



**Bimodal breeding phenology in the parsley frog  
*Pelodytes punctatus* as a bet-hedging strategy in an  
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20 ABSTRACT

21 When environmental conditions are unpredictable, expressing alternative phenotypes spreads  
22 the risk of failure, a mixed strategy called bet-hedging. In the southern part of its range, the  
23 Parsley frog *Pelodytes punctatus* breeds from autumn to spring. Our aim was to study the  
24 breeding phenology and reproductive success associated with the use of those two seasonal  
25 niches to understand how this breeding strategy can be maintained. Field surveys revealed  
26 that breeding phenology was typically bimodal with a higher breeding effort in autumn. More  
27 importantly, in spring, the survival rate of offspring was severely reduced by the presence of  
28 autumn tadpoles, indicating a clear priority effect. However, the autumn cohort often failed to  
29 survive over winter, in which case spring cohorts were often successful. Based on those  
30 results, we constructed a model in which females can allocate a variable portion of eggs to  
31 each season and added a priority effect. We conclude that the existence of the two breeding  
32 seasons may indeed constitute a bet-hedging strategy.

33

## INTRODUCTION

Breeding phenology is one of the key components of adaptation to temporally variable environments. Temporal dynamics of both the biotic and abiotic environment impose selective constraints on parental development and physiological state (to be able to reproduce) as well as offspring survival (at the various developmental stages until they reach sexual maturity and start to reproduce) (Rand 1973). There is a vast amount of literature on intraspecific variation of breeding patterns in relation to environmental conditions, in particular latitude, altitude and climate. In the context of current climate change, many species in temperate regions have advanced their breeding time (e.g. Brown et al., 2016; Frederiksen et al., 2004; Møller, 2008), as a result of microevolutionary changes and/or of phenotypic plasticity (Charmantier & Gienapp 2014). Most of these studies concern species with a single reproductive peak in the year, which has to match as precisely as possible a seasonal peak of resource availability in order to maximise reproductive success (e.g. caterpillar availability for tits). The exact date of the resource peak may vary from year to year and species usually rely on cues to anticipate it and plastically delay or advance the onset of reproduction every year. However, in some cases reproductive success depends on even irregular and/or unpredictable conditions. In such situations, species face the risk of complete reproductive failure at any given breeding attempt, a regime that favors expressing alternative phenotypes to spread the risk, a strategy known as bet-hedging (Cohen 1970; Slatkin 1974; Philippi & Seger 1989; Leimar 2005; Venable 2007).

Theory predicts that in stochastic environments, selection favors life history traits that reduce temporal fitness variation even if they result in lowered arithmetic mean fitness (Philippi & Seger 1989). In temporally variable environments, long-term fitness of a genotype is measured by the geometric mean of the fitness contribution over successive years at the individual level or over successive generations for a particular genotypic (Lewontin & Cohen

1969; Olofsson *et al.* 2009; Simons 2011; Yasui & Yoshimura 2018). This geometric mean fitness is highly impacted by low values; hence, traits with lower fitness variation may have higher long-term fitness. In principle, lower variation in fitness can be achieved either using the same low-risk strategy (conservative bet-hedging), or displaying several strategies, either at once or over several instances (diversified bet-hedging). While there is an abundant theoretical literature on bet-hedging, empirical studies have provided limited evidence so far (Simons 2011), and the most comprehensive examples concern the timing of germination/diapause and the fraction of dormancy seeds/diapausing eggs (Venable 2007; Gremer *et al.* 2016; García-Roger *et al.* 2017; Wang & Rogers 2018). Some studies even report experimental evolution of bet-hedging traits in response to unpredictable environment (Beaumont *et al.* 2009; Maxwell & Magwene 2017; Tarazona *et al.* 2017).

One of the best examples of stochastic, unpredictable environments are temporary ponds, alternating between inundation and drought where each breeding event is a bet as habitat desiccation can occur before the end of the breeding cycle. Several examples of bet-hedging occur in temporary ponds. Fairy shrimps (Anostraca) produce drought-resistant eggs showing asynchronous hatching at different hydroperiods (Saiah & Perrin 1990; Simovich & Hathaway 1997; Wang & Rogers 2018). Similarly, rotifers produce diapausing eggs to overpass unfavorable planktonic growing season and only a fraction of those eggs hatch when conditions are suitable (García-Roger *et al.* 2017; Tarazona *et al.* 2017).

