Detecting spatial ontogenetic niche shifts in complex dendritic ecological networks

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Abstract. Ontogenetic niche shifts (ONS) are important drivers of population and community dynamics, but they can be difficult to identify for species with prolonged larval or juvenile stages, or for species that inhabit continuous habitats. Most studies of ONS focus on single transitions among discrete habitat patches at local scales. However, for species with long larval or juvenile periods, affinity for particular locations within connected habitat networks may differ among cohorts. The resulting spatial patterns of distribution can result from a combination of landscape-scale habitat structure, position of a habitat patch within a network, and local habitat characteristics—all of which may interact and change as individuals grow. We estimated such spatial ONS for spring salamanders (Gyrinophilus porphyriticus), which have a larval period that can last 4 years or more. Using mixture models to identify larval cohorts from size frequency data, we fit occupancy models for each age class using two measures of the branching structure of stream networks and three measures of stream network position. Larval salamander cohorts showed different preferences for the position of a site within the stream network, and the strength of these responses depended on the basin-wide spatial structure of the stream network. The isolation of a site had a stronger effect on occupancy in watersheds with more isolated headwater streams, while the catchment area, which is associated with gradients in stream habitat, had a stronger effect on occupancy in watersheds with more paired headwater streams. Our results show that considering the spatial structure of habitat networks can provide new insights on ONS in long-lived species.

Key words: centrality; Gyrinophilus porphyriticus; network topology; occupancy; salamander; species distribution.

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INTRODUCTION

Life history responses to environmental variation include changes in niche position as individuals mature, and these ontogenetic niche shifts (ONS) contribute to regulation of population growth and community structure (Werner and Gilliam 1984, Claessen and Dieckmann 2002, Nakazawa 2014, van Leeuwen et al. 2014), especially for species with complex life cycles or delayed maturity. For these species, consequences of habitat choice in early life stages may influence population growth rates by allowing access to differentially available resources (e.g., Mittelbach et al. 1988) or via differential sensitivity of life stages (e.g., Crouse et al. 1987, Rudolf and Lafferty 2011, Regnier et al. 2012, Burton and Metcalfe 2014). Resource levels within different habitats may depend on the position of sites within the landscape (Turner et al. 1997, Laurance 2008), complicating the assessment of ONS at small spatial scales. Most studies of ONS focus on changes in
diet or habitat use during a single transition within
the lifespan of an organism, such as changes asso-
ciated with maturation or achieving a size thresh-
old (Graham et al. 2007, Ayllón et al. 2010). Some
studies have included variation in local resources
among sites, but they generally do not describe
how different cohorts vary in their response to
broader patterns of habitat structure and isolation
within a landscape. Specifically, although studies
often compare resource levels associated with
ONS, how landscape structure mediates variation
in resources is poorly understood. A new focus on
spatial ONS, which accounts for the position of a
habitat patch within landscapes as well as the geo-
metric configuration of landscapes, can provide
new insights into the mechanisms affecting popu-
lation distribution and persistence.

The geometry of stream ecosystems drives
resource distributions (Benda et al. 2004), results in
predictable gradients of habitat isolation, and con-
strains movement for aquatic-obligate species,
making these good systems for studying spatial
ONS. Unlike terrestrial environments, streams are
formed by geomorphic processes that result in pre-
dictable gradients in habitat and resource distribu-
tions (Vannote et al. 1980, Rodriguez-Iturbe and
Rinaldo 2001, Rosi-Marshall and Wallace 2002,
Raymond et al. 2016). For example, resource distri-
butions may change at confluences, and species
that inhabit small streams may respond to this spa-
tial variation in resources by moving to areas with
higher resource levels (Uno and Power 2015). The
ability to take advantage of these spatially dis-
tributed resources depends on the degree to which
species are restricted to moving within or along
stream channels (Vannote et al. 1980, Heino et al.
2005). Further, the overall structure of the land-
scape changes the relative importance of isolation
(Andren 1994, Carrara et al. 2012). As individuals
age and require different resources for survival,
growth, or reproduction, their ability to access
complementary habitats can be strongly influenced
by the stream network geometry. Therefore, con-
sidering both the structure of stream networks and
the isolation of habitat within networks is im-
portant to describe spatial ONS.

