Assembly-History Dynamics of a Pitcher-Plant Protozoan Community in Experimental Microcosms

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Abstract

Background: History drives community assembly through differences both in density (density effects) and in the sequence in which species arrive (sequence effects). Density effects arise from predictable population dynamics, which are free of history, but sequence effects are due to a density-free mechanism, arising solely from the order and timing of immigration events. Few studies have determined how components of immigration history (timing, number of individuals, frequency) alter local dynamics to determine community assembly, beyond addressing when immigration history produces historically contingent assembly.

Methods/Findings: We varied density and sequence effects independently in a two-way factorial design to follow community assembly in a three-species aquatic protozoan community. A superior competitor, Colpoda steinii, mediated alternative community states; early arrival or high introduction density allowed this species to outcompete or suppress the other competitors (Ptociorochromonas malhamensis and Eimeriidae gen. sp.). Multivariate analysis showed that density effects caused greater variation in community states, whereas sequence effects altered the mean community composition.

Conclusions: A significant interaction between density and sequence effects suggests that we should refine our understanding of priority effects. These results highlight a practical need to understand not only the “ingredients” (species) in ecological communities but their “recipes” as well.

Introduction

Community assembly has been a prominent concept in ecology; a variety of sometimes divergent views have reflected different assumptions and a confusing array of terminology. At one extreme, communities have been viewed as the product of random dispersal events, after which deterministic species sorting overrides immigration history. For example, Diamond [1] outlined a set of “assembly rules” of limited membership for the local fauna of bird communities in New Guinea that set limits on which species from the regional source pool could coexist. At the other extreme, the final community structure can be viewed as a historical artifact of the precise order of species’ arrival. Although not supporting such an extreme role for historical contingency, Drake [2], used aquatic microcosms to show that community assembly depends on potentially complex ways on the identities and sequence of arrival of species as communities develop.

Empirical efforts to understand historical forces driving community assembly have included observational comparisons of natural communities at different localities at various disturbance levels (see, e.g., Urban [3], Weslien et al. [4]) and experimental perturbations of naturally recovering communities [5–7]; these empirical studies complement theoretical investigations into alternative stable states (e.g., Shurin et al. [8]) and transient states [9]. Communities from a wide range of habitats have been shown to be affected by the direct manipulation of immigration history (e.g., acacia ants, by Palmer et al. [10]; amphibians by Wilbur and Allord [11]; aquatic protists by Robinson and Dickerson [12] and Fukami [13]; ectomycorrhizal fungi by Kennedy et al. [14]; drosophilids by Shorrocks and Bingley [15]; wood-decaying fungi by Fukami et al. [16]). Along with empirical insights, theoretical work suggests that the context in which communities assemble can be altered by regional factors (e.g., large regional species pools, low rates of connectivity) and local factors (e.g., high productivity and low disturbance) [17]. These studies have explored aspects of the effects of immigration history on historically contingent assembly, but do not separate how various components of immigration history (timming, number of individuals, frequency) alter local dynamics to determine community assembly. No empirical studies have rigorously identified mechanisms by which local dynamics interact with immigration history.

History drives community assembly by two potentially independent mechanisms, density effects and sequence effects. Density effects are predictable dynamics that follow directly from different initial abundances of competitors and the time for unimpeded growth between colonizing events. For example, simple Lotka-Volterra models predict that, when conditions for a stable two-species equilibrium occur, communities will reach the same final equilibrium state regardless of the initial abundances of species,
but when parameters create an unstable equilibrium, differences in
species’ abundances at the time when later colonists arrive
determine which species outcompetes the other [18]. Density
effects are independent of the history of other species and are
firmly anchored in population-dynamics principles.

In contrast, sequence effects occur through differences that are
unrelated to density but are due purely to the order in which
species arrive. Possible mechanisms of sequence effects would
include delayed life-history effects [19] and ecosystem engineering
that alters fitness landscapes of competing species [20]. Note that,
by our definitions, the widely used term “priority effects” (sensu
Wilbur and Alford [11], Young et al. [21]) confounds density and
sequence effects, even though theory gives reason to suspect that
density and sequence effects on community assembly can differ (cf.
Lotka [18], Connell and Slatyer [22]). Our separation of density
and sequence effects is therefore essentially a claim that we should
refine interpretations of priority effects. Previous experimental
studies (e.g., Drake [2], Fukami [9], Robinson and Dickerson [12],
Kennedy et al. [14], Collinge and Ray [23]) have shuffled the
sequence of species introduction, but because they did not
factorially vary the intensity of immigration (density of species)
crossed with sequence of arrival, the underlying mechanisms
leading to historically contingent community structure remain
undetermined.

