increased the duration of the larval period, and again was marginally significant ($F_{1,43} = 6.41$, $P = 0.015$). Both main factors exerted a similar effect in intensity and direction on date of metamorphosis at the higher stress levels (brackish water or crowding), extending the length of the larval period.

A similar effect was observed with the interaction of water salinity and the presence of *B. bufo*. *B. calamita* delayed metamorphosis 10.4 d when *B. bufo* was added, independently of water salinity.

2. *Intraspecific density*.—Intraspecific density did not contribute significantly to reduce survival of *B. calamita* larvae. In turn, the length of the larval period was increased and the mean mass at metamorphosis was reduced at high intraspecific densities (see Table

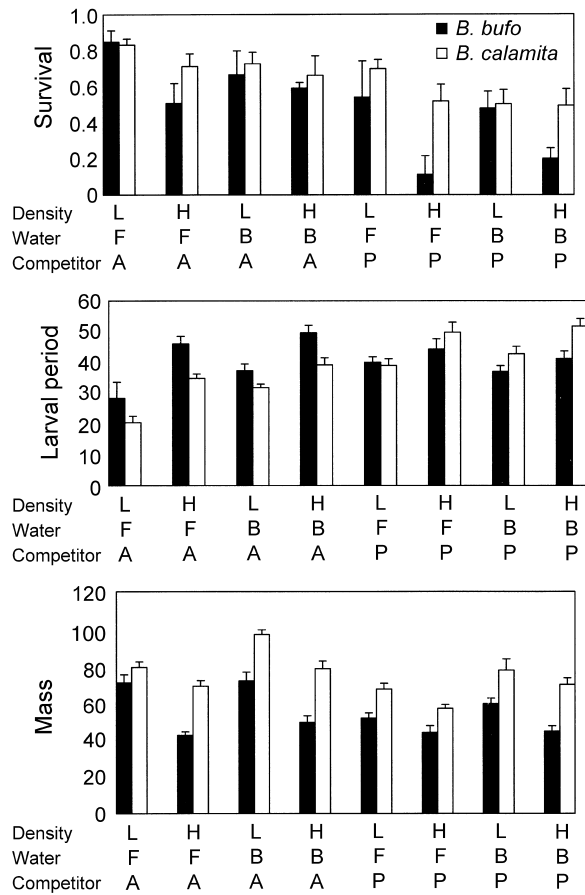


FIG. 1. Means of the response variables at the species level across all experimental treatments. L and H stand for low and high density treatments; F and B for fresh and brackish water treatments; and A and P for absence and presence of heterospecific tadpoles. Larval period is measured since the beginning of the experiment. Mass was measured at tail reabsorption (Gosner stage 46). Error bars indicate ± 1 SE.

2). The mean delay in the mean date at metamorphosis was of 12.1 d ($F = 17.09$, $P < 0.001$). Mean mass at metamorphosis was lowered from 87 to 72 mg, a 16.8% loss of body mass ($F_{1,45} = 14.9$, $P < 0.001$).

3. *Interspecific density*.—The presence of *B. bufo*

significantly lowered the survival probability of *B. calamita* ($F_{1,91} = 11.94$, $P < 0.001$; Fig. 1). Nevertheless, when the effect was tested separately for the two density levels, results were not significant at the adjusted significance level. The interaction between intraspecific density and presence of *B. bufo* was not significant, implying that the effect of the combined stresses was additive. Duration of the larval period was increased by 15 and 13 d, respectively, at low and high density ($F_{1,45} = 25.67$, $P < 0.001$; $F_{1,43} = 30.21$, $P < 0.001$, respectively) in the presence of *B. bufo*. Mass at metamorphosis decreased 18% at low and 13% at high density, though the latter test was not significant ($F_{1,45} = 14.13$, $P < 0.001$; $F_{1,43} = 7.13$, $P = 0.01$, respectively). As in the case of the two previous variables discussed, there was no significant interaction between intraspecific and interspecific densities, suggesting that the effect of these two sources of stress is additive.

Bufo bufo.—

1. *Water salinity*.—Water salinity did not significantly affect the survival and development of *B. bufo*. Even though the contrast analysis failed to show a direct significant effect of this factor on any of the three response variables, it behaved additively in relation to the other two sources of stress, contributing to reduce the survival rate of *B. bufo* at either intraspecific density.