For amphibian species breeding in temporary ponds, drought can cause 100% mortality of eggs or larvae, resulting in complete failure of one breeding event. One way to reduce the risk of losing a breeding opportunity entirely is to spread this risk at a spatial scale, partitioning brood into various pools as done by the Neotropical poison frog *Allobates femoralis* (Erich *et al.* 2015). Another bet-hedging strategy could be to split the breeding effort at a temporal scale and exploit all suitable temporal windows.

From an ecological point of view, such temporal niche partitioning is expected to reduce inter and intra-specific competition as well as resource depletion (Carothers & Jakšić 1984). For example, species may share the same habitat but have opposite activity patterns (nocturnal versus diurnal species), as is the case in Neotropical felid community or in grassland ants (Albrecht & Gotelli 2001; Di Bitetti *et al.* 2010). The same type of temporal segregation at a daily scale is also observed within species: brown trout *Salmo trutta* reduces competition for a limiting resource by sequential use of foraging areas (Alanärä *et al.* 2001). Voltinism in insect is another well-studied example of temporal partitioning at the annual scale which is an adaptation to predictable seasonal cycles (Kivelä *et al.* 2013; Zeuss *et al.* 2017; Forrest *et al.* 2019)

However, if successive seasonal cohorts overlap, fitness gains may be asymmetric, because progeny produced by late breeding may suffer from competition or even predation from earlier cohorts (Morin 1987; Ryan & Plague 2004; Eitam *et al.* 2005). Those priority effects are often difficult to disentangle from seasonal effects due to environmental differences experienced by the temporal cohorts (Morin *et al.* 1990). If priority effects are strong, late breeders may select breeding sites in order to limit the competition by conspecifics (Halloy & Fiaño 2000; Halloy 2006; Sadeh *et al.* 2009) and this may restrict late breeders to poorer sites (Crump 1991).

In amphibians, this temporal partitioning of breeding activity is thought to regulate community dynamics through interspecific competition (Lawler & Morin 1993; Gottsberger & Gruber 2004; Richter-Boix *et al.* 2006b, 2007c). Similarly, community composition may depend on species arrival and priority effects whereby species arriving earlier monopolize available resources and gain a competitive advantage over late species (Morin *et al.* 1990; Blaustein & Margalit 1996; Urban & De Meester 2009). In Mediterranean regions, climatic conditions are characterized by dry, hot summers and mild winters, with the maximum

rainfalls in autumn and spring. This leads to large breeding asynchrony observed between and within amphibian species (Diaz-Paniagua 1988; Jakob, Poizat et al. 2003; Richter-Boix, Llorente et al. 2006; Vignoli, Bologna et al. 2007): whereas most species typically breed in spring, some species breed earlier at the end of winter, and some even breed in autumn in addition to spring.

The Parsley Frog *Pelodytes punctatus* is a small sized Anuran distributed in Spain and most of France (locally reaching neighboring countries). It has a broad ecological niche but has poor competitive abilities and is sensible to fish predation (Morand & Pierre 1995; Crochet et al. 2004; Richter-Boix et al. 2007b); it thus prefers seasonally flooded habitats to than large permanent water bodies (Guyétant et al. 1999; Salvidio et al. 2004; Richter-Boix et al. 2007a). In Spain the parsley frog shows a bimodal breeding pattern with higher reproductive effort in spring than in autumn (Guyétant et al. 1999; Richter-Boix et al. 2006b). In France, in addition to spring breeding, autumnal breeding is also observed in Mediterranean regions and areas with mild oceanic climate (Guyétant et al. 1999; Jakob et al. 2003; Richter-Boix et al. 2006a; Cayuela et al. 2012) but the importance of autumn versus spring reproduction has not been quantified. In the rest of the range and in altitude, only spring breeding occurs.

In the Mediterranean areas of southern France, the parsley frog uses temporary ponds that refill in September and October but may dry during autumn or later in late spring. Adults have thus to deal with very unpredictable environmental conditions for their future offspring. In addition to this unpredictable risk, tadpole hatched in autumn or spring are exposed to very different environmental conditions, the most obvious being that the autumn tadpoles overwinter while the spring ones do not. This should result in drastically different developmental trajectories but also in different offspring survivals. Both seasonal cohorts may also interact, leading to a potential competitive advantage to the earlier cohort over the later, i.e. a priority effect. The relative success of each breeding period and the outcome of the

interaction between cohorts are key parameters to understand the persistence of this two-peaks breeding strategy. In fact, several pieces of information are still lacking in order to understand the evolutionary basis of this seasonally variable breeding strategy. Do we have a single protracted breeding season or a really bimodal reproduction generated by the coexistence of alternative breeding timing (autumn or spring)? If so, what is the relative importance of autumn versus spring reproduction? What is the survival of offspring produced at the two breeding periods and how is it affected by the presence of conspecifics? Once this basic knowledge is obtained, it can be fed into theoretical models for the evolution of mixed breeding strategies.

