Foraging selectivity by larval yellow perch (*Perca flavescens*): implications for understanding recruitment in small and large lakes

Richard S. Fulford, James A. Rice, Thomas J. Miller, Fred P. Binkowski, John M. Dettmers, and Brian Belonger

Abstract: Growth and survivorship of larval yellow perch (*Perca flavescens*) have been examined in many systems but can conclusions from well-studied perch populations in smaller lakes be applied to populations in meso-oceanic systems like Lake Michigan, USA? Laboratory experiments were conducted with yellow perch (hatch to 35 mm total length) to develop an empirical selectivity function based on Chesson’s $\alpha$ to describe larval diet as a function of changes in prey community composition. This function was used in an individual-based foraging and growth model (IBM) to describe changes in foraging decisions resulting from changes in prey composition between different systems. Larval perch made three selective transitions during ontogeny. Initial positive selection for rotifers and the relative selectivity for cladocerans vs. copepods in late-stage larvae were both dependent on prey composition. Larvae exposed to prey assemblages differing only in composition had different diets. The empirically based IBM accurately predicted these dietary differences and resulting differences in larval growth and likelihood of starvation between systems at equal prey density. The importance of feeding behavior to larval survival will differ between Lake Michigan and smaller lakes, and these results are important for comparisons of recruitment dynamics between large and small systems.

Résumé : La croissance et la survie des larves de la perchaude (*Perca flavescens*) ont été étudiées dans plusieurs systèmes, mais il reste à savoir si les conclusions tirées de populations bien analysées dans les lacs plus petits sont applicables aux populations de systèmes méso-océaniques, tels que le lac Michigan, É.-U. Nous avons mené des expériences de laboratoire avec des perchaudes (de l’éclosion à 35 mm de longueur totale) afin de mettre au point une fonction de sélection empirique basée sur l’$\alpha$ de Chesson pour décrire le régime alimentaire des larves en fonction des changements dans la communauté de proies. Cette fonction sert dans un modèle de la recherche de nourriture et de la croissance basé sur l’individu (IBM) à décrire les changements dans les décisions de recherche de nourriture resultant de variations de la composition des proies dans les divers systèmes. Les larves de perchaude traversent trois périodes de transition dans leur sélection alimentaire durant leur ontogénie. Une sélection initiale positive pour les rotifères et une sélection relative pour les cladocères par rapport aux copépodes chez les larves avancées sont toutes deux reliées à la composition des proies. Les larves exposées à des ensembles de proies qui diffèrent seulement par leur composition ont des régimes alimentaires différents. Dans des conditions de densité constante des proies, le modèle empirique IBM prédit de façon exacte ces différences de régime, ainsi que les différences qui en résultent dans la croissance larvaire et la probabilité de mourir de faim dans les divers systèmes. L’importance du comportement alimentaire pour la survie des larves diffère dans les petits lacs et dans le lac Michigan et ces résultats sont d’importance pour comparer la dynamique du recrutement dans des systèmes de grande et de petite taille.

Introduction

It is well-known that variability in mortality of larval fishes is often a significant factor driving variability in fishery recruitment (Sharp 1987). Mortality during the larval stage is affected by a variety of factors, including predation (Cowan and Houde 1993; Rice et al. 1993) and feeding success (Cushing 1990), and predicting larval mortality has proven elusive. In particular, interannual variation in survival of larval yellow perch (*Perca flavescens*) has been linked to pre-


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tion (Shroyer and McComish 2000), but the interaction of feeding success, growth, and survival has not been well described, particularly in larger systems.

Feeding success in fish larvae is affected by endogenous factors such as sensory ability (Blaxter 1986) and swimming performance (Webb and Weh's 1986; Fuiman et al. 1999) and by exogenous factors such as prey size and prey community composition (Kerfoot et al. 1980; DeVries et al. 1998). It is likely that larval fishes select prey based on a combination of these factors that is changing rapidly during ontogeny. Descriptions of larval diet based on endogenous factors only or on simplistic, exogenous models based only on prey size may fail to predict how changes in the prey community composition affect selection. Interactions of prey community structure with larval selectivity are particularly relevant for an analysis of feeding success of an opportunistic feeder such as larval yellow perch.