2. *Larval density*.—High intraspecific density only significantly decreased survival when the heterospecific competitor was present ($F_{1,46} = 14.0$, $P < 0.001$). Interspecific competition only affected survival of *B. bufo* significantly at high density, reducing it from 0.55 to 0.25 ($F_{1,46} = 8.81$, $P < 0.005$). At low density only a slight tendency towards mortality increase could be detected. Duration of the larval period was considerably increased when intraspecific density was raised ($F_{1,37} = 23.17$, $P < 0.001$; see Table 2), metamorphosing a week later than their controls at low density. Date of metamorphosis remained unaffected by the addition of *B. calamita* at either density. Mass at metamorphosis was clearly diminished when intraspecific density was raised ($F_{1,34} = 21.1$, $P < 0.001$), falling

TABLE 2. Summary of effects of the main experimental factors on the response variables at the species level.

Response variable	<i>B. calamita</i>			<i>B. bufo</i>		
	Salinity	Intraspecific density	Presence of heterospecific	Salinity	Intraspecific density	Presence of heterospecific
Survival	-24.5
Larval period	55.9	69.8	90.7	...	62	40.8
Mass at metamorphosis	22.8	-12.7	-15.2	...	-40.9	-26.8

Notes: Only significant effects are shown. Values shown are percentages by which the response variable was affected as contrasted with the experimental controls in freshwater, low density, and absence of competitors. Positive values represent increases, while negative values show decreases.

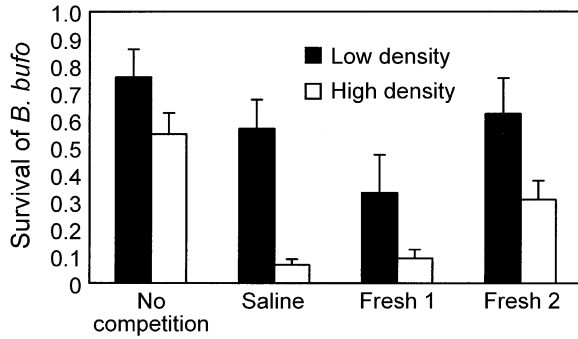


FIG. 2. *B. bufo* survival in the absence of *B. calamita*, and in competition with each of the three *B. calamita* populations. Error bars indicate +1 SE. Fresh 1 and Saline have the strongest impact, while the competitive effects of Fresh 2 are negligible. Fresh 1 reduced survival of *B. bufo* 47% at low population density and 82% at high density. The effect of Saline was only significant at high density where it represented a 92% decrease in survival of *B. bufo*.

from a mean 71.2 to 45.9 mg. Interspecific competition was only relevant in reducing the mass at metamorphosis when it occurred under low density conditions ($F_{1,34} = 17.12, P < 0.001$). Since mean mass at metamorphosis was sensitive to both intra- and interspecific density, we conducted a contrast for equivalence of competitors, which was found to be significant ($F_{1,34} = 9.44, P < 0.005$). Conspecifics caused a more severe reduction of body mass of *B. bufo* than heterospecific competitors (35.4 and 19.9%, respectively; Fig. 1).

Population level responses

Differential effect on *B. bufo*.—The three *B. calamita* populations affected differently the performance of *B. bufo*. Survival of *B. bufo* was significantly affected by Fresh 1 at low and high density ($F_{1,44} = 10.60, P < 0.003; F_{1,44} = 9.63, P = 0.003$, respectively), by Saline at high density ($F_{1,44} = 9.83, P = 0.003$), and was unaffected by Fresh 2 in all cases (see Fig. 2). The duration of the larval period of *B. bufo* remained unaffected in all three cases, while mass at metamorphosis was significantly reduced by Saline at low density ($F_{1,44}$

$= 15.74, P < 0.001$). This effect was marginally significant in the case of Fresh 1 and Fresh 2 ($F_{1,44} = 8.45, P = 0.006$ and $F_{1,44} = 8.86, P = 0.005$, respectively).

Saline.—Neither the increase in intraspecific density, the addition of an interspecific competitor, or the switch from freshwater to saline water seemed to affect the survival or the mass at metamorphosis of *B. calamita* larvae from the brackish pond of Saline (see Table 3). In turn, the larval period length was considerably dependent on the two biotic interactions. Increased intraspecific density significantly delayed duration of the larval period ($F_{1,17} = 10.72, P < 0.005$), as did competition with *B. bufo* (mean 17.2 and 19.6 d, respectively), though the effect was only significant at low density ($F_{1,17} = 17.26, P < 0.001$). The contrast of equivalence between competitors, opposing those treatments with 30 Saline larvae in it against those with 15 Saline larvae plus 15 *B. bufo* larvae was not significant.