In this paper, we characterize the breeding phenology (temporal dynamic, relative proportion of each breeding period) of parsley frog in a French Mediterranean area based on results from a 3-year field survey. We monitored the survival of offspring produced in each season to estimate the success of this breeding strategy. We also investigated the factors influencing breeding and tadpole survival, in particular whether there is a priority effects between seasonal cohorts. Finally, using an analytical model adapted from Cohen (Cohen 1966) we tested whether the coexistence of the two breeding periods can be interpreted as a bet-hedging strategy.

## MATERIAL AND METHODS

### **Field survey**

The field study was carried out from September 2007 to August 2010 in 19 ponds situated around Montpellier, southern France (Annex 1). Those ponds are man-made environments, often dug out to provide drinking water for livestock (sheep and cows) or for game. The ponds surveyed included temporary and permanent sites. We define here the autumn breeding season as the period spanning from September to December and the spring breeding season



from January to April. We surveyed each pond twice each month. During each visit, we recorded the depth of the pond.

## **Sampling methods**

At every visit (mostly diurnal), we looked for newly deposited egg masses throughout the entire water body and classified the egg masses as small, medium and large, corresponding to an average of 75, 150 and 250 eggs per mass, respectively (Salvador & Paris 2001) and personal observation). The parsley frog's embryonic period ranges from 5 days at 15°C to 15 days at 10°C (Toxopeus *et al.* 1993). Moreover, embryos stay attached to the jelly for several additional days (Guyétant *et al.* 1999). Thus, with an interval of 15 days between two successive visits, we may have missed a few masses but we have avoided double-counting masses since 15-day old masses can readily be distinguished from new ones based on the developmental stages of the embryos. In only 2% of the larval cohort produced, small larvae were observed in ponds where we did not notice the presence of egg masses before. Note that the probability of detection of an egg mass, even if not perfect, was similar in autumn and in spring.

We estimated the number of amphibian larvae and invertebrates present in the ponds using 5 to 10 dipnet sweeps (depending on the pond size). The anuran community of the area consists of 7 species: *Pelodytes punctatus*, *Pelobates cultripes*, *Alytes obstetricans*, *Bufo bufo*, *Epidalea calamita*, *Hyla meridionalis*, and *Pelophylax sp.*, (*P. ridibundus* and/or *P. perezi* & *P. kl. grafi*, depending on the sites). Potential predators of tadpoles are urodeles and aquatic invertebrates. Two urodele species (*Lissotriton helveticus* and *Triturus marmoratus*) were recorded in the ponds but due to the rare occurrence of *Triturus marmoratus*, only *Lissotriton helveticus* was included in subsequent analyses (as adults as well as larvae).

We also surveyed dragonfly larvae (Anisoptera) and backswimmers (Heteroptera, Notonectidae) that are potential predators of tadpoles (Richter-Boix *et al.* 2007a) except during the first year. Diving beetles (Coleoptera, Dytiscidae) are also known to prey on tadpoles but were very rare in the studied ponds and thus not considered for this study.

We divided the total counts for each amphibian larvae and invertebrate predators captured in each pond by the number of dipnet sweeps taken in each pond. This procedure yielded a crude proxy for density on the basis of catch per unit effort and could therefore be compared across localities.

### **Reproductive effort and offspring survival**

Reproductive strategy of parsley frog was described by two measures: the probability of spawning and the breeding effort. Spawning probability indicates if any new egg mass was observed during a visit. Breeding effort measures the quantity of eggs produced when there was at least one egg mass. We normalized the number of egg masses by their size (e.g. a small egg mass equals  $\frac{1}{2}$  medium egg mass).

For each breeding event, we estimated the hatching rate as the ratio of the number of small tadpoles (Gosner stage 26, free swimming tadpole) to the number of eggs spawned. Similarly, we quantified the survival from egg to metamorph as the ratio of the number of metamorphs (Gosner stage 42-43) to the number of eggs spawned. When hatching was successful (i.e. in about one third of the breeding event), we could calculate the survival during larval stage as the product of the two former ratios.