Yellow perch has a pelagic phase lasting 30–40 days that is generally associated with the ontogenetic period from hatch to the acquisition of juvenile characteristics, although the transition to demersal habitat generally occurs slightly later (Auer 1982). Yellow perch are widely distributed throughout North America in aquatic systems from 0.1 to 52,000 km² in area (Jenkins and Burkhead 1994). Such a wide range in habitat size suggests that perch may be exposed to a wide variety of foraging conditions throughout their distribution. Ontogenetic changes in taxonomic and size selectivity have been observed for larval perch within a variety of systems (surface area 0.1–392 km², mean depth 5–12 m; Bulkley et al. 1976; Mills et al. 1984; Wahl et al. 1993). In a comparative study of two lakes, Siefert (1972) found that yellow perch larvae in a shallow eutrophic lake (1.51 km², maximum depth 3.1 m) followed a typical selective pattern, first feeding on copepod nauplii, then on copepodes, and finally on the larger copepods and cladocerans; however, perch larvae in a deeper, oligotrophic lake (8.1 km², maximum depth 34 m) selected for rotifers at first feeding and never showed positive selection for cladocerans. Such observed differences in selective pattern for larval perch between systems suggest that larval yellow perch can respond to variance in prey composition by changing their feeding behavior independent of ontogenetic development. Such plasticity in feeding behavior may be important to comparisons of larval feeding, growth, and survival between systems that differ in prey community composition.

Yellow perch larval feeding has been studied in lakes of widely varying sizes (0.1–392 km²), and prey community composition is not often cited as a potential cause of annual variation in survival for larval yellow perch within particular systems. Still, there are major differences between even the largest of these lakes and a more meso-oceanic system like Lake Michigan (52,000 km², mean depth 89 m), such as the rotifer- and copepod-dominated zooplankton community of Lake Michigan (Madenjian et al. 2002), and more spatial variation in the abundance of larger zooplankton (Fulford 2003). The significance of these differences for comparisons of yellow perch feeding and growth between Lake Michigan and other systems, and the corresponding implications for larval survival, are unknown.

We conducted a series of laboratory experiments to quantify larval selectivity as a function of both larval size and prey type using a broader range of natural prey choices than has been used in previous selection studies on larval yellow perch. We then used these data to develop an empirical function designed to describe the diet of larval yellow perch with minimal assumptions regarding optimality of prey. This function combined Chesson’s alpha (α) as a measure of the larval preference and prey community composition from field data as a measure of prey availability to predict prey selection of individual larva. We used this function to build an individual-based foraging and growth model (IBM) uniquely suited to addressing how differences among prey communities may affect foraging selectivity and diet of larval yellow perch. We used this IBM to ask how larval prey selection and diet may change between Lake Michigan and a smaller regional system (Green Bay) and how these differences may translate to differences in larval growth rate and survival between systems.

Green Bay (4212 km², mean depth 20 m) is a shallow, productive embayment connected to Lake Michigan at the Lake’s northwest corner and has been the focus of research into the growth and survival of larval yellow perch (Bremigan et al. 2003). Green Bay is generally considered to be a large system, but it is closer to the small end of the spectrum in comparison with Lake Michigan and has a zooplankton composition similar to that observed in smaller systems described above, where yellow perch feeding selectivity has been studied, such as Oneida Lake, New York (Hansen and Wahl 1981; Mills and Forney 1981), and Lake Mendota, Wisconsin (Schaeel et al. 1991). Comparisons of perch foraging dynamics between the main body of Lake Michigan and Green Bay should be very informative regarding differences in larval yellow perch foraging behavior between different sizes of systems. Further, Green Bay is located adjacent to Lake Michigan and is exposed to similar regional conditions. The proximity of the two systems offers an opportunity to address differences between systems resulting from differences in system size without the confounding effect of larger-scale factors such as climate or geography.

