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Tadpole co-occurrence in ponds: When do guilds and time matter?

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ABSTRACT

An extensive meta-analysis study suggested that co-occurrence in communities composed mostly of ectotherm species would not be segregated in space. However, there is some evidence of segregated occurrence for ectotherms when guilds were considered in the analysis. Therefore the apparent randomness found in co-occurrence analyses may be a consequence of the pooling of heterogeneous groups. We tested the following hypotheses for tadpole pond communities: i) an analysis including all species will indicate random co-occurrence, ii) co-occurrence will differ between seasons, and iii) analysis of co-occurrence within guilds will reveal a segregated pattern. We sampled tadpoles in Caçapava do Sul, southern Brazil, in two seasons (spring, summer). Species co-occurrence patterns were analyzed using the C-score index and three null models. We collected 10,852 tadpoles of 21 species, which belonged to four guilds: benthics, nektonics, suspension-raspers, and suspension-filterers. The last two guilds contained only a single species each, and therefore the analyses were restricted to the benthic and nektonic guilds. We obtained contrasting results depending on the null model. We used results only from non-degenerate simulated matrices. The co-occurrence of all species analyzed together was random in both seasons. Benthic tadpoles showed a tendency to segregated co-occurrence in spring, and random co-occurrence in summer. Nektonic tadpoles showed random co-occurrence for the total and the spring matrices, and segregated co-occurrence in summer. Our results indicate that the strength of negative co-occurrence may depend on season and guild. We conclude that apparent random co-occurrences of ectotherm communities may actually result in a segregated structure when restricted to a single guild, which is in concordance with other studies where guilds were considered, and supports the hypothesis that an apparent lack of negative associations might be related to the study of heterogeneous groups.

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1. Introduction

To understand the relevance of assembly rules for community structure is one of the major challenges for ecologists. Why do some species never, and other species always occur together? Are competitive interactions patterning communities? A landmark in this debate was Diamond’s (1975) proposal that competition structures the distributions of birds on the islands of the Bismarck Archipelago, and the resulting controversy regarding how to detect competition from distributional data (Connor and Simberloff, 1979). Thirty years after Diamond’s work, many developments in theoretical frameworks and analytical tools have appeared, although these still need further investigation of their general applicability (Gotelli and McCabe, 2002).

In a meta-analysis of 96 presence-absence matrices, Gotelli and McCabe (2002) evaluated the co-occurrence patterns of different taxonomic groups, and found evidence of structure in most communities, mainly within homeotherms. They did not find indications of structured communities for fishes, amphibians, reptiles, and most invertebrates, except for ant assemblages. Based on this difference between homeotherms and ectotherms, it was suggested that the co-occurrence pattern of animal communities is ordered in a “continuum” (Gotelli and McCabe, 2002; Gotelli and Rohde, 2002). In this way, small animals, with less mobility and with small populations would be less subject to competitive pressures, and therefore would show co-occurrence patterns similar to...
those expected by chance. In contrast, large and/or highly mobile animals with large populations would tend to show segregated or checkerboard distributions (Gotelli and Rohde, 2002). It should be noted, however, that negative associations among species pairs may result from groups that use different habitats (Diamond and Gilpin, 1982). Species may not co-occur because they exploit different habitats (Sfenthourakis et al., 2005). Additionally, analyses including assemblages that are heterogeneous regarding resource use would dilute competitive effects and tend to show a lack of competition, when it in fact exists (Diamond and Gilpin, 1982). This last confounding factor is particularly relevant, because previous analyses of co-occurrences usually included all species in the assemblage and not only those for which a plausible competitive hypothesis could be suggested.