The number of tadpoles in a pond was estimated using the mean number of tadpoles caught per dipnet sweep scaled to a sampling surface of 1 m<sup>2</sup> (we estimated that one dipnet sweep sampled a surface of 0.5 m<sup>2</sup>, taking the dipnet size and the length of the haul into account) and then multiplied by the surface of the pond. This should not be taken as an attempt to estimate

precisely the number of tadpoles present in a pond at a given time but as an index of abundance that can be compared between ponds and between breeding events. It was sometimes impossible to follow the larval development and metamorphosis of offspring from a particular breeding event. Indeed, parsley frogs may breed three to four times during each seasonal breeding event. In these cases, the successive sub-cohorts produced are undistinguishable after a few weeks, and we summed the eggs counted in two or three successive visits to evaluate survival from a combination of breeding events within a given season (and within a site). Survival measures should be viewed as an index to assess the differences of reproductive success between seasons as there is no reason to expect any seasonal bias in this index.

### **Explanatory variables**

Explanatory variables for the breeding probability and breeding effort are the depth of the pond as well as the presence of conspecific and inter-specific competitors (larvae of anuran species) and predators (invertebrates and adult newts) in the pond. The two last categories of variables were also applied to explain the success (offspring survival) of breeding events. We summed the density of competitors and similarly the density of predators despite the differences in competitive performance and predation pressure of the various species toward parsley frog tadpoles.

To assess the potential impact of predation and competition on survival rates, we evaluated the mean density of predators and competitors encountered by parsley frog tadpoles during their larval development. More precisely, data from literature indicates that only small tadpoles (<12 mm snout-vent length) have lower survival due to predation by aquatic invertebrates (Tejedo 1993). Above this size, the predators will only injure them or even fail to catch them. Larvae laid in autumn reached this limit size in about 3 months, whereas only

1.5 month is necessary for larvae laid in spring (personal observation). Thus, we used the mean density of predators and competitors over a period of 3 months after spawning date for autumn tadpoles and 1.5 months for spring tadpoles.

### **Statistical analyses**

All statistical analyses were performed on R 3.4.1 (R Core Team 2018). To assess if pond characteristics differed between seasons, we apply a linear model for the depth of the pond and generalized linear models with a quasi-poisson family for all other variables to account for overdispersion. Breeding probability and breeding effort were analysed using a generalized mixed model with site as random effect, with a binomial family or a negative binomial family (to account for overdispersion), respectively. The survival rates were often zero hence we decided to analyse them as binary variables using a generalized mixed model with site as random effect and a binomial family. Those variables, called breeding success and hatching success, are the probability of producing at least one metamorph or one hatchling. The significance of fixed effects were tested using  $\chi^2$  tests to compare nested models (Zuur *et al.* 2009).

### **Bet-hedging model**

Finally, we wondered if the coexistence of two breeding periods could result from a bet-hedging strategy, with the optimal strategy being to split the breeding effort between the two favourable seasons to spread the risk of complete failure (Seger, J. and Brockman 1987). The following model is derived from Saiah and Perrin (Saiah & Perrin 1990) on the hatching probability of fairy shrimp seasonal cohorts. This model was primarily inspired by Cohen (Cohen 1966), reviewed by (Seger, J. and Brockman 1987) on the optimal reproduction

strategy of an annual plant whose seeds can either germinate or remain dormant. In our case, there are two strategies: autumn breeding with initial success (i.e. the ability of offspring to persist until spring) depending on the environmental conditions and spring breeding with success depending mainly on the presence of autumn tadpoles, hence on the initial success of autumn breeding (as suggested by the results on success of autumn and spring breeding events, see below).

Let  $c$  be the proportion of eggs laid in autumn (thus  $1-c$  in spring) – we assume, in agreement with our data (see results), that  $c$  represents a fixed strategy, i.e. the frogs cannot predict failure in advance to avoid laying in autumn, nor can they avoid laying eggs in spring when an autumn cohort is present. As mentioned above, the autumn cohort is assumed to succeed or fail, at random, with probability  $q$  and  $1-q$  respectively. When it succeeds, a fraction  $s_1$  of the offspring survive to reproductive age. The spring cohort completely fails whenever the autumn cohort has survived in a pond (a reasonable simplification based on our survival rates estimates, see below ), otherwise a proportion  $s_2$  of spring tadpoles survive. Overall the mean number of individuals produced per female is  $c s_1$  when the autumn cohort doesn't fail and  $(1 - c) s_2$  when it does.