We test for spatial ONS by characterizing shifts
in distributions of different cohorts of aquatic-
obligate larval spring salamanders (*Gyrinophilus
porphyriticus*) in headwater stream networks.
Spring salamanders have a prolonged larval period
that can last 4 years or more (Bruce and Castanet
2006, Resetarits 1995), have continuous growth,
and multiple age classes may be present in a given
stream segment. We identify age cohorts based on
size frequency data of captured individuals, and
use the occurrence patterns of each age class to
test two predictions regarding spatial ONS in
spring salamanders. First, we predict that occu-
pancy patterns will differ among cohorts and be
related to measures of network geometry. Second,
we predict that any response to stream network
position will be mediated by the overall structure
of the network. The role of space in population
dynamics has long been appreciated; the study of
ONS can also benefit from new approaches that
account for the isolation of resources in land-
scrapes.

**METHODS**

*Field surveys*

To detect spatial ONS, we sampled 59 stream
segments across 10 randomly selected watersheds
within the Shenandoah National Park, Virginia,
United States. The park ranges in elevation from
400 to 1200 m above sea level along a north–south
ridge that runs over 150 km, and is primarily cov-
ered by hardwood forest. We selected five water-
sheds with outlets at the park boundary that were
drained by at least a second-order stream. We also
sampled an adjacent watershed on the opposite
side of the main ridge that divides the park to
control for any effect of aspect on *Gyrinophilus por-
phyriticus* distribution.

Stream segments were sampled for spring sala-
mander larvae during June and July 2012. We
sampled each 20-m stream segment with three
consecutive passes, and individual larvae were
placed into a plastic bag at the time of their cap-
ture. Stream segments were located 10 m and
150 m upstream of confluences on the main chan-
nel and tributary, and they contained a range of
stream habitats (e.g., riffles, runs, pools). Larval
salamanders were measured to the nearest mil-
liter (snout-vent and total lengths, Heyer 1994)
before being released near their point of capture.

We quantified in-stream habitat at the time of
surveys by measuring stream width and stream
depth at three random locations along the sam-
pling transect. We also estimated percent of each
transect covered by silt, sand, pebble, gravel,
cobble, and boulder in the streambed (Hauer and Lamberti 2007). We log-transformed stream width and stream depth and standardized variables prior to conducting a principal components analysis to describe major gradients in stream habitat.

**Characterizing stream network topology**

To characterize spatial ONS of larval salamanders, we calculated three metrics that describe the geometry of stream networks (i.e., network topology). First, the catchment area of a stream segment (the total area drained by a stream segment) is correlated with stream size and associated gradients in stream habitat (Vannote et al. 1980). We also derived two centrality metrics that characterize the position of stream segments within a network relative to potential dispersal behavior of animals (Estrada and Bodin 2008). First, betweenness centrality reflects the degree to which a location acts as a central connection for pathways through a network. Betweenness centrality quantifies the proportion of pathways between all the sites in a network that pass through a given site. We hypothesized that betweenness could be related to spatial ONS because downstream movements would put individuals in areas with higher betweenness centrality, which may be advantageous for exploration of the network (Estrada and Bodin 2008, Altermatt 2013). Second, farness centrality reflects the average distance from one stream segment to all other stream segments within the network (Altermatt 2013). This metric may characterize spatial ONS because downstream movements out of first-order streams would see individuals moving into stream segments with lower farness centrality. Farness centrality can be calculated with weighted distances to allow for upstream or downstream bias in movement, and we calculated it by weighting downstream movements twice as costly as upstream movements based on a prior study of dispersal behavior of this species (Lowe 2002). Both centrality metrics were calculated after deriving stream networks from a 10-m resolution digital elevation model (www.nationalmap.gov; accessed 2 March 2015), splitting networks into 20-m stream segments, and normalizing values by dividing centrality by the total number of stream segments within a network. We used the r.watershed, r.stream.stats, and v.net.centrality functions in GRASS GIS for these analyses (version 7.0, GRASS Development Team 2012).

To allow for an effect of the overall watershed structure, we also calculated two metrics of stream branching structure at the level of the entire network (Dodds and Rothman 1999): Horton’s length ratio (or the ratio of the length of tributaries to main channels, $R_T$), and the average number of major tributaries within a stream network ($T_1$), which is defined as being one Strahler order less than the main channel (e.g., where major tributaries are defined as being first-order streams that flow into a second-order stream; Fagan et al. 2010). These metrics are calculated based on all streams within a watershed, but they are appropriate for characterizing aspects of first-order streams because of the fractal nature of rivers (Rodriguez-Iturbe and Rinaldo 2001). For species that inhabit first-order streams, lower $R_T$ and higher $T_1$ reflect more paired first-order streams, which are associated with higher probabilities of occupancy and abundance for headwater-associated species (Lowe and Bolger 2002, Grant et al. 2009).