Methods

Selectivity experiments
A population (hereafter main population) of yellow perch larvae was established in the laboratory from egg skeins collected in Lake Michigan. Skeins were manually stripped from ripe females collected in gill nets about 1.2 km from shore at a depth of 10–20 m. Collected skeins were fertilized in the boat with milt from three to six males and returned to the lab for incubation. Hatching occurred within 12–14 days of fertilization, and larvae were maintained in the lab in a 2.4 m diameter tank at 15–18 °C under flow-through conditions. Larvae were initially fed a tank culture of rotifers and small zooplankton four to six times per day. Starting 5 days posthatch (dph), larvae were fed *Artemia* nauplii four times per day. At 10 dph, food changed again to a commercial pellet fed four times per day from automatic feeders.

Zooplankton prey for the experiments were collected from two sites: nearshore Lake Michigan and Lake Nagawicka (3.7 km², mean depth 11 m), which is located 40 miles (1 mile = 1.609 km) west of Milwaukee, Wisconsin. Zoo--
Zooplankton from nearshore Lake Michigan were collected 1.2 km from shore in waters 25 m deep by towing a 1 m diameter, 64 µm mesh plankton net in a circular pattern 2 m below the surface. Two 15 min tows were completed at each of two sites about 1 km apart along the same depth contour. Zooplankton from Lake Nagawicka were collected at a single site 0.5 km from shore in waters 20–30 m deep as described above, with the exception that net diameter and tow length were reduced to 0.5 m and 5 min, respectively. All zooplankton were returned to the lab and maintained under mild aeration.

Experimental trials were conducted in 2000 and 2001. The length range for larval yellow perch used in selectivity trials combined over both years was 5.5–35 mm total length (TL; 2–50 dph). In 2000, trials were conducted with three larval ages: 2 dph (mean length = 5.5 mm, standard deviation (SD) = 0.41 mm), 15 dph (8.6 mm, 0.91 mm), and 30 dph (12.3 mm, 1.4 mm). To begin each trial, 200 larvae were transferred from the main population to each of five 38 L aquaria and allowed to acclimate for 7 days. These larvae were maintained at 15–18 °C under flow-through conditions and were fed to satiation with live zooplankton collected from Lake Nagawicka for the first 5 days of the acclimation period. Feeding stopped on the sixth day of acclimation to ensure larvae would feed during the trial period and to allow pretrial zooplankton to be flushed from the system. Pretrial water samples indicated that no zooplankton remained in the trial tanks at the beginning of each trial period.

For this experiment, the objective was to present yellow perch larvae with as wide a range of prey choices as possible to assess larval selectivity under optimal conditions. Therefore, to begin a trial, Lake Nagawicka and nearshore Lake Michigan zooplankton were mixed equally and introduced into each trial tank at a target density of 250 organisms·L⁻¹. The number of zooplankton introduced into each tank was standardized based on zooplankton mass according to a predetermined mass–density relationship for mixed zooplankton (R. Fulford, unpublished data).

In 2001, trials were conducted at four larval ages: 15 dph (mean length = 7.9 mm, SD = 0.6 mm), 30 dph (11.6 mm, 1.3 mm), 40 dph (15 mm, 1.4 mm), and 50 dph (21.5 mm, 3.0 mm). Seven days prior to each trial date, larvae were moved from the main population to the trial tanks, and the acclimation process proceeded as described above for the 2000 trials. In 2001, we wished to measure larval selectivity changes in response to differences in the zooplankton community between nearshore Lake Michigan and an inland lake (Lake Nagawicka). Therefore the experimental system was expanded to 10 aquaria and two zooplankton treatments: nearshore Lake Michigan and Lake Nagawicka. Zooplankton from nearshore Lake Michigan and Lake Nagawicka were collected and maintained as described for the 2000 trials. For each trial, tanks containing acclimated larvae were randomly assigned to one zooplankton treatment or the other (nearshore Lake Michigan or Lake Nagawicka, n = 5 per treatment). Zooplankton were introduced into each tank at a target density of 250 organisms·L⁻¹. The number of zooplankton introduced into each tank was standardized based on zooplankton mass and a predetermined mass–density relationship established separately for each treatment.