If each frog reproduced only during one year, the optimal strategy would maximize the geometric mean of the annual reproductive outcome (Dempster 1955) which is

$$W = (c s_1)^q ((1 - c) s_2)^{1-q}$$

Or, equivalently

$$\text{Log}(W) = q \text{Log}(c s_1) + (1 - q) \text{Log}((1 - c) s_2)$$

However, reproductive life lasts more than one year in frogs (say,  $n$  years), which in itself is a way to spread the risk of failure among successive cohorts of offspring – an uncertainty remains however, for each frog, on how many ( $k$ ) of the  $n$  breeding years will not allow the

autumn cohort to survive. For each individual,  $k$  is distributed binomially with probability  $1-q$  so that

$$\text{Log}(W) = \sum_{k=0}^n \binom{n}{k} (1-q)^k q^{n-k} \text{Log}(c s_1 (n-k) + (1-c) k s_2)$$

where  $\binom{n}{k}$  represents the number of possible repartitions of the  $k$  years with autumn failure among the total number of breeding years  $n$ .

The selection gradient is

$$G[c] = \sum_{k=0}^n \binom{n}{k} (1-q)^k q^{n-k} \frac{(s_1 + s_2) k - n s_1}{c(k(s_1 + s_2) - n s_1) - k s_2}$$

We traced the fitness curves and the selection gradients using Mathematica (Wolfram Research Inc. 2018) based on the following parameter combinations. We set survival probabilities based on our estimates of survival from egg to metamorphose:  $s_1 = 0.047$  (estimated among breeding events producing offspring that survived until spring) and  $s_2 = 0.038$  (in the absence of autumn tadpoles). We assumed that survival and fecundity were equal for both seasonal cohorts for the rest of the life cycle. We set the number of reproductive years  $n = 3$  to 5, according to a study of age structure of a breeding population in Spain (Esteban *et al.* 2004). Note that this model applies at the individual level (as developed above) as well as at the genotype level.

## RESULTS

### Characteristics of temporal niches

Pond depth was not significantly different between the autumn (here from September to December) and spring (here from January to April) breeding seasons (Table 1). The densities of amphibian larvae (other than parsley frog) were not significantly different. In autumn, extreme densities of *Epidalea calamita* tadpoles were recorded in some small ponds whereas

the well-known spring breeding-species (*Hyla meridionalis*, *Pelophylax* sp., *Triturus marmoratus*, *Lissotriton helveticus*) reproduce later than the parsley frog, hence their larvae are only present from April onwards. The density of potential invertebrate predators was higher in autumn than in spring ( $\chi^2_1 = 37.17$ , p-value = 0.005) with the lowest density being from December to March. On the contrary, the number of adult newts (potential predators of parsley frog tadpoles) was higher in spring than in autumn ( $\chi^2_1 = 369.36$ , p-value =  $2.2 \times 10^{-16}$ ).

### **Breeding phenology**

We registered 184 breeding events, 79 in autumn and 105 in spring. The breeding effort showed a bimodal pattern with a peak in October and another in February (Figure 1). Note that in two sites, one breeding event was recorded in May.

The spawning probability (finding at least one egg mass when visiting a pond) was not significantly different between the two seasons (0.18 (0.02), mean (S.E) per visit in autumn and 0.22 (0.02) per visit in spring, (over 429 and 470 visits, respectively)  $\chi^2_1 = 2.31$ , p-value = 0.128, see also Annex 2). This spawning probability was not affected by the presence of anurans from other species (larvae), nor by the presence of predators (invertebrates or adult newts). It was positively related to the depth of the pond ( $\chi^2_1 = 20.40$ , p-value =  $6.3 \times 10^{-6}$ ). The spring spawning probability was not affected by the presence of autumn tadpoles ( $\chi^2_1 = 0.03$ , p-value = 0.875).

The breeding effort was higher in autumn than in spring (23.0 (4.0) egg masses per breeding event in autumn and 13.7 (2.4) in spring;  $\chi^2_1 = 9.25$ , p-value = 0.002, Fig. 2 and Annex 3). As a result, autumn breeding contributed slightly more than spring breeding to the production of egg masses (57% versus 42.9%).

### **Breeding success**

329 Hatching success (i.e. the percentage of breeding events producing at least one larvae) was  
330 higher in autumn than in spring (68.4% and 43.8% respectively,  $\chi^2_1=11.12$ , p-value= 0.001).  
331 Drought (pond totally dried up) caused the total failure of 7 breeding events (9% of the  
332 breeding events) in autumn and of 5 breeding events in spring (4.8%) over the 3 year-survey  
333 and the 19 sites. Drought caused mortality of offspring at different developmental stages  
334 (mostly eggs for autumn cohort and tadpoles for spring cohort). As a consequence, breeding  
335 success (i.e. the percentage of breeding events producing at least one metamorph) was not  
336 significantly different between the two seasons (34.2% in autumn and 29.8% in spring  $\chi^2_1=$   
337 0.39, p= 0.531).