The guild approach is traditionally used in community ecology, and is based on the idea that communities are built from groups of species that share ecological and/or phylogenetic similarities (Blondel, 2003). It is not expected that all species will show competitive interactions, but only those that use resources in a similar way (Simberloff and Dayan, 1991). Therefore, the characterization and designation of an ecological guild constitute a fundamental step for the competitive interaction analysis, although the two processes are carried out independently (Gotelli and Entsminger, 2007). In fact, recent studies that evaluated co-occurrence within guilds found evidence of structure in some ectotherm communities. For example, Sanderson (2004) evaluated the confamilial co-occurrence of amphibians, and found checkerboard patterns for some cases that had been considered random in a previous analysis (Hofer et al., 1999). Hofer et al. (2004) found evidence of structure for lizards in a null-model analysis in which species traits were taken into account. Luiselli (2006) reviewed snake community studies and found evidence of competition for habitat depending on the snake guild and the geographical region. Luiselli (2008) showed that competition should be important to freshwater turtle communities. Heino (2009) studied stream macroinvertebrates and found non-random patterns of co-occurrence within guilds, and Dilling et al. (2007) reported segregation for some insect guilds associated with the hemlock Tsuga canadensis. Jenkins (2006) studied co-occurrence of zooplankton communities and found evidence of segregated patterns in a null-model analysis in which species was proportional to area. All these studies based on ectotherm communities challenge the continuum hypothesis, even though random co-occurrence has been reported for other ectotherm guilds (e.g., Krüger et al., 2010).

Pond communities provide a good opportunity to study assembly rules, because they have easily recognizable limits, similarly to islands, where assembly rules have historically been studied (Chase and Leibold, 2003). The presence of tadpoles in a pond relies, in the first instance, on the site that adults choose for egg deposition. The type of habitat selected strongly influences the development of most species (Alford, 1999; Ryan and Winne, 2001). Larvae that develop in permanent habitats are less exposed to hydric stress, but are generally exposed to a wide variety of potential predators and competitors (Alford, 1999). Thus, different traits are expected to be required to occupy ponds in the different positions along the permanence gradient. Changes in composition of species across gradients of pond permanence have been documented (Richter-Boix and Llorente, 2007; Snodgrass et al., 2000; Welborn et al., 1996).

The tadpole community structure in ponds has been attributed mostly to predation, competition, and degree of permanence of the pond. Heyer et al. (1975) suggested that habitat permanence would determine species composition in ephemeral ponds, whereas competition and other biotic interactions would have major roles in more-permanent habitats. Wilbur (1987) manipulated these three factors experimentally, and concluded that the importance of each factor depends on the species studied and on the habitat, in agreement with Heyer et al. (1975). Additionally, the results of Wilbur (1987) indicated that competition is a patterning pressure in small and short-permanence habitats, and that predators attenuate competition in more-permanent ponds.

Subsequent experimental and observational studies that assessed segregation patterns of tadpoles attributed different values to competitive interactions. Laboratory and mesocosm experiments indicated that competition has an important role in structuring tadpole communities (e.g., Morin, 1987; Morin and Johnson, 1988; Parris and Semlitsch, 1998). On the other hand, observational studies tended to assign less importance to competitive interactions (e.g., Eterovick and Barros, 2003; Rossa-Feres and Jim, 1996; Wild, 1996). In fact, some authors have questioned the prevalence of competition for communities in natural conditions (Chase and Leibold, 2003), and hypothesized that under such conditions, heterogeneity and predation would attenuate and/or prevent competition (Lawler and Morin, 1993; Morin, 1983).

Amphibian breeding activities are strongly related to time-structured variables (see Both et al., 2008; Canavero and Arim, 2005; Prado et al., 2005; Toft, 1993). Studies that assessed the temporal occurrence of amphibians, unsurprisingly, found segregated co-occurrence patterns (e.g., Canavero et al., 2009; Sanchez et al., 2009). This seasonal component of amphibian communities could prevent both adult and larval species co-occurrence (Glasser, 1989), and should be taken into account even in spatial studies.

We investigated the co-occurrence of tadpoles in pond assemblages considering all-species records, and also restricting analyses to within guilds and seasons. We hypothesized that: i) in the search for structure using all species, the co-occurrence pattern would be random as a result of the dilution effect; ii) pond conditions vary greatly in time, and thus co-occurrence patterns may differ between seasons; and iii) within guilds, the co-occurrence models would reveal a segregated pattern.