We varied density and sequence effects independently in a two-
way factorial design to follow community assembly of an inquiline
protozoan community in experimental microcosms. The commu-
nity originates from the water-filled leaves of the purple pitcher
plant, Sarracenia purpurea; in this ecosystem, energy is derived from
allochthonous material in the form of insects that fall into the
water-filled leaves and drown [24]. Bacteria make up the bottom
trophic level as communities develop through immigration of
protozoans, rotifers, and top predators [25]. This well-studied
community has rapid dynamics and is ideal for studying assembly.
We specifically tested the hypothesis that density and sequence
effects interact to determine the mean and the variation (i.e., beta
diversity) in community structure of protozoans in experimental
microcosms.

**Materials and Methods**

**Ethics statement**

No specific permits were required for the described field and
laboratory studies, as protozoa and bacteria sampling is freely
allowed in natural areas in the Apalachicola National Forest. None
of our studies involved endangered or protected species.

**Study organisms**

Poterioochromonas malhamensis (species A), Colpoda steinii (species B),
and Eimeriidae gen. sp. (species C) are protozoans commonly
found in water-filled leaves of the purple pitcher plant, Sarracenia
purpurea, in the Apalachicola National Forest, Florida, USA. The
first is a suspension feeder; the latter two use both suspension
feeding and grazing [26]. All three are generalist bacterivores and
compete with each other for shared food in pitcher leaves and in
experimental microcosms. We make the simplifying assumption
that bacterial population dynamics are fast enough relative to
protozoan dynamics that one can, without too much error, model

![Figure 1. Experimental design for microcosms of the three protozoan species.](https://example.com/Figure1.png)

![Figure 2. Single-species population growth curves of the three protozoan species used in the community-assembly experiment.](https://example.com/Figure2.png)
the consumer-resource dynamics in a reduced phase space that
considers only consumer protozoans ("ecological abstraction"
"sensu Schaffer [27]).

Experiment design and procedure

We assembled three-species protozoan communities in 10-mL
microcosms, created as described below, by adding 0.50-mL
aliquots from appropriate stock monocultures sequentially in a
two-way factorial design (3 density levels x 6 sequence levels),
replicated in three temporal blocks, each of which contained 18
microcosms. Successive blocks were started at 1-hour intervals (i.e.,
54 microcosms were started over 3 hours). Sequence treatment
included six categorical levels, corresponding to all possible orders
for introduction of the three species (Fig. 1). For each density
treatment, we produced three scenarios, in which introduction
density was negatively correlated, positively correlated, and
uncorrelated with the order of introduction (Fig. 1). A negative
correlation would result in stronger density effects of early-arriving
species on later-arriving species, whereas a positive correlation
would be expected to produce weaker density effects of early
species on later species. Lack of correlation was intended as a
control. A preliminary study showed that single-species popula-
tions of the three species readily attain quasi-steady states within
72 hours under the same immigration density setting (Fig. 2) and
allowed us to estimate microcosm equilibrium carrying capacities
for each species. Differences in maximum population density
(carrying capacity) among the three species possibly reflect
differences in body size. The immigration densities and times of
introduction were therefore determined so that initial high-
intermediate-, and low-density introductions of the first colonizer
result in its having 100, 50, and 25% of equilibrium density at the
time of the introduction of the third species.

Microcosms initially contained 10 mL (within the range of
natural leaf pitcher volumes) of sterile water that was inoculated
with bacteria by addition of 1.6 mg of Tetramin fish food (Tetra
Werke, Germany) and exposure to open air for 24 hours before
the introduction of protozoan species. The first, second, and third
species were added from stock monocultures after 0, 12, and
24 hours, respectively. Differences in introduction density were
created by dilution of high-density stock cultures with medium
similarly inoculated with bacteria; Table 1 gives the densities of the
three protozoan species in the 0.50-mL aliquots used for
introduction. After the initial 24 hours, 0.10-mL subsamples were
taken 12, 24, 48, and 72 hours after establishment of each
microcosm in order to monitor the subsequent abundances of the
three species. We used a phase-contrast microscope at 100x
to census all or a part of each subsample by means of a Palmer
counting cell; all counts were converted to cells per 0.10 mL.
During the census period, microcosms were provided with food
semincontinuously by addition of 0.55 mg of Tetramin every
6 hours.