338 Neither breeding success nor hatching success were explained by interspecific competition  
339 (the density of other amphibian larvae) or by predation (density of potential invertebrate  
340 predators or number of adult newts).

341 Survival rates are represented in Figure 3 and Annex 4. The survival rates from egg to  
342 metamorph were similarly low (autumn: 2.24 % (0.61) and spring: 1.97 % (0.73)), resulting in  
343 a higher contribution (74.6%) of autumn breeding in the overall production of metamorphs  
344 per site and per year (due to the higher breeding effort in autumn, see above).

345 The autumn cohort persisted until spring in 34/79 breeding events (43%, corresponding to the  
346 rate of initial success,  $q$ , see bet-hedging model). In those cases, tadpoles laid in spring  
347 coexisted in their pond with tadpoles from the autumn cohort. From the point of view of  
348 spring breeders, in 28/57 cases, they found autumn tadpoles in the pond. The presence of an  
349 autumn cohort of parsley frog tadpoles significantly reduced the success of spring breeding  
350 event to 18.4% in presence of autumn tadpoles, versus 40.0% in absence of autumn tadpoles,  
351  $\chi^2_1=10.60$ , p-value= 0.005). This reduction effect was not significant for the hatching success  
352 (32.6% in presence of autumn tadpoles, versus 53.6% in absence of autumn tadpoles  
353  $\chi^2_1=4.63$ , p-value=0.099). Accordingly, all survival rates were reduced in the presence of



autumn tadpoles and this effect was most pronounced for the survival from egg to metamorphs (3.77 (1.4) versus 0.16 (0.08) in absence versus presence of autumn tadpoles, Figure 4 and Annex 5).

Finally, the figure 5 summarize the breeding strategy of parsley frog showing the presence of egg masses, tadpoles and the outcome of the breeding event (production of metamorphs) in each studied site, over the three years of survey. It illustrated the quasi-exclusion between the two cohorts.

### **Maintenance of spring breeding**

Selection gradients based on our bet-hedging model predict that a mixed strategy is maintained when the rate of initial success of the autumn cohort ( $q$ ) is between 0.2 and 0.8 for a number of reproductive years  $n = 3$ . In this condition, a pure autumn strategy is predicted above 0.8, and a pure spring strategy below 0.2. (Figure 4). The maintenance of this strategy is less probable if the number of reproductive years increases ( $n = 5$  years of breeding), with a reduced range of  $q$  leading to a stable mixed strategy.

## **DISCUSSION**

### **Cost and benefits of a bimodal breeding phenology**

We used field surveys to describe the breeding phenology of the parsley frog in the French Mediterranean region but also to quantify the relative contribution and success of each seasonal reproduction (autumn and spring reproduction). This quantification, rarely achieved in the wild (but see (Licht 1974; Banks & Beebee 1988; Gascon 1992; Wheeler *et al.* 2015), is essential to understand the evolution of this bimodal breeding strategy. We confirmed the existence of two distinct seasonal peaks in breeding activity, probably mediated by cold temperature in December and January as adult parsley frogs tend to breed in mild and rainy

periods as was previously observed (Toxopeus *et al.* 1993; Guyétant *et al.* 1999; Jakob *et al.* 2003). However breeding episodes occurred even in the absence of rainfall as long as ponds were filled with water (personal observations and Richter-Boix *et al.* 2006b)).

The breeding effort in our population was higher in autumn than in spring. This is in apparent contradiction with Richter-Boix *et al.* (2006b) who found that spring breeding effort was four-fold higher than autumn breeding effort in the northeast of the Iberian Peninsula. While we don't have a definitive explanation for this difference, we suggest it could be related to higher competition among anuran larvae in autumn in north-east Spain compared to France. In our study area in southern France, larvae of *Pelodytes punctatus* are typically the only anuran larvae found after the summer drought in the ponds in autumn. In contrast, four other species of Anura (*Hyla meridionalis*, *Epidalea calamita*, *Alytes obstetricans* and *Pelophylax perezi*) have tadpoles in autumn and three of them (i.e. all except *E. calamita*) can have overwintering tadpoles in Spanish ponds (Richter-Boix *et al.* 2006b). These authors also showed that *Pelodytes punctatus* tadpoles suffer from interaction with *Hyla meridionalis* (Richter-Boix *et al.* 2007b). It is thus possible that increased competition for *Pelodytes punctatus* larvae in autumn and winter compared to our study area make the autumn niche less favourable in northeastern Spain compared to southern France and reduce parsley frog investment in autumn breeding there.