2. Materials and methods

2.1. Study area and sampling

The study area comprised rural sites in the municipality of Caçapava do Sul, Rio Grande do Sul, southern Brazil (between 30° 57' and 31° 03' S, 53° 25' and 53° 39' W), located in the Pampa biome (IBGE, 2004). The landscape is composed by natural grasslands and pastures, with small forest patches on hills and in valleys. The grasslands are mostly used for cattle and sheep grazing. The climate is classified as Temperate Humid, with an annual mean temperature of 16.8 °C and annual precipitation of 1588 mm (Maluf, 2000).

We sampled tadpoles in 38 natural ponds of different sizes and degrees of permanence. Each pond was sampled twice: in November 2007 and January 2008 (respectively austral spring and summer). All ponds were sampled within seven consecutive days in spring, and five days in summer. We measured the area and depth of each pond in each collection event. For depth, we took the mean of three to six measurements obtained in each pond, depending on its size. A detailed description of pond descriptors is available in Both et al. (in press). Of the 38 ponds with water in the spring sampling, only 21 still contained water in the following summer sampling event. The volume reduction or drying of ponds observed in summer months is related to the increase of evaporation rates in subtropical regions (Both et al., 2009). Tadpoles were collected with a dip net (0.4 × 0.3 m; 0.5 mm mesh), which was swept all around the pond once only. The collected tadpoles are housed in the...
herpetological collection of the Universidade Federal de Santa Maria (ZUFSM), Brazil.

2.2. Presence/absence matrices

First, we compiled two presence/absence matrices: i) a spring occurrence matrix, containing the species recorded in 38 ponds and ii) a summer occurrence matrix, including the 15 species found in 21 ponds that retained water. In order to test the hypothesis that competitive interactions would be best reflected in intra-guild groups, each of the two matrices was divided according to guilds. Criteria used for guild designation followed Alig and Johnston (1989) and are detailed in Both et al. (in press).

2.3. Co-occurrence patterns

We tested if the co-occurrence pattern observed in each matrix differed from that expected by chance, comparing it with 30,000 matrices randomized according to a null model. We used the C-score index of co-occurrence (Stone and Roberts, 1990), which measures the average number of checkerboard units (CU) between all pairs of species (Gotelli and Entsminger, 2007). In a competitively structured community or one in which species are segregated according to habitat, the observed C-score should be significantly higher than scores expected by chance according to a null model.

The choice of the null-model algorithm is a crucial step, where biological and statistical restrictions should be carefully taken into account. We built the null models based on three different algorithms. In the first null model, we fixed the total number of species occurrences and the species richness in each site. This is a traditional although very conservative null model (Gotelli, 2000; Gotelli and Graves, 1996). This model maintains the observed rarity or evenness of the original matrix. Similarly, this model maintains the pattern of rich and poor sites in the randomized matrices. Fixing total species occurrences and species richness at each site results in a very conservative null model because usually few combinations of simulated matrices can be produced (Gotelli, 2000). As an alternative null model, we fixed species occurrences, but weighted species richness in sites by their mean depths. We chose mean pond depth to weight models based on the results of Both et al. (in press), which showed that depth was the best variable that explained tadpole richness and guild composition of the present data. This fixed-weighted model is less conservative and takes into account the importance of the environmental descriptor, in addition to competition, in structuring the community (Heino, 2009; Jenkins, 2006). This procedure, however, can produce degenerated matrices, where some sites contain no species, which in turn may increase Type I error (Connor and Simberloff, 1979). One can argue that degenerated matrices are not too artificial because ponds can be naturally “degenerated”, i.e., we can find filled ponds with no tadpoles. Taking these considerations into account, we built a third null model fixing species occurrences, weighting sites by mean depth, but excluding degenerated matrices. The null matrices were generated by sequential swapping (Gotelli and Entsminger, 2003, 2007). The analyses were carried out using EcoSim 7.72 software (Gotelli and Entsminger, 2007).

3. Results

We collected 10,852 tadpoles of 21 species, belonging to five families: Cyclocephalidae (1), Hylidae (8), Leiuperidae (5), Leptodactylidae (6), and Microhylidae (1). The 21 species were initially assigned to four ecomorphological guilds: benthics, nektonics, suspension-raspers, and suspension-filterers. Two of the guilds were composed of a single species each. Phyllomedusa iheringii is a suspension-rasper tadpole, and Elachistocleis bicolor is a suspension-filterer. Both species were included only in the all-species analyses. The species list and guild designations are presented in Table 1.

The co-occurrence of all species analyzed together was random in spring ($P = 0.673$) and in summer ($P = 0.512$) in the fixed–fixed null model (Table 2). However, species co-occurrences were non-random in both seasons when using the fixed-weighted model which included degenerated matrices ($P = 0.032$, spring; $P = 0.033$, summer). In the fixed-weighted model discarding degenerated matrices, all-species co-occurrence was random in spring ($P = 0.140$) and tended to be non-random in summer ($P = 0.064$). The co-occurrence for benthic tadpoles in spring also presented a low probability of randomness ($P = 0.058$), although it was not different from that expected by chance in summer ($P = 0.864$) in the fixed–fixed model. In the fixed-weighted model including degenerated matrices, benthic tadpoles also showed a segregated occurrence in spring ($P < 0.001$), and a random pattern in summer ($P = 0.158$). In the fixed–weighteded model excluding degenerated matrices, benthic tadpoles showed a random co-occurrence in summer. It was not possible to obtain enough non-degenerated null matrices based on the benthic occurrence matrix in spring.

Nektonic tadpoles showed a random co-occurrence in spring ($P = 0.404$), and only a weak evidence of segregated co-occurrence in summer ($P = 0.087$) (Table 2) in the fixed–fixed model. In the fixed-weighted model, the co-occurrence of nektonics was non-random for the spring and summer matrices ($P < 0.001$ for both cases). In the fixed-weighted model without degenerated matrices, nektonics showed a segregated pattern ($P = 0.002$). It was not possible to build enough non-degenerated null matrices for the nektonic occurrence matrix in spring.

4. Discussion

Tadpole co-occurrence in these ponds showed contrasting patterns depending on the null model used. The results obtained based on the fixed–fixed and fixed-weight excluding degenerated matrices models were mostly congruent. They tended to indicate that random co-occurrence patterns prevail when all species are analyzed together (Table 2). However, it was not possible to generate the null universe without degenerated

| Species Gallus Guild Frequency |
|-------------------------------|-------------------|
| Limmomedusa macroglossa | Benthic | 3 |
| Dendropsophus minutus | Neotonic | 16 |
| Hypsiboa albo punctatus | Benthic | 4 |
| Hypsiboa pulchellus | Benthic | 10 |
| Phyllomedusa iheringii | Suspension-rasper | 5 |
| Pseudins minuta | Neotonic | 7 |
| Scinax fusovarius | Neotonic | 12 |
| Scinax granulatus | Neotonic | 11 |
| Scinax nasicus | Neotonic | 10 |
| Physalaemus biligonigerus | Benthic | 12 |
| Physalaemus cvieri | Benthic | 11 |
| Physalaemus graciilis | Benthic | 23 |
| Physalaemus riograndensis | Benthic | 4 |
| Pseudopaludicola falipes | Benthic | 7 |
| Leptodactylus chaquensis | Benthic | 2 |
| Leptodactylus fuscus | Benthic | 1 |
| Leptodactylus graciilis | Benthic | 3 |
| Leptodactylus latinosus | Benthic | 3 |
| Leptodactylus latrans | Benthic | 5 |
| Leptodactylus mystacinus | Benthic | 2 |
| Elachistocleis bicolor | Suspension-filterer | 15 |
and (***) when rejected are indicated by (*) when the weighted II they were not permitted. Models in which the null hypothesis was rejected are indicated by (***) when the P-value was less than 0.05 (**) when $P < 0.01$ and (***) when $P < 0.001$.

### Table 2

Values of observed C-score and mean of C-scores obtained in 30,000 null simulations. Models fixed-weighted I, include degenerated matrices and in the fixed-weighted II they were not permitted. Models in which the null hypothesis was rejected are indicated by (*) when the P-value was less than 0.05 (**) when $P < 0.01$ and (***) when $P < 0.001$.

<table>
<thead>
<tr>
<th>Species/guild</th>
<th>Season</th>
<th>C-score (obs)</th>
<th>Model</th>
<th>C-score (sim)</th>
<th>$P (obs &gt; sim)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>All species</td>
<td>Spring</td>
<td>Fixed–fixed</td>
<td>20.08</td>
<td>0.673</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Fixed–weighted I</td>
<td>17.54</td>
<td>0.032*</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>Fixed–weighted II</td>
<td>18.63</td>
<td>0.140</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Summer</td>
<td>Fixed–fixed</td>
<td>8.53</td>
<td>0.0512</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Fixed–weighted I</td>
<td>7.30</td>
<td>0.033*</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Fixed–weighted II</td>
<td>7.57</td>
<td>0.064</td>
<td></td>
</tr>
<tr>
<td>Benthics</td>
<td>Spring</td>
<td>Fixed–fixed</td>
<td>14.79</td>
<td>0.058</td>
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<tr>
<td></td>
<td></td>
<td>Fixed–weighted I</td>
<td>11.50</td>
<td><strong>&lt;0.001</strong>*</td>
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<tr>
<td></td>
<td></td>
<td>Fixed–weighted II</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Summer</td>
<td>Fixed–fixed</td>
<td>4.54</td>
<td>0.864</td>
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<td>Fixed–weighted I</td>
<td>3.30</td>
<td>0.158</td>
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<tr>
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<td></td>
<td>Fixed–weighted II</td>
<td>4.28</td>
<td>0.369</td>
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<td>Neptotics</td>
<td>Spring</td>
<td>Fixed–fixed</td>
<td>32.76</td>
<td>0.040</td>
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<td>Fixed–weighted I</td>
<td>18.36</td>
<td><strong>&lt;0.001</strong>*</td>
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<tr>
<td></td>
<td></td>
<td>Fixed–weighted II</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Summer</td>
<td>Fixed–fixed</td>
<td>15.67</td>
<td>0.087</td>
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<td></td>
<td></td>
<td>Fixed–weighted I</td>
<td>9.79</td>
<td><strong>&lt;0.001</strong>*</td>
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<tr>
<td></td>
<td></td>
<td>Fixed–weighted II</td>
<td>13.42</td>
<td>0.002*</td>
<td></td>
</tr>
</tbody>
</table>

matrices for the benthic and nektontic matrices in spring. This was due to the presence of a proportionally large number of rare species and/or sites with only one species, i.e., a large number of zeros.

The fixed-weighted model with degenerated matrices indicated that five of the six matrices were non-random and showed low probabilities. Also, all probabilities were notably low in this model when compared with the other two models, which either prevented or discarded degenerated matrices. Based on the fixed-weighted model with degenerated matrices, we would conclude that tadpole communities are segregated independently of the guild and the season. The effect of degenerated matrices in reducing probabilities was first noted by Connor and Simberloff (1979). Gotelli (2000) tested this effect and found that for most matrices it was less severe than the effect of the type of null model used. In this study, the degenerated effect was strong and probably due to the large proportion of zero values in the matrices. Accordingly, in the remaining discussion we will focus on the results from the other two models.

The results produced by the fixed–fixed model for both seasons supported our first hypothesis of lack of a segregation pattern when all species are analyzed together. Using the fixed-weighted non-degenerated model this hypothesis was also supported, although a weak evidence for segregation was observed in summer season. In the case of the all-species analysis, non-competitive species pairs should dilute the effects of negative associations among interacting species. The comparison restricted to species within guilds reinforced the suggestion of co-occurrence analysis for subsets of potentially interacting species (see Diamond and Gilpin, 1982; Gotelli and McCabe, 2002; Simberloff and Dayan, 1991). A guild approach reduces the variability and makes co-occurrence analysis more realistic because it considers species similarity (Diamond and Gilpin, 1982; Kelt and Brown, 1999) and evaluates associations within the habitat shared by species (Gotelli and Graves, 1996). This makes the results easier to interpret. On the other hand, in an analysis including species from all guilds, segregated patterns could appear based on the habitat checkerboard. This is a plausible hypothesis for the segregated co-occurrence pattern found for all species in the fixed-weighted non-degenerated model. For instance, $P. iheringii$, a suspension-rasper, is associated with deep ponds (Both et al., in press), and benthic species of Leptodactylus are associated with ephemeral ponds. Combining these species in a single matrix would result in a checkerboard distribution, although it would not be a consequence of competition.