| Table 1. Abundances of the three protozoan species in 0.50-mL aliquots of stock cultures (used for introduction of protozoans
into the microcosms) in medium inoculated with bacteria. |
|---------------------------------|-----------------|----------------|
| High density | Poterioochromonas malhamensis (species A) | 5000 | 1600 | 300 |
| Intermediate density | Colpoda steinii (species B) | 100 | 80 | 50 |
| Low density | Eimeriidae gen. sp. (species C) | 10 | 10 | 10 |

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Figure 3. Population dynamics of the three protozoan species over 72 hours in the density (negative, control, and positive
treatments; see Fig. 1) and sequence-of-introduction treatments. Densities were log_e (x+1)-transformed and then scaled to log-transformed
carrying capacity for each species, and three temporal blocks were averaged. Circles, species A; triangles, species B; crosses, species C. According to
Fig. 2, we used 11000/mL for species A, 2500 for species B, and 550 for species C as estimates of carrying capacity for illustrative purpose.

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Statistical analysis

Community-wide analysis. We used permutational MANOVA (PERMANOVA) to test for density and sequence effects on the final community structure of the three protozoan species in microcosms on day 3 (72 hours after microcosm establishment). PERMANOVA uses an additive partitioning of a pairwise distance metric (e.g., Bray-Curtis index) according to a multifactorial ANOVA design, with significance testing by permutation to accommodate the frequent violation of the assumptions of MANOVA in community data [28]. p-values were obtained from separate sets of 999 permutations that were performed across only the pair of groups being compared, and Bonferroni correction adjusted experimentwise error rates for multiple comparison of the arrival density treatments.

Although PERMANOVA can test for differences in the centroids of multivariate species composition between levels of a factor, we were also interested in differences in the among-plot variability in species composition (i.e., beta diversity, Anderson et al. [29]) across different levels of a factor. We performed an analysis of multivariate homogeneity of group dispersions with the nonparametric PERMDISP [30], then calculated the averaged Bray-Curtis distance of group members to the group centroid within density and sequence treatments. To examine the significance of specific treatment combinations, we used Tukey’s HSD.

Pairwise a posteriori analysis. We tested for the effects of the identities of first-arriving and late-arriving species, density effects, and the interaction between first-arriving/late-arriving effect and density effects, using appropriate subsets of the data for pairwise a posteriori PERMANOVA. For example, the difference between the first-arrival effects of species A and B was based on a subset of sequence treatments in which either species A or species B arrived first (i.e., the first, second, fourth, and fifth columns of Fig. 1). Statistical analysis was performed in R 2.13 [31] with the community ecology package vegan [32].

Results

Direct manipulation of density and sequence in the initial 24 hours resulted in large variation in the subsequent protozoan community structure (Fig. 3). For example, in the treatment where the order of invasion was species C, then A, then last B (hereafter CAB) and the ACB treatment, crossed with positive density effects, species B was suppressed to an extremely low density or went extinct. Species A dominated numerically in 83% of the treatments but not in control BC or positive-density treatments crossed with BCA and BAC. Replicate communities displayed strong consistency among temporal blocks (PERMANOVA, Table 2).
Community-wide effects of density and sequence

Density treatments differed significantly in average community states (PERMANOVA, $F = 11.86$, df = 2, $p = 0.001$), as did sequence treatments ($F = 11.28$, df = 5, $p = 0.001$), and notably, these two effects interacted to determine the protozoan community structure (PERMANOVA, $F = 5.986$, df = 10, $p = 0.001$). Strong density effects also caused greater dispersion in the protozoan community structure (PERMDISP, Tukey's HSD, $p < 0.001$; Table 5). When either species A or species C was the first colonist (i.e., in pairwise comparisons, species B compared to A and species B compared to C), density and sequence effects interacted to determine the final community structure (Table 4). Only the main effects of density and sequence were significant when either species B or species C was the late-arriving species, no main effects of density or sequence were detected, but the density-by-sequence interaction was significant (PERMANOVA, $F = 3.365$, df = 2, $p = 0.007$; Table 3). When either species C or species A was the late-arriving species, density effects primarily controlled the community structure (PERMANOVA, $F = 5.578$, df = 2, $p = 0.003$; Table 5).