Offspring survival (from egg to metamorph) was low in both seasons. The combination of breeding effort and survival rates eventually resulted in a higher contribution of autumn breeding to the overall production of metamorphs. The overall low survival rates of offspring that we found is in line with previous field studies in anurans (e.g. Licht 1974; Banks & Beebee 1988) and can be caused by pond desiccation, predation, inter and intra-specific competition for food and parasitism or pathogen infections. Our study revealed no obvious effect of variation in predation on tadpole survival even if the predation pressure encountered

by tadpoles at the beginning of their development varies from site to site (but not between seasons). This may seem surprising since many studies experimentally demonstrated that predation cause substantial mortality to tadpole populations (e.g. (Tejedo 1993; Van Buskirk & Arioli 2005)). This may be due to the lack of information about predation during the first year of survey which reduced our statistical power or to the fact that causative factors are numerous and more complex to identify in the field. However, other studies reported no effect of predation on tadpole survival (Hartel *et al.* 2007) or even a positive effect (Barandun & Reyer 1997) probably due to predator-induced phenotypic plasticity. Nevertheless, our results suggest that the predation pressure is probably not a stronger constraint in one season than in the other.

Spring tadpoles should be exposed to more competitors during their development than autumn tadpoles since the majority of amphibian species in the local community breed in March and April. Nevertheless, we found no effect of interspecific competition on survival for any of the two seasonal tadpole cohorts. This seems surprising since parsley frog is a poor competitor as a tadpole compared to most species of the anuran community, in particular *Hyla meridionalis* and *Rana perezi*, *perezi* present in spring in permanent ponds (Richter-Boix *et al.* 2007b). On the contrary, in small temporary ponds and during autumn and winter season, Parsley frog tadpoles encounter mostly *Bufo bufo* and *Epidalea calamita* with even lower competitive abilities (Richter-Boix *et al.* 2007b). We hypothesized that interspecific competition effect was not detected in our study due to numerous uncontrolled sources of variation.

#### **Priority effects**

We revealed a striking negative effect of the presence of conspecific autumn tadpoles on the survival of spring tadpoles in the Parsley Frog. Previous studies have demonstrated the occurrence of such intraspecific priority effect in amphibians in experimental settings (Morin

*et al.* 1990; Eitam *et al.* 2005; Murillo-Rincón *et al.* 2017) but as far as we know, our study is the first evidence for intraspecific, inter-cohort competition in amphibians in nature. In the field, we observed in several occasions that large autumn tadpoles were eating freshly laid eggs of their own species, which could partly explain the lower hatching rate of spring eggs in presence of autumn tadpoles. Moreover, (Tejedo 1991) previously described how parsley frog tadpoles predates *Epidalea calamita* eggs. In this latter study, predaceous tadpoles were exclusively old tadpoles and they could cause a loss of 50 to 100% of the eggs. Oophagy has also been demonstrated to be responsible for interspecific priority effect between *Scaphiosus couchii* and *Bufo speciosus* (Dayton & Fitzgerald 2005). Intraspecific oophagy has been described on some anuran species (Summers 1999; Dayton & Wapo 2002) and has been proposed as an energetic opportunistic response in food shortage in temporary ponds.

However, the presence of autumn tadpoles also affect the larval survival (post-hatching) of spring tadpoles. This may reflect competition for resources between large autumn and small spring tadpoles as shown in *Rana arvalis* (Murillo-Rincón *et al.* 2017). Interference competition mediated by microorganism may also play a role: smaller tadpoles could display coprophagy instead of feeding on higher quality resources (Beebee & Wong 1992; Baker & Beebee 2000). This large priority effect between the two seasonal tadpole cohorts of parsley frog has a great impact on the overall efficiency of breeding: in most ponds, there could be only one successful breeding period, autumn or spring. Nonetheless, we found no indication that spring breeders select their oviposition site to avoid conspecifics, as other amphibian species sometimes do (Sadeh *et al.* 2009). Accordingly, the spawning probability was also unaffected by the presence of potential competitors or predators.