Fresh 1.—The survival and mass at metamorphosis of the individuals belonging to the population of Fresh 1 was not affected by any of the factors considered (see Table 3). In turn, increasing the intraspecific density clearly elongated the larval period ($F_{1,20} = 16.46, P < 0.001$), postponing the date of metamorphosis a mean of 9.3 d. The addition of heterospecific competitors delayed the date at metamorphosis but the effect was only significant at low density ($F_{1,20} = 19.6, P < 0.001$). The mean date of metamorphosis was delayed a maximum of 20 d when combined with a high density of conspecifics. The contrast of equivalence between competitors was conducted and found not to be significant. As in the previous case, the saline environment fails to show a statistically significant effect on the duration of the larval period.

Fresh 2.—As in the case of the other two *B. calamita* populations, survival of the larvae from Fresh 2 was not significantly affected by any of the experimental treatments, and no contrast analysis was thus conducted on this variable. On the other hand, all three main factors significantly conditioned the duration of the larval period and all of them caused delays of varying length in the timing of metamorphosis. Salinity of the water

TABLE 3. Population-level summary of main effects on the three *B. calamita* populations.

Factor	Response variable	Populations		
		Saline	Fresh 1	Fresh 2
Salinity	survival
	larval period	+100.6
	mass at metamorphosis	+24.7
Density	survival
	larval period	+74.9	+41.5	+112.1
	mass at metamorphosis	-16.1
Presence of <i>B. bufo</i>	survival
	larval period	+106 (L)	+49.8 (L)	+139.4 (L), +63.7 (H)
	mass at metamorphosis	-10 (L), -20.6 (H)

Notes: Only significant effects are shown, and in the case of interspecific interaction, the intraspecific density level (L, low; H, high) at which the effect is significant is specified. The values indicate the changes in the mean of the response variable, expressed as a percentage of the experimental control mean under the different experimental treatments.

was responsible for a mean delay of 10.3 d ($F_{1,19} = 10.33$, $P < 0.005$). Differences in the experimental intraspecific densities accounted for a mean delay of 12.1 d ($F_{1,19} = 20.79$, $P < 0.001$). There was also a net interaction effect between the salinity of the water and the intraspecific density ($F_{1,19} = 8.33$, $P < 0.01$). The presence of heterospecific competitors also increased the duration of the larval phase, and it was so for both experimental densities. At the lowest one, *B. calamita* metamorphosed a mean of 17.4 d later than in absence of *B. bufo* ($F_{1,19} = 30.51$, $P < 0.001$) and also interacted significantly with water salinity, further increasing the larval period ($F_{1,19} = 10.06$, $P < 0.01$). At high density the delay reached 20.3 d ($F_{1,19} = 10.49$, $P < 0.005$). Mean mass at metamorphosis was clearly diminished when intraspecific density was increased ($F_{1,19} = 9.11$, $P < 0.01$), from a mean mass of 91 mg to 76 mg. The presence of *B. bufo* also reduced the body mass at metamorphosis, and the effect was significant at both densities. Interspecific competition at low densities caused a reduction of ~16% of the body mass, while at high densities the reduction was ~14%. The effect of water salinity over the mean mass at metamorphosis was statistically significant ($F_{1,19} = 14.64$, $P < 0.002$) and represented an increase in the mass of larvae under brackish conditions.

DISCUSSION

Species level

Water salinity caused a delayed metamorphosis in *B. calamita* (see Table 2). In turn, development of *B. bufo* was not significantly affected by osmotic stress. *B. bufo* is generally believed to be poorly resistant to water salinity, partly because it is usually absent from saline environments. Thus it is surprising to find that salinity as main effect did not affect either of the response variables of *B. bufo* in this study. The fact that *B. bufo* tadpoles were exposed to the brackish conditions at a later developmental stage than natterjacks may have reduced the negative effects of salinity. However, according to our data, the absence of *B. bufo* from brackish environments cannot be explained solely in terms of detrimental effects of salinity on the larvae. The impact of salinity on the embryos seems to be more acute in *B. bufo* when compared with higher tolerance in natterjack embryos (I. Gómez-Mestre and M. Tejedo, *unpublished data*) although other explanations such as breeding site selection may explain the pattern. On the other hand, the present experimental design may have low statistical power to detect the effect of water salinity by itself and we could only perceive it as a synergistic effect with other factors. The significant increase in mass at metamorphosis caused by water salinity in *B. calamita* is likely to be an indirect consequence of prolonging the larval period while food availability remains constant. Contrary to our predictions, the abiotic factor studied did not qualitatively

alter the outcome of the competitive interaction, but merely intensified it, as has been the case in other systems studied (Pehek 1995).