In summary, whichever species arrived first had an advantage and more strongly influenced later-colonizing competitors, regardless of its initial density. A superior competitor, species B, mediated alternative community states; when arriving first, it outcompeted or suppressed the other two species, whereas it was vulnerable to extinction under the simultaneous disadvantages of late arrival and low density. In contrast, species A and C were able to overcome any disadvantage due to low initial density.

**Discussion**

Our experiment and analyses demonstrated that density and sequence effects were distinct ecological mechanisms that differed qualitatively in their impacts on assembly of a three-species protozoan community and, most importantly, that density and sequence effects on assembly interacted. Density effects caused greater dispersion in the protozoan community structure without...
substantially changing the average community states, whereas sequence effects often altered the community states, possibly through changing the locations of the community attractors themselves (Fig. 4). Historical contingency in the protozoan community therefore arises from three sources: (1) whether or not initial densities differ sufficiently to cause density effects when the immigration sequence and times of arrival are fixed, (2) whether sequence effects determine community structure even when initial densities do not differ substantially, and (3) effects of assembly history that arise from interactions between density and sequence effects.

Although most of our experimental protozoan communities appeared to have stabilized after 72 hours (approximately 9 generations), we cannot be certain that they do not actually represent transient community states. We suggest that the durations of our experiments were generally sufficient, on the basis of the criteria presented by Grover and Lawton [33]—the intervals between invasions was longer than the generation times, the invasion interval was shorter than the time necessary for maximum population densities to develop, and the total duration of the experiment was much longer than the generation times. However, even if our final communities did not represent stable or near-stable states, theoretical work suggests that historical contingency can be important for understanding transient dynamics as well [9].

In prior studies, priority effects have often been invoked as a post hoc explanation for the observed community changes, as the investigator looked back to initial conditions to interpret current conditions [see, e.g., Robinson and Dickerson [12], Kennedy et al. [14]]. However, because early-arriving species in these studies are likely to be more abundant by the time later species arrive, the effects of sequence and density are confounded. Our work not only suggests a refined interpretation of priority effects in principle but also provides a wider framework that might be useful in decision making in practical restoration projects. For restoration ecologists, the vague concept of priority effects does not reveal when, or how many individuals of, a particular species should be introduced, because most previous studies of priority effects inherently confounded density and timing. We propose that a theory of assembly history could better guide restoration efforts if density and timing are considered separately and interactively.

Sequence effects may be characteristic of particular types of systems—they may lead us forcefully to dissect purely historical processes into trait-based mechanisms (see, e.g., Beckerman et al. [19]) for the practical purpose of gaining specific predictions about the target systems. Separating density and sequence effects can thus contribute theoretical guidance to harnessing contingency behind community assembly or at least clarifying the information demands in previous studies that have relied heavily on purely empirical, case-by-case approaches. Sequence effects may less important in other systems, however, especially over the long term: for example, Collinge and Ray [23] used a restoration project in vernal plant communities to test for historically contingent assembly but found that the order and intensity of seeding influenced plant communities only transiently, within a decade of early community formation.

An important future challenge will be to determine whether such historical forces scale up to more complex situations. Natural experiments often involve many uncontrolled variables and may require using multiple sources of information to rule out alternative hypotheses of assembly-history dynamics. Reconstructing population-genetic structure by analyzing current populations, for example, may allow us to use proxies for density effects and sequence effects of the unwitnessed past. Accumulating quantitave facts about the components of immigration history (timing, number, frequency, etc.) in island restoration, biocontrol management, and biological invasion continues to be important for understanding a large-scale imprint of assembly-history dynamics. Although our study was of a competitive community, further mechanistic lines of inquiry into assembly-history dynamics for predator-prey interactions, mutualisms, and multi-trophic food webs will enrich our understanding not only of the ingredients (the species) but also of the recipes (timing and numbers of individuals) for ecological communities in an invasion-driven world.

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Author Contributions
Conceived and designed the experiments: KK BDI TEM. Performed the experiments: KK. Analyzed the data: KK. Wrote the paper: KK BDI TEM.

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