Trophic-specific responses to migration in empirical metacommunities

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Abstract
The metacommunity, as it evolved from Levins’s metapopulation, provides a framework to consider the spatial organization of species interactions. A defining feature of metapopulations and metacommunities is that organisms (populations or communities) are connected via migration. An important result from Levins’s metapopulation work – that increasing migration lowers regional extinction probability – is often incorporated into conceptions of metacommunities; however, this may not hold true for multiple interacting metapopulations (metacommunities). We report results from a metacommunity field experiment conducted with a tropical terrestrial leaf litter macro-arthropod community. We show that migration induces regional extinctions of predators without changing the predator community composition. For non-predators we found no evidence of regional extinctions, but a significant change in community composition. Our result corroborates the findings of a prior similar metacommunity experiment with a temperate forest leaf litter community. The concordance between these experiments, even with vastly different communities, highlights the importance of considering trophic and non-trophic community structure to understand metacommunity dynamics, and suggests a potential connection between migration rates and trophic-specific responses in ecological communities.

Keywords: metacommunity, metapopulation, migration, non-trophic interactions, space
Introduction

The theory of metapopulations has become a standard way of thinking about simple population dynamics, and its success has stimulated a seemingly obvious extension, the metacommunity (Levins 1969; Wilson 1992). As originally envisioned, a metacommunity is a collection of interacting populations of different species in which extinction and migration occur on a regular basis. This results in patchiness that creates subcommunities, which may be distinct in species composition. It might be argued that MacArthur and Wilson’s theory of island biogeography was the first metacommunity theory, and perhaps the most elegant, in which patchiness is provided by the existence of islands (MacArthur and Wilson 1964). From these formulations, an important conclusion of metapopulation theory has been tacitly incorporated as an obvious corollary of metacommunity theory—that increasing migration lowers overall extinction probability. While it may be a reasonable proposition at first glance, further reflection on the assumption suggests that the expectation of lowered extinction is not universal in the metacommunity context (Vandermeer et al. 1980; Caswell and Cohen 1991).

Huffaker’s classic experiment might be thought of as a canonical case study that supports a link between increasing inter-patch migration and lower extinction. When isolated to a small feeding area, both predator and prey mites go extinct (predator eats prey to extinction then itself goes locally extinct)—yet when many smaller feeding areas were arranged to facilitate local dispersal of prey (but not predators), apparently stable oscillations result (Huffaker 1958). A similar experimental set-up that yielded stable oscillations also included predator-focused dispersal limitations (Huffaker 1958). A similar pattern was also observed in Gause’s experimental system of protozoans twenty years prior (Gause et al. 1936). In these examples, migration appears to decrease the probability of extinction.

Alternatively, it is not difficult to imagine the reverse outcome in other systems: a case in which migration might increase extinction probability. For example, in a two-predator one-prey situation in which spatial structure allows for a segregation of the two predators in space, increasing the predator migration rate could increase intraguild antagonism, leading to one of the predators dominating, and a reduction of total species diversity from three to two. Even in cases without intraguild competition, the simple dispersal of any intermediate predator can theoretically cause trophic instabilities due to more lag in population dynamics (Jansen 1995). Thus, elevated migration rates could result in either increased or decreased species diversity, depending on the strength or timescale of antagonistic (or even facilitative) interactions across ecological guilds (Guzman et al. 2019). The simple migration-extinction equilibrium of island biogeography and metapopulation theory may yield predictions that are not generalizable for more complex community structures.

The role of migration in rescuing unstable populations has been highlighted so often that it is usually taken for granted, but that result is not theoretically inevitable (Simberloff and Cox 1987). The ability to simulate a wide range of metacommunity dynamics suggests the need for an empirical approach. One of the key shortcomings of much of the experimental metacommunity work lies in its simplification of community interactions (Polis et al. 1989), where studies often only consider a subset of species with simple trophic structure (e.g. consumer-resource pairs) Warren 1996; Shurin 2001; Kneitel and Miller 2003; Cadotte 2006; Fox et al. 2017). Although there are some notable experiments which attempt to include some of the trophic and non-trophic realism of communities (Neill 1974; Vandermeer et al. 1980), surprisingly few experimental studies have focused on empirically realistic metacommunities.