#### **Seasonal partitioning of breeding: a bet-hedging strategy?**

453 The temporal partitioning of breeding activity could reflect several evolutionary processes: 1)  
454 the existence of two specialized phenotypes either genetically determined (in which case we  
455 would expect temporal genetic differentiation between cohorts) or set by early environmental  
456 cues (phenotypic plasticity); 2) a use of alternative strategies by some or all individuals (bet-  
457 hedging). We previously demonstrated that the two temporal cohorts do not reflect two  
458 genetically distinct temporal populations (Jourdan-Pineau *et al.* 2012) but breeding phenology  
459 may still be set once for good for each individual. In this case, breeding in autumn or in spring  
460 could be determined by the physiological state (and sexual maturity) of the breeder and  
461 maintained year after year, by physiological constraints (typically the case for a capital  
462 breeder species which stores energy for future reproduction *e.g.* (Lardner & Loman 2003)). In  
463 a diversified bet-hedging strategy, individual breeding activities could vary from year to year  
464 (each year, individuals would “choose” one breeding season) or individuals could split their  
465 breeding effort between the two seasons in some or most years.

466 Based on our field survey, it appears that the bimodal breeding phenology of parsley frog is a  
467 typical diversified bet-hedging strategy. The large priority effect between the two seasonal  
468 cohorts, combined with high unpredictability of conditions that result in failure or success of  
469 entire cohorts, results in frequency dependent-selection and favour risk-spreading strategies:  
470 the best option is to develop in ponds with the smaller number of conspecific competitors.  
471 These conditions are found partly in autumn, when the habitat becomes favourable after the  
472 dry summer period, or in spring, as some of the autumn cohorts have died in the winter,  
473 leaving the habitat free. Poethke *et al.* (Poethke *et al.* 2016) developed a theoretical model in  
474 which they outlined this impact of competition on the evolution of bet-hedging strategy.  
475 Using a model for optimal germination fraction, based on field data on desert plants, Gremer  
476 and Venable (Gremer & Venable 2014) also showed that density-dependence could explain  
477 the observed bet-hedging strategy of germination spread in time (i.e. not all seeds at once).

478 In the parsley frog, our model shows that the observed mixed breeding strategy is maintained  
479 if the rate of initial success of the autumn cohort ( $q$ ) is between 0.2 and 0.8 (if females have  
480 on average 3 years of breeding in their lifetime) or between 0.35 and 0.65 (for 5 years of  
481 breeding). Those conditions are fulfilled according to our field estimates (0.43). We estimated  
482 the proportion of eggs laid in autumn by all breeders (0.57) but could not estimate this  
483 proportion at the individual level. Survival rates set in the model were based on our field  
484 estimates of survival from egg to metamorphosis; hence, we assumed similar survival after  
485 metamorphosis of the two cohorts. Unfortunately, we have no information about survival of  
486 parsley frog during its adult terrestrial life. However, the adult survival is an important  
487 parameter in our model since it determines the number of reproductive years. The mixed  
488 breeding strategy is less stable when the number of breeding opportunities per lifetime  
489 increases – as the risk is now spread over several successive years. Indeed, experiencing  
490 variation in reproductive success among those opportunities is less harmful when it is possible  
491 to try again the next year. A skeletochronology study conducted in a upland population in  
492 Spain indicated that the mean age of sampled parsley frog females was 5.01 years (with a  
493 standard deviation of 1.99) (Esteban *et al.* 2004). Assuming a minimal age at first  
494 reproduction of 1 year (as done by Esteban *et al.* 2004), this translates into an average number  
495 of reproductive years or females of  $n = 4$ . Our evaluation of the bet-hedging strategy with  $n =$   
496 5 is thus probably conservative.

497 We previously showed that the parsley frog successfully exploits two temporal niches in the  
498 Mediterranean region thanks to a high phenotypic plasticity of tadpole development to face  
499 very different seasonal environments (Jourdan-Pineau *et al.* 2012). Recently, the combination  
500 of phenotypic plasticity and bet-hedging has been theoretically investigated, suggesting that  
501 phenotypic plasticity could further minimize fitness variances caused by mismatches between  
502 phenotype and environment (Haaland *et al.* 2020; Rádai 2020). Interestingly, in the wolf

503 spider, temperature and day length leads to alternative developmental types within broods.  
504 This cohort splitting is both probabilistic and sensitive to environment , a phenomenon  
505 proposed as being a plastic bet-hedging strategy by Rádai (Rádai 2020). In this case, the  
506 various plastic phenotypes, triggered by environmental variations, constitute a bet-hedging  
507 response to grassland habitats with substantial and unpredictable year-to-year variation.  
508 The breeding strategy of parsley frog seems to constitute an original example of bet-hedging  
509 strategy driven by high environmental stochasticity and large inter-cohort priority effect.  
510 Characterizing adult survival and individual breeding pattern (using mark-recapture and  
511 parentage assignment of egg masses) would allow further refining our model and seeing how  
512 it can apply to other anuran species.

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