**Niche modeling**

Because the sampled population contains individuals of multiple age classes, we fit mixtures of normal distributions to size frequency data from individuals captured in occupancy surveys to identify age cohorts (MacDonald and Pitcher 1979). We verified the size distribution data using annual growth rate estimates from mark-recapture data (Appendix S2). The presence of each age class was related to covariates using occupancy models (MacKenzie et al. 2006), fit with JAGS (http://mcmc-jags.sourceforge.net/) using the jagsUI library in R (Kellner 2015, R Core Team 2015). These models account explicitly for the probability of detecting an individual of a given age class, given it is present during a sampling occasion, and thus provide unbiased estimation of occupancy and the relationship between occupancy and our covariates (Royle and Dorazio 2008). We fit models for each age class separately, and related occupancy to $R_T$, $T_1$, betweenness centrality, farness centrality, catchment area, and the first three principal components of stream habitat (Appendix S1), which accounted for 76% of the variation in stream habitat. We also included the first three principal components of stream habitat as covariates for detection probability because we believed larval salamanders were more difficult to capture in certain microhabitats (Crocker et al. 2007).
We visually inspected trace plots for evidence of good mixing among chains, and we examined the $R$ statistic to evaluate model convergence (Kery and Schaub 2012). We ran three chains for 100,000 iterations with a burn-in period of 20,000 iterations, and we did not thin the results (Eaton and Link 2011). We evaluated covariate effects on occupancy by examining credible intervals of posterior distributions. More precise estimates have narrower credible intervals. We considered effects to be strong when 95% of the mass of a posterior distribution does not include 0.

RESULTS

Mixture models identified four age classes of larvae present across our 59 stream networks. Growth rates estimated from mark–recapture data for this species (Appendix S2) confirmed that these mixture models provided an accurate representation of the age cohorts within the population. The probability of detecting each age class during one sampling pass varied: $P_{class1} = 0.05$ (95% credible interval [0.02–0.15]); $P_{class2} = 0.13$ (95% credible interval [0.06–0.27]); $P_{class3} = 0.37$ (95% credible interval [0.30–0.45]); $P_{class4} = 0.34$ (95% credible interval [0.25–0.50]). In general, principal components of stream habitat (Appendix S1) were weakly related to detection probability. The only exception was the second principal component, which was associated with shallow, narrow stream reaches and which had a negative effect on the detection probability of age 2 larvae ($-0.95$, 95% credible interval $[-1.65, -0.32]$).

Age classes varied in their response to stream habitat (Fig. 1). Age classes 1 and 3 did not show a strong response to the first principal component of stream habitat (58.2% and 18.1% of posterior mass <0, respectively), but age classes 2 and 4 were less likely to occur in areas with higher amounts of cobble (99.4% and 95.9% of posterior mass <0, respectively). Age classes 2, 3, and 4 were more likely to occur in shallow, narrow reaches (99.9%, 73.4%, and 95.5% of posterior mass >0), but age class 1 was less likely to occur in this habitat (17.4% of posterior mass >0). Age classes 1 and 3 did not show a strong response to the third principal component, which was associated with reaches dominated by gravel and pebble streambeds (25.7% and 59.3% of posterior mass >0, respectively), while age class 4 was more likely to occur in this habitat (97.8% of posterior mass >0) and age class 2 was less likely to occur in this habitat (0.3% of posterior mass >0). Results for age classes 1 and 2 should be interpreted with caution because of the low numbers of individuals that were captured from these age classes. Age classes varied in their response to stream network structure and within-network position. Catchment area had a weak negative relationship with occupancy of age classes 3 and 4 (80.0% of posterior mass <0 for class 3, 92.5% of posterior mass <0 for class 4). Betweenness centrality was unrelated to

![Fig. 1. Parameter estimates and 95% credible intervals for parameters predicting occupancy of four age classes of larval spring salamanders (Gyrinophilus porphyriticus) in the Shenandoah National Park, Virginia, United States (PC1: first principal component of stream habitat; PC2: second principal component of stream habitat; PC3: third principal component of stream habitat; BC: betweenness centrality; FC: farness centrality; CA: catchment area; RT: Horton's length ratio; T1: average number of major tributaries). Panels display results for age classes 1–4 (top to bottom).](image-url)
the presence of age 3 individuals, but positively associated with the presence of age class 4 (58.2% of posterior mass >0 for age class 3, 90.0% of posterior mass >0 for age class 4), while farness centrality showed a positive relationship with the presence of age classes 3 and 4 (93.4% of posterior mass >0 for age class 3, 98.6% of posterior mass >0 for age class 4). RT was negatively associated with the occupancy of both age class 3 (5.3% of posterior mass >0) and age class 4 (92.6% of posterior mass <0), but T1 did not have a strong relationship with either age class 3 (71.6% of posterior mass <0) or age class 4 (67.2% of posterior mass <0).