One of the early attempts to study the role of migration in empirical metacommunities used leaf litter macro-arthropod communities and found that the predator guild (defined taxonomically) decreased in richness when random migration was induced, while non-predator richness was unaffected by migration (Vandermeer et al. 1980). These results suggest that conclusions about metacommunity structure may, at least in some contexts, be trophic-specific. The dependence of community dynamics on trophic structure has been noted in some well-known debates, for example, in considering whether communities tend to be controlled primarily by consumers (top-down) or producers (bottom-up) (Hairston et al. 1960), and why trophic cascades operate differently in aquatic versus terrestrial systems (Strong 1992). Clearly, a wide range of ecological processes interact with migration in real metacommunities, including higher-order interactions (or trait-mediated indirect interactions), which recent work suggests may be more determinant of community structure than the more direct, lower-order or pairwise species interactions (Werner and Peacor 2003; Bairey et al. 2016; Grilli et al. 2017; Terry et al. 2017).

The empirical result that experimental migrations make a difference in metacommunity structure for predators, but not for non-predators (Vandermeer et al. 1980), was found in a species-poor temperate deciduous forest (Michigan, USA). Here we revisit the experiment conducted by Vandermeer et al. (1980) in a more speciose montane tropical agroecosystem. We considered that in the tropics, the hypothesized stability-preserving aspect of high biodiversity (McCann 2000) could overwhelm any special effect of a strongly antagonistic predator species, which was proposed to explain the results of the earlier temperate zone study. Accordingly, we sought to investigate how migration affects leaf litter community richness and composition, focusing especially on the effects at different trophic levels. Based
on underlying assumptions about the importance of migration in maintaining biodiversity, we hypothesized that curtailing local migration would reduce local species diversity, and that this effect would be observed at all trophic levels.

**Methods**

**Study region and design**

This study was conducted at Finca Irlanda, an organic shaded coffee agroecosystem in the Soconusco region of Chiapas, Mexico. The study site was on a subset of land recently transitioned from rustic coffee production to a forested reserve. The experimental set-up was positioned adjacent to a patch of invasive golden bamboo, the litter of which created a uniform mat.

Leaf litter was collected from a well-forested area of the reserve, homogenized, and separated into 10 mesocosms, each of which were 0.5 m² in area and separated from one another by 1 meter. Five mesocosms were positioned on either side of a walking trail. No physical barriers prevented migration between mesocosms. Inter-patch migration was therefore possible, but we assumed it was sufficiently infrequent such that it could be ignored. We assumed organisms would not prefer to leave a mesocosm of leaf litter of the same composition from which they were originally collected to migrate across a relatively inhospitable mat of dried litter from an invasive bamboo species.

Half of the mesocosms were assigned as controls and half as treatments. To simulate migration among treatments, one-quarter of the litter in each mesocosm was removed and replaced with the same amount of litter from a different mesocosm. The transfer schedule was set so that a different quarter of the mesocosm was migrated during each transfer event, and the replacement pattern was randomized so that each mesocosm received and contributed litter to a different, randomly assigned mesocosm. Transfers were done every 4 days for 16 days. Migration was not manipulated among control mesocosms, though one-quarter of the litter was lifted, agitated, and replaced in the same mesocosm every four days to control for the disturbance of the litter transfers between treatment mesocosms. All mesocosms were harvested on day 20. This time scale is comparable to Vandermeer et al.’s 1980 experiment which ran for 30 days.

After harvesting, the litter was sieved using 3-mm meshes to remove coarse detritus. Each sample was searched by four people for 20 minutes, and all encountered organisms were individually removed and placed in alcohol. This technique did not likely capture all organisms found within the mesocosms, but we expect that any bias toward certain groups of organisms was standardized across all samples, which we assured by blinding the sample labels throughout the sorting process. Individuals were sorted into orders or families and identified to morphospecies. Morphospecies were then classified as either predators or non-predators, where predators included spiders, Staphylinidae beetle larvae, pseudoscorpions and centipedes.

**Statistical methods**

To compare the number of species in our control and migration treatments, individual-based rarefaction curves were calculated for the whole dataset and for each trophic group (predators and non-predators) separately. Rarefactions followed the now standard methodology of resampling the list of species observations with replacement at increasing numbers of individuals (Gotelli and Colwell 2001). 100 resamples were conducted for each level of individuals sampled, and the mean number of species for a given density of individuals was calculated.