In both years, larvae were allowed to feed for 30 min, and then they were removed from the tank, euthanized in tricaine methanesulfonate (MS-222), and preserved in 95% ethanol for stomach analysis. Zooplankton were sampled at the beginning and end of each 30 min feeding trial by lowering a 4 cm diameter polyvinyl chloride tube onto four randomly placed rubber stoppers on the bottom of each tank to collect four replicate 250 mL samples. Collected zooplankton were preserved in 95% ethanol for identification and enumeration. Data collected were used to calculate larval diet selectivity and the difference in zooplankton composition between treatments. Zooplankton samples collected at the end of each trial were also used to check resource depletion during the trial period. The objective was to keep resource depletion to less than 50% in any single trial. Maximum proportion of zooplankton consumed during any trial was 40% (mean proportion 22%).

Sample analysis
Mean total length on each trial day was calculated from length measured to the nearest 0.1 mm for 20 euthanized larvae from each trial tank. Stomach contents of preserved larvae (n = 30) were identified and enumerated by taxonomic group at 8× magnification. Taxonomic zooplankton groups were defined as rotifers, copepod nauplii, cyclopoid copepods, calanoid copepods, small cladocerans, and Daphnia spp. (Table 1). We chose to separate zooplankton into these groups based on consideration of differences in zooplankton size, morphology, and what is typically reported in the diet of larval yellow perch (Bulkley et al. 1976; Schael et al. 1991). The small cladoceran prey group was composed almost entirely of Bosmina spp. and Eubosmina spp. Lengths of whole prey items found in each stomach were measured to the nearest 0.1 mm with a digital imaging system (ImagePro 5.1, Media Cybernetics Inc., Silver Spring, Maryland).

Zooplankton in samples collected from each tank were also identified and enumerated by taxonomic group based on a complete count of samples (n = 8 per tank) from both the beginning and end of the trial period. Mean size of zooplankton from the tanks was measured to the nearest 0.1 mm for common taxa with a digital imaging system. Differences in zooplankton community composition among treatments were examined with a MANOVA (multivariate analysis of variance) comparison (SAS Institute Inc. 2002).

Diet selectivity
Larval selectivity was quantified with the Chesson’s α statistic (Chesson 1983). Selectivity of larva k for prey type i is defined as

\[ \alpha_{i,k} = \frac{r_{i,k}/p_i}{\sum_{j=1}^{m} r_{j,k}/p_j} \]

where \( r_{i,k} \) is the proportion of prey type \( i \) in the gut of larva \( k \), and \( p_i \) is the proportion of prey type \( i \) in the experimental tanks for \( m \) different prey types. Lockwood (1998) recommended using proportional data rather than absolute number for each prey item in experiments where it is difficult to con-
control for variation in total consumption with a time-independent stopping rule.

Chesson’s \( \alpha \) is a useful metric of selectivity for modeling because it is numerically associated with a measure of attack and capture probability. Unlike some selectivity indices that have either a clear numerical or biological definition, Chesson’s \( \alpha \) has both and it can be used in the IBM to predict larval diet; this point will be expanded in a later section. Chesson’s \( \alpha \) was calculated for each individual fish (\( n = 30 \) per tank), and a mean of these values for each tank represented a replicate value of \( \alpha \) for each treatment (\( n = 5 \) per treatment). For the analysis of selectivity patterns, selection was interpreted as neutral if the 95% confidence interval for \( \alpha \) at a particular larval size included \( m^{-1} \) and either positive or negative if the 95% confidence interval was higher or lower than \( m^{-1} \), respectively.