We created two sets of maps to illustrate the variation in patterns of occupancy for different age classes. First, we mapped the predicted occupancy for each age class based on models with average stream habitat conditions in the watersheds that had the highest and lowest RT (Fig. 2). Age classes 3 and 4 had similar occupancy patterns in a watershed with the lowest RT, but age class 4 was less likely than age class 3 to occur in tributaries near the outlet of the watershed that had the highest RT.

Second, we compared maps of the predicted occupancy for age class 4 in these two watersheds with maps of predicted occupancy that included one measure of network position and both measures of network structure, while fixing the other two measures of network position and measures of stream habitat at their average values (Fig. 3). This was done because the relative isolation of a site within a habitat network might be less important in a landscape where paired headwaters are more common. When a variable has a strong effect on occupancy patterns, a map showing the predicted effect of that variable with other predictors held at average conditions will resemble the map produced with all terms within a model, which is similar to calculating partial correlation coefficients for parameters of a regression model. The predicted pattern of occupancy from the full model was most closely matched by the effect of catchment area in the watershed with the lowest RT and the effect of farness centrality in the watershed with the highest RT.

**DISCUSSION**

Habitat choice during early life stages has implications for overall population dynamics and trends, though detecting these preferred niches using static distribution data may be difficult for species with overlapping generations, long larval life stages, and for species living in continuous habitat networks. Theoretical models for ONS often focus on the role of body size and transitions between different habitat types (Nakazawa 2014), but network structure may place constraints on these optimal ONS by restricting movement among habitat types (Carrara et al. 2014, Bertuzzo et al. 2015). Although the spatial scale at which populations respond to habitat has been examined for some species (Ayllón et al. 2010), the importance of the topological relationships among patches within habitat networks remains poorly understood. We used size frequency data to identify age classes in populations of the stream-associated spring salamander, which allowed us to identify spatial ONS related to local- and landscape-scale factors. Our findings illustrate that ONS can be driven by habitat structure at large spatial scales.

The location of sites within stream networks was an important predictor of ONS in larval spring salamanders. While betweenness centrality and farness centrality were more strongly associated with the presence of the fourth age class than with that of the third age class, catchment area had a consistent negative effect on the presence of these older ages. When predicted occupancy is mapped across networks, it is apparent that the branching structure of stream networks mediates the effect of the position of stream segments within networks (Fig. 3). In a watershed with more paired headwaters (Big Run, Fig. 3), occupancy patterns are well represented by the response to catchment area, but in a watershed with less complex topology (Jeremy’s Run, Fig. 3), farness centrality reflects the variation in predicted occupancy. This indicates that the isolation of stream segments matters more in a watershed that contains more isolated first-order streams. While a previous study of stream fish found that extinction risk declined as Horton’s length ratio, RT (the ratio of the length of tributaries to main channels) increased (Fagan et al. 2010), we found lower occupancy rates for stream salamander larvae in watersheds with higher RT. This difference may result from different preferences for habitat position (e.g., main channel vs. tributary) among species, suggesting that habitat isolation is more important for headwater species. In watersheds with higher RT, first-order streams...
are more isolated from one another, while larger-order streams are always adjacent to one another. This difference is compounded by the upstream bias in movement of many species that inhabit first-order streams (Grant et al. 2007). All age classes of spring salamanders have an upstream bias in movement, especially the adults (Lowe 2003), so selection for network positions with higher centrality allows for better access to headwater habitats, which are the main sites for reproduction. Occupancy site selection of younger age classes may indicate that those positions in the

Fig. 2. Predicted occupancy for age classes 1–4 (top to bottom) of larval spring salamanders (*Gyrinophilus porphyriticus*) in Big Run (panels a–d, lowest Horton’s length ratio) and Jeremy’s Run (panels e–h, highest Horton’s length ratio).
stream network are better for survival and growth (Cecala et al. 2009).