Interspecific competition did occur between *B. calamita* and *B. bufo* in our experiment. The mutual effects were asymmetric, but led to a decrease in survival, a retardation of the larval period, and a reduced mean mass of the emerging toadlets for both species, reducing their fitness. Similar consequences of interspecific competition have also been reported for other competing anuran species (Wilbur 1982, Morin 1983a, b, Werner 1986, Warner et al. 1993, Baker and Beebee 1997, Bardsley and Beebee 1998a).

According to our predictions, *B. bufo* had a marked competitive effect (*sensu* Goldberg and Fleetwood 1987) over *B. calamita* (increased mortality, delayed metamorphosis, and decreased mass at metamorphosis), and also exhibited a stronger tolerance to competition than *B. calamita*. The competitive effect was asymmetric since survival of *B. bufo* was not significantly affected by *B. calamita*. *B. bufo* survival was only diminished when competition occurred at high density as an interaction with the increase in intraspecific competition. Differences in response to competition are also remarkable because *B. calamita* experienced an increase in the duration of the larval period more than twice the one experienced by *B. bufo* when they were set to compete (see Table 2). The relatively minor decrease in mass at metamorphosis showed by *B. calamita* with respect to *B. bufo* may be at least partially attributable to relaxed intraspecific density caused by mortality.

Competitive superiority among species is thought to be size dependent to a large extent (Connell 1983, Thompson 1988, Werner 1994). In the case of anuran larval guilds of phylogenetically close taxa, interspecific relative-size relationships are conditioned primarily by the time span between the breeding of the different species. Early breeders attain large sizes by the time other species lay their clutches and that difference may be translated into competitive superiority throughout the entire larval period (Alford and Wilbur 1985, Faragher and Jaeger 1998). In southern Spain *B. bufo* usually breeds earlier than *B. calamita*, when the autumn–winter rainfall is abundant enough, but it is also common that their breeding seasons greatly overlap when rainfall is more scarce or starts later. In our experiment, *B. bufo* tadpoles were two weeks older than *B. calamita*. Thus, *B. calamita* larvae could be expected a priori to be an inferior competitor in all interactions with *B. bufo*, a situation described for several UK populations (Bardsley and Beebee 1998a, b). However, considering a resource basis of competition, if one species of a competing pair has a clear impact on the resource which is the object of the competition, it is conceivable to expect a certain degree of reciprocal effect, however slight it might be (Woodward 1982, Connell 1983, Alford and Wilbur 1985). Although the

design employed in this study does not allow us to distinguish between resource and interference competition, it was observed that *B. bufo* also experienced negative effects resulting from the interaction with *B. calamita*. However, in the case of *B. bufo*, the detrimental effects of intraspecific competition exceeded those caused by interspecific competition, while the reverse was true for *B. calamita* (see Table 2), as was predicted. Superiority against heterospecific competitors suggests a strong competitive ability and has been shown to correlate with intense intraspecific competition (Wilbur 1982, Semlitsch and Walls 1993). Thus, if *B. bufo* is forced to spawn in more ephemeral ponds, the concomitant increase in intraspecific density may compromise its competitive superiority over *B. calamita*, a species customary to highly temporary ponds. Pond duration can alter the outcome of competition by reducing survival, even though the intensity of the interaction in terms of relative growth rate remains constant (Petranka and Sih 1986). As temporary ponds dry out, density increases, which may affect *B. bufo* more strongly than *B. calamita*, and ultimately modify the relative ratio of metamorphic success. A high density of conspecifics renders *B. bufo* more vulnerable to interspecific competition and to abiotic stress conditions. Similar phenomena have been demonstrated for other species, e.g., *Bufo terrestris* in relation to water temperature and dissolved oxygen (Travis and Trexler 1986), and the competing pair *Hyla gratiosa*–*Hyla femoralis* in relation to pH (Warner et al. 1991, 1993). Overall, interactions at different levels, synergism, and antagonism among biotic and abiotic factors (Newman 1998) may be the rule rather than the exception in generating diversification in ecological processes.