### Zooplankton sampling

Data regarding the zooplankton community of Lake Michigan were obtained from samples collected every 3 days from June to August in 2000 and 2001. Sampling was conducted with triplicate vertical hauls of a 0.5 m diameter, 64 \( \mu \)m mesh plankton net at each of four sites between 1 and 5 km from shore. Samples were collected from the top 10 m of the water column at all sites. Zooplankton were preserved in 95% ethanol and returned to the lab for identification and enumeration of taxonomic groups (Table 1). Three 5 mL subsamples from each main sample were enumerated at 20x magnification in a counting wheel, and the combined count for all three subsamples was converted to density for each prey type based on a constant sample volume (sample depth x net mouth area).

Zooplankton community data for Green Bay were based on samples collected in 1998 and 1999. Samples were collected in a similar manner weekly in May and June at a site near Little Tail Point, Wisconsin (B. Belonger, unpublished data). The sampling period in Green Bay was earlier than in Lake Michigan because the larval period for yellow perch begins about a month earlier in Green Bay. Zooplankton were identified and enumerated as described for samples from Lake Michigan (J. Dettmers, unpublished data). Data from different years were used because comprehensive zooplankton data were not available for both systems in the same years. All zooplankton groups collected in zooplankton samples in Green Bay and Lake Michigan were present in collections used in laboratory experiments.

## Table 1. Mean length and dry mass of the six larval prey types.

<table>
<thead>
<tr>
<th>Prey item</th>
<th>Length (mm)</th>
<th>Intercept (( a ))</th>
<th>Exponent (( b ))</th>
<th>Mass (( \mu )g)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rotifers</td>
<td>0.13</td>
<td>1.84</td>
<td>1.44</td>
<td>0.10</td>
<td>Dumont 1975</td>
</tr>
<tr>
<td>Copepods</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nauplii</td>
<td>0.2</td>
<td>3.0</td>
<td>1.71</td>
<td>0.19</td>
<td>Culver et al. 1985</td>
</tr>
<tr>
<td>Calanoid</td>
<td>0.54</td>
<td>6.19</td>
<td>1.96</td>
<td>1.85</td>
<td>Culver et al. 1985</td>
</tr>
<tr>
<td>Cyclopoid</td>
<td>0.43</td>
<td>6.66</td>
<td>2.89</td>
<td>0.58</td>
<td>Culver et al. 1985</td>
</tr>
<tr>
<td>Small cladocerans(^a)</td>
<td>0.3</td>
<td>17.74</td>
<td>2.22</td>
<td>1.2</td>
<td>Culver et al. 1985</td>
</tr>
<tr>
<td><em>Daphnia</em> spp.</td>
<td>0.9</td>
<td>7.50</td>
<td>1.56</td>
<td>6.4</td>
<td>Culver et al. 1985</td>
</tr>
</tbody>
</table>

Note: Dry mass was calculated from mean length with relationships taken from the literature. Function used is mass = \( a \times \) length\(^b\).

\(^a\)Small cladocerans were predominantly from the genus *Bosmina*; the length–mass function used is for *Bosmina longirostris*.

### Individual-based foraging model

To examine the relationship between diet and growth for larval yellow perch, we adapted a general, larval fish, foraging IBM (Letcher et al. 1996) to predict growth and starvation rates of larval yellow perch from hatch to 45 dph (Fig. 1). This model predicted daily consumption of each prey type (Table 1) for each individual larva \( k \) based on larval length (\( l \), mm) and a stochastic, sized-based encounter rate (\( E_R_{i,k} \)), handling time (\( H_{T,i,k} \)), and probability of attack and capture (\( Q_{i,k} \)). Encounter rate and handling time were calculated using the general model with parameters specific to larval yellow perch (Table 2); full details of these calculations can be found in Fulford (2003).

Probability of attack and