Our modeling approach shows that larval salamanders choose sites based on local habitat after accounting for network topology. Both the third and fourth age classes were associated with shallower, narrower reaches. Although the third age class was not associated with reaches having

Fig. 3. Predicted occupancy for age class 4 of larval spring salamanders (*Gyrinophilus porphyriticus*) in Big Run (panels a–d, lowest Horton’s length ratio) and Jeremy’s Run (panels e–h, highest Horton’s length ratio) from the full model (top) and predicted occupancy (ψ) when considering only network structure and catchment area (second row), network structure and farness centrality (third row), and network structure and betweenness centrality (fourth row).
more cobble, gravel, or pebble substrates, the fourth age class was positively associated with these conditions. Indeed, we find that habitat characteristics of stream segments were unrelated to their position within stream networks (Appendix S1). The vegetation structure of terrestrial habitats may be affected by patch area and patch isolation via factors like edge effects or seed dispersal limitation (Roderwald 2003, Laurance 2008). In contrast to this, stream habitats are often the result of stream flow and other fluvial processes, so the range of habitats associated with smaller streams can be similar in watersheds that vary in their size or the relative isolation of headwater streams (Benda et al. 2004).

Understanding spatial ONS improves our ability to identify intraspecific variation in distribution, can elucidate cryptic spatial dynamics in populations with multiple life history stages, and presents new opportunities for empirical tests of ecological theory. The complexity of a habitat network is related to metapopulation capacity (Bertuzzo et al. 2015) and persistence (Grant 2011) in theoretical landscapes. In species with prolonged larval or juvenile life stages, spatial ONS may further enhance metapopulation capacity and persistence by maximizing survival in complementary locations in the habitat network. Our results show that larval salamanders appear to have distributions that shift upstream as they age, which is likely to put them in areas with fewer predators and lower resource levels.

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LITERATURE CITED

Altermatt, F. 2013. Diversity in riverine metacommunities: a network perspective. Aquatic Ecology 47:365–377.
Andren, H. 1994. Effects of habitat fragmentation on birds and mammals in landscapes with different proportions of suitable habitat: a review. Oikos 71:355–366.
Ayllón, D., A. Almodovar, G. G. Nicola, and B. Elvira. 2010. Ontogenetic and spatial variations in brown trout habitat selection. Ecology of Freshwater Fish 19:420–432.
Benda, L., N. L. Poff, D. Miller, T. Dunne, G. Reeves, G. Pess, and M. Pollock. 2004. The network dynamics hypothesis: How channel networks structure riverine habitats. BioScience 54:413–427.
Bertuzzo, E., I. Rodriguez-Iturbe, and A. Rinaldo. 2015. Metapopulation capacity of evolving fluvial landscapes. Water Resources Research 51:2696–2706.
Burton, T., and N. B. Metalffe. 2014. Can environmental conditions experienced in early life influence future generations? Proceedings of the Royal Society B 281:20140311.
Carrara, F., F. Altermatt, I. Rodriguez-Iturbe, and A. Rinaldo. 2012. Dendritic connectivity controls biodiversity patterns in experimental metacommunities. Proceedings of the National Academy of Sciences USA 109:5761–5766.
Carrara, F., A. Rinaldo, A. Giometto, and F. Altermatt. 2014. Complex interaction of dendritic connectivity and hierarchical patch size on biodiversity in river-like landscapes. American Naturalist 183:13–25.
Cecala, K. K., S. J. Price, and M. E. Dorcas. 2009. Evaluating existing movement hypotheses in linear systems using larval stream salamanders. Canadian Journal of Zoology 87:292–298.
Claessen, D., and U. Dieckmann. 2002. Ontogenetic niche shifts and evolutionary branching in size-structured populations. Evolutionary Ecology Research 4:189–217.
Crocker, J. B., M. S. Bank, C. S. Loftin, and R. E. Jung Brown. 2007. Influence of observers and stream flow on the northern two-lined salamander (Eurycea bislineata bislineata) relative abundance estimates in Acadia and Shenandoah National Parks, USA. Journal of Herpetology 41:323–329.
Crouse, D. T., L. B. Crowder, and H. Caswell. 1987. A stage-based population model for loggerhead sea turtles and implications for conservation. Ecology 68:1412–1423.
Dodds, P. S., and D. H. Rothman. 1999. Unified view of scaling laws for river networks. Physical Review E 59:4865.
Eaton, M. J., and W. A. Link. 2011. Estimating age from recapture data: integrating incremental growth measures with ancillary data to infer age-at-length. Ecological Applications 21:2487–2497.