While there are standard methods to extrapolate the number of species (Colwell and Coddington 1994; Chao et al. 2014) and compare the overall shape of rarefaction curves (Cayuela et al. 2015), we were interested in the statistical differences between our rarefaction curves across the range of resampled levels. To assess differences between the control and migration treatments, we conducted a bootstrap procedure to compare the difference in the mean number of species at every re-sampling level, \( x_i \), along the rarefaction curves. For a given \( x_i \), 100 random draws from the observed datasets for the control and migration, \( D_c \) and \( D_m \), were used to calculate the mean observed number of species, \( \bar{S}_c \) and \( \bar{S}_m \), for sampling level \( x_i \). These values were used to calculate the observed difference in the number of species, \( \bar{S}_c - \bar{S}_m = \Delta_{OBS} \) for resampling level \( x_i \). The observed data, \( D_c \) and \( D_m \), were then pooled together to create \( D_p \), which was then randomly partitioned into null data sets 100 times for the control and migration treatments, \( N_c \) and \( N_m \). \( N_c \) and \( N_m \) were then randomly sampled 100 times at \( x_i \) sampling level to calculate mean number of species, \( S_{N_c} \) and \( S_{N_m} \), sampled at \( x_i \) for both null data sets. These values were then used to calculate the null difference in the mean number of species sampled, \( S_{N_c} - S_{N_m} = \Delta_{NULL} \). This gives us a distribution of \( \Delta_{NULL} \) which was then compared to \( \Delta_{OBS} \) to calculate the probability of observing \( \Delta_{OBS} \) for a given \( x_i \) if \( D_c \) and \( D_m \) come from the same statistical population. The probability, \( p \), is calculated by \( p = 1/n \sum q \) where \( q \) is the number of times that \( \Delta_{OBS} \geq \Delta_{NULL} \) and \( n \) is the number of \( \Delta_{NULL} \) values in the distribution. This procedure was repeated for every value of \( x_i \), where the rarefactions of both treatments overlap. The supplementary material contains a graphical walkthrough of the statistical test (Figure S1) and a link to a repository with the R code for the test.

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To look at patterns in community composition for both treatments (increased migration and control) we used both Bray-Curtis and Jaccard distances as measures of dissimilarity at the patch scale. This allowed us to look at the data as weighted by abundance of morpho-species (Bray-Curtis) as well as just looking at the presence-absence (Jaccard). Analysis of Similarity tests (ANOSIM) were used to calculate statistical differences in community composition for both dissimilarity measures between our control and migration treatment. The NMDs plots, ANOSIM tests, and calculations of dissimilarity measures were implemented with the ‘vegan’ package (Oksanen et al. 2010) in R (R Core Team 2019). The ‘anosim()’ function of the ‘vegan’ package used with 5000 permutations to calculate the R statistic and the p-values. For the Bray-Curtis distance ANOSIM we created a community matrix where the rows are the separate patches, the columns are morpho-species, and the entries are the number of morpho-species for a given patch. With the Jaccard distance ANOSIM we created a similar matrix, but where the entries are the presence (1) or absence (0) of morpho-species.

To quantify the impact of migration in the communities across treatments we infer the local extinctions dynamics in our experiment by using the distribution of rare species across treatments. Given that the leaf-litter substrate was homogenized prior to the experimental set-up we assume that observed differences result from the dynamics in different treatments. We defined rare species in two ways here. First, by being a singleton (i.e. having an abundance of 1 in a single patch) across all patches (control and migration), and second by having an abundance that is less than the mean morpho-species abundance in the community (“relatively rare species”) (5.12 for predator community and 4.44 for non-predator community). This comparison of ‘rare’ morpho-species is done with both the community of predators and non-predators separately to understand how dynamics differ across trophic position across treatments.

Results

While there was no statistically significant difference between the control and migration treatments for the whole community (Figure 1A) or the non-predator community (Figure 1C), we did observe a significant difference in species richness between the control and migration treatments within the predator community (Figure 1B). This difference between control and migration treatments for the predators starts at just nine individuals sampled and remains significant for the rest of the overlap between the two curves.

Community analysis

Analysis of similarity (ANOSIM) showed no differences between the community composition for the predators for both Bray-Curtis (R=0.12, p=0.182) and Jaccard distance (R=0.16, p=0.081), while the non-predator community showed significant differences for both Bray-Curtis (R=0.478, p=0.0172) and Jaccard distance (R=0.332, p=0.0523) (Figure 2). The amount of species overlap between control and migration treatments was 51% for predators and 40% for non-predators. Additionally, we see that rare morpho-species, as defined by being singletons and less abundant than the mean morpho-species abundance in the community are more common the non-migration treatment for predators with no apparent difference in the non-predator community Table 1. This suggests that relatively rare morpho-species in the predator community are more prone to local extinctions than rare morpho-species in the non-predator community when migration occurs.