Population level

Our results confirmed the prediction that geographic variation exists among populations of *B. calamita* in their response to the biotic and abiotic factors under study, as a reflection of the local selective pressures. Water salinity had no effect on either Saline or Fresh 1 populations. *B. calamita* from the Saline population seem to have undergone processes of local adaptation to brackish water environments (I. Gómez-Mestre and M. Tejedo, unpublished data), and thus it was expected to be less sensitive to water salinity than those populations coming from freshwater environments. Lack of effect of osmotic stress on Fresh 1 could not be predicted in terms of its natural environmental conditions. Osmotic stress resistance could have evolved in Fresh 1 as selective response to particular environmental conditions, but given the relative geographic proximity of Fresh 1 and Saline, gene flow estimates would be required to exclude the possibility of immigration of salinity-resistant genotypes. In contrast, Fresh 2 markedly delayed metamorphosis when reared under osmotic stress. Delaying metamorphosis in a brackish temporary pond forces tadpoles to face the risk of des-

iccation of the pond and also the steady increase in salinity as water evaporates, greatly reducing survival probability.

Because different populations of the same species experience large variations in life history and morphological traits, interactions between species are not immutable (Thompson 1994, 1999, Fauth 1998), but they vary in both space and time as species coevolve and adapt to the regionally predominant environmental conditions. Interpopulational variation among amphibian species in competitive and predator–prey interactions is not uncommon (Fauth 1998). The three *B. calamita* populations examined here also differed in their response to competition with *B. bufo*. As indicated in Table 3, Fresh 1, naturally syntopic of *B. bufo*, experienced by far the least increase in larval period. Saline tadpoles showed a less acute retardation of larval period in the presence of heterospecifics than Fresh 2, which in addition decreased its mean mass at metamorphosis when *B. bufo* was present. Approaching the issue from another perspective, we can score the competitive effect of each of the three *B. calamita* populations on *B. bufo*. Only Fresh 1 and Saline reduced *B. bufo* survival, and only Fresh 1 gave this result under low density conditions (see Fig. 2). Furthermore, mean mass at metamorphosis of *B. bufo* was also more reduced by Fresh 1 than by the other two populations. Competitive effect and competitive response are not necessarily positively correlated (Goldberg and Landa 1991), but in the case of these *B. calamita* populations, the greater the effect on the heterospecific, the greater the resistance to interspecific competition.

Saline and Fresh 1 are naturally exposed to more extreme environments, suggesting that more resistant phenotypes have been selected in these populations. Saline breeds in a brackish pond with low predation and competitive intensities (*B. bufo* is virtually absent), but is exposed to a high degree of osmotic stress while Fresh 1 breeds in a mountain stream, with a shorter benign period for larval development and crowded with competitors, particularly *B. bufo*. Fresh 2, in turn, breeds in temporary freshwater ponds where *B. bufo* is absent. These differences among populations result from diverse local-differentiation processes under the specific environments where they have been evolving.

Our data support the notion that populations within any given species are not geographically homogeneous subsets sharing the same pattern and trait distribution (Thompson 1988). Rather, populations seem to vary in structure and dynamics, in their degree of specialization to the different environmental conditions found, both biotic and abiotic, and thus the type, intensity, and outcome of their interactions with other organisms are likely to be variable and environment specific. Most biotic interactions show a geographic mosaic dynamics at both the ecological (Tilman and Kareiva 1997, Bascompte and Solé 1998) and evolutionary time scales (Herrera 1988), resulting from the combination of phy-

logenetic constraints, the biogeography of the interacting species, and stochastic and deterministic genetic phenomena acting on them (Thompson 1994, 1999, Gomulkiewicz et al. 2000). In that sense, we conclude that even though competitive superiority of *B. bufo* over *B. calamita* is generally true, the outcome of competition between these two species is to a great extent dependent on the particular surrounding abiotic conditions and is strongly mediated by density-dependent effects. Furthermore, the modulation of these interactions varies geographically among populations both because of the different environments experienced and because of the particular natural histories of the populations involved.

Future research should broaden the spectrum of populations screened to generalize the observed pattern in competitive ability and to estimate their underlying genetic variance within and among populations, while gaining insight on the proximate mechanisms mediating the competitive interaction.

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