Estrada, E., and O. Bodin. 2008. Using network centrality measures to manage landscape connectivity. Ecological Applications 18:1810–1825.

Fagan, W. F., E. H. C. Grant, H. Lynch, and P. J. Unmack. 2010. Riverine landscapes: ecology for an alternative geometry. Pages 85–100 in S. Cantrell, C. Cosner, and S. Ruan, editors. Spatial ecology. CRC Press, Chicago, Illinois, USA.

Graham, B. S., D. Grubbs, K. Holland, and B. N. Popp. 2002. Landscape-scale spatial population dynamics in human-impacted stream systems. Environmental Management 30:225–233.

Lowe, W. H. 2003. Linking dispersal to local population dynamics: a case study using a headwater salamander system. Ecology 84:2145–2154.

Lowe, W. H., and D. T. Bolger. 2002. Local and landscape-scale predictors of salamander abundance in New Hampshire headwater streams. Conservation Biology 16:183–193.

MacDonald, P. D. M., and T. J. Pitcher. 1979. Age-groups from size-frequency data: a versatile and efficient method of analyzing distribution mixtures. Journal of the Fisheries Research Board of Canada 36:987–1001.

MacKenzie, D. I., J. D. Nichols, J. A. Royle, K. H. Pollock, L. L. Bailey, and J. E. Hines. 2006. Occupancy estimation and modeling: inferring patterns and dynamics of species occurrence. Academic Press, San Diego, California, USA.

Mittelbach, G. G., C. W. Osenberg, and M. A. Liebold. 1988. Trophic relations and ontogenetic niche shifts in aquatic ecosystems. Pages 219–235 in B. Ebenman and L. Persson, editors. Size-structured populations. Springer, Berlin, Germany.

Moll, J. D., and J. S. Brown. 2008. Competition and coexistence with multiple life-history stages. American Naturalist 171:839–843.

Morrongiello, J. R., and R. E. Thresher. 2014. A statistical framework to explore ontogenetic growth variation among individuals and populations: a marine fish example. Ecological Monographs 85:93–115.

Nakazawa, T. 2014. Ontogenetic niche shifts matter in community ecology: a review and future perspectives. Population Ecology 2014:1–8.

R Core Team. 2015. R: a language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. http://www.R-project.org/
Journal of the American Statistical Association 92:894–902.
Rosi-Marshall, E. J., and J. B. Wallace. 2002. Invertebrate food webs along a stream resource gradient. Freshwater Biology 47:129–141.
Royle, J. A., and R. M. Dorazio. 2008. Hierarchical modeling and inference in ecology: the analysis of data from populations, metapopulations, and communities. Academic Press, San Diego, California, USA.
Rudolf, V. H. W., and K. D. Lafferty. 2011. Stage structure alters how complexity affects stability of ecological networks. Ecological Letters 14:75–79.
Turner, M. G., W. H. Romme, R. H. Gardner, and W. W. Hargrove. 1997. Effects of fire size and pattern on early succession in Yellowstone National Park. Ecological Monographs 67:411–433.
Uno, H., and M. E. Power. 2015. Mainstem-tributary linkages by mayfly migration help sustain salmonids in warming river networks. Ecology Letters 18:1012–1020.
Urban, D. L., E. S. Minor, E. A. Treml, and R. S. Schick. 2009. Graph models of habitat mosaics. Ecology Letters 12:260–273.
van Leeuwen, A., M. Huss, A. Gardmark, and A. M. de Roos. 2014. Ontogenetic specialization in predators with multiple niche shifts prevents predator population recovery and establishment. Ecology 95:2409–2422.
Vannote, R. L., G. W. Minshall, K. W. Cummins, J. R. Sedell, and C. E. Cushing. 1980. The river continuum concept. Canadian Journal of Fisheries and Aquatic Sciences 37:130–137.
Werner, E. E., and J. F. Gilliam. 1984. The ontogenetic niche and species interactions in size-structured populations. Annual Review of Ecology and Systematics 15:393–425.
Whitlock, M. C. 2001. Dispersal and the genetic properties of metapopulations. Pages 273–282 in J. Clobert, E. Danchin, A. A. Dhondt, and J. D. Nichols, editors. Dispersal. Oxford University Press, Oxford, UK.

Supporting Information

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