Our second hypothesis predicted that co-occurrence patterns for each guild would differ over time, as previous studies have shown distinct amphibian breeding seasonality in subtropical regions (Both et al., 2008; Canavero et al., 2008; Sanchez et al., 2009). Of the 21 species found, 18 occurred in spring and 15 in summer, showing that some species pairs did not co-occur. Since tadpole guilds are intrinsically linked with the water column and/or their swimming abilities (Alrig and Johnston, 1989), environmental changes in ponds that occur across seasons constitute a resource change for tadpoles from each guild. As expected, co-occurrence varied between the two seasons, but the specific pattern of co-occurrence was dependent on guilds.

The nektontic tadpoles showed a random distribution in spring, but presented a segregated pattern in summer based on the fixed-weighted non-degenerated model. The fixed–fixed model did not reject the null hypothesis, but showed a tendency of segregation given by the low probability of randomness ($P = 0.087$). The ample availability of deep ponds in spring may prevent competition among nektontic species. Species could share the plentiful resources and habitat represented by deep ponds. Conversely, the reduction in volume or even complete drying of ponds in summer may restrict occupancy to a few suitable sites, and thus competition may prevail. Instead, the benthic tadpoles tended to show a segregated distribution in spring, and a random one in summer. Shallow ponds should be scarce in spring, and benthic species would compete for ephemeral ponds or the margins of deep ponds. As the volume of ponds decreases in summer, the plentiful availability of shallow sites prevents competition among species. It should be noted that the preference for shallow ponds of benthics in this study is probably related to the nature of the species belonging to this guild, which were mostly from the families Leptodactylidae and Leiopteridae.

The importance of depth in modulating tadpole interactions has been documented in previous field studies. Eterovick and Fernandes (2001) analyzed the microhabitat use of tadpoles in streams and found that the position in the water column and depth, together with time of occurrence, were the best variables to distinguish microhabitat use. Species segregation in the water column was suggested by Rossa-Feres and Jim (1996), and Both et al. (in press) reported guild replacement along gradients of depth and permanence of water.

Welborn et al. (1996) suggested that the gradient of pond permanence represents a stress gradient for tadpoles. Deep ponds in spring and shallow ones in summer would represent stress for, respectively, benthic and nektontic species. It has been suggested that under stress conditions, co-occurrence would be positive or random because competition would be ameliorated (Bertness and Callaway, 1994). Our analyses, however, indicated potential segregated occurrences during stress conditions. Similarly to our results, evidence of segregated occurrences under stress conditions was found for sand-dune plant communities (Maltez-Mouro et al., 2010), soil arthropods (Pitzalis et al., 2010), and Mediterranean tadpole communities (Richter-Boix et al., 2007b).

Direct competition among tadpoles is not the only explanation for the observed patterns. Segregation among tadpoles that occupy small and ephemeral ponds may be the result of the choice of breeding sites by adults. There is experimental evidence that species typical of temporary ponds are usually poor competitors in permanent ponds (Richter-Boix et al., 2007a). Other studies showed that adults of some species choose breeding sites taking into account the presence of predators and competitors, in addition to the availability of oviposition sites (Resetaritis and Wilbur, 1991).
and that the order of occupancy should influence the outcome of competitive interactions (Lawler and Morin, 1993). Therefore, the effects of one species on others would not be solely related to resource depletion and its consequences for growth and survival rates, as observed in classical competition experiments (e.g., Tilman et al., 1981). Instead, the observed segregation could be a result of interference (inhibition) mediated by priority colonizer effects.

Historical processes or phylogeny are currently employed to explain segregated distribution patterns, mainly in regional- to studies (Bowers and Brown, 1982). On this scale, allopatric speciation can generate checkerboard patterns for some species pairs. However, phylogeny could explain segregation even at small scales (Taylor and Gotelli, 1994). The relationship between guild and phylogeny was not tested for tadpoles, and deserves future investigation (McDiarmid and Altig, 1999), as well as the role of phylogeny was not tested for tadpoles, and deserves future investigation of seasonal variation in amphibian activity. J. Nat. Hist. 43, 45–48.