Discussion

Our results showed that, even after scaling up the biodiversity background to a tropical leaf litter community with over 100 species, predator guild species diversity decreased significantly with migration. It is notable that even the coarsest distinction of trophic complexity (predators and non-predators) provides insights that do not emerge when analyzing the community as a whole. These findings echo those of the earlier study (Vandermeer et al. 1980) which was done with a lower-biodiversity temperate leaf litter community. Results from both this and the earlier study seemingly contradict a main conclusion of basic metacommunity theory: that migration increases species’ persistence, and thus also regional richness. While theoretical treatments of metacommunities acknowledge the potential complexity of community structure and its effect on migration (Caswell and Cohen 1991; Mouquet and Loreau 2002; Econom and Keitt 2008), it remains that simplified metacommunity theory generates the prediction that migration will tend to cause species diversity to increase, a result in concordance with the original MacArthur-Wilson, Levins-Heatwole framework (Heatwole and Levins 1972; 1973; Levins et al. 1973).

Importantly, there is also no evidence, to our knowledge, that suggests that leaf litter communities in the temperate or tropical zones are organized in such a way that predisposes them to the results of both of these studies. These consistent results with distinct communities in distinct regions suggest that there may be some generality in the way that migration impacts trophic guilds.

While it is of particular interest that effects of migration fall along the lines that delineate trophic position in the community, that does not necessarily imply a trophic mechanism to explain the observed
In our experiment, we manipulated migration rates such that there were equivalent potential migration rates for predators and non-predators. This could be an important caveat to our study’s generalizability, given the potential differences in the realized predator and non-predator migration rates. Migration should increase diversity initially, as homogenization occurs, but trophic and non-trophic interactions can act to increase or decrease diversity after this initial homogenization. Our result of lower predator richness with migration runs counter to this expectation of increased diversity with homogenization, and thus indicates that our experimental time frame was appropriate to assess changes in these communities. We found no change in the richness of non-predators, but have no reason to suspect
that homogenization effects would operate on a different time scale for predators and non-predators in the leaf litter community, particularly given that we manipulated the potential migration rates to be equal across trophic levels.

The work reported herein sits comfortably with the current enthusiasm for metacommunities, a framework originally suggested by Wilson (1992). It is substantially similar to the framework of MacArthur and Wilson’s original offering, in which 1) ecological dynamics occur locally, with species interactions (of various forms) determining which species will survive and which will locally perish, while 2) the more regional process of migration continually feeds these local communities, counteracting local extinctions with regional migrations to provide the expected equilibrium (MacArthur and Wilson 1967). Eschewing some recent complexities (Leibold et al. 2004), we consider a metacommunity as structured in the original sense of Wilson (1992), wherein ecological dynamics occur at a local level, but local patches affect one another through dispersal. Our experiment interrogates the consequences of migration, but more specifically explores the interaction between community structure and the dynamics of migration. Our results highlight the importance of considering trophic and non-trophic structure when evaluating metacommunity dynamics.

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References


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Supplementary material (available online as Appendix oik-06342 at <www.oikosjournal.org/appendix/oik-06342>). Appendix 1

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Figure Legends

**Figure 1:** (A-C) Individual-based rarefaction curves for A.) whole community (green), B.) predators (red), and C.) non-predators (blue). Treatments are shown in lighter colors (control) and darker colors (migration). One standard deviation (based on the 1000 random draws) is plotted in the shaded areas around the curves. The vertical dashes above the curves in B.) represent a statistically significant (p < 0.05) difference in the number of species for a given number of individuals sampled between the control and migration treatments.
Figure 2: NMDS plots for predator community and non-predator community. a.) and b.) are made using Bray-Curtis dissimilarity and c.) and d.) are using Jaccard. ANOSIM showed no differences between control and migration treatments for the predators (Bray-Curtis $R=0.12$; $p=0.182$ & Jaccard $R=0.16$ $p=0.081$) and significant differences between control and migration treatments for non-predators (Bray-Curtis $R=0.478$; $p=0.0172$ & Jaccard $R=0.332$, $p=0.0523$). The same analysis was conducted using Jaccard distance and is reported in the results section. C1-C5 refer to the control patches while M1-M5 refer to the migration patches.
Table Legend

**Table 1**: Shows the percentage of rare species in the predator and non-predator communities for the control and migration treatment of the experiment. Two definitions of “rare” are used here: first, a morpho-species as a singleton in the dataset (top-half) and second, a given morpho-species as less abundant than the mean abundance of all morpho-species in the community (“relatively rare”).

<table>
<thead>
<tr>
<th>Predator Community</th>
<th>%</th>
<th>Non-Predator Community</th>
<th>%</th>
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<tr>
<td><strong>Singletons</strong></td>
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<td><strong>Singletons</strong></td>
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<tr>
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<td>Control</td>
<td>49</td>
</tr>
<tr>
<td>Migration</td>
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<td>Migration</td>
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<td></td>
<td><strong>Relatively Rare</strong></td>
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<td>Control</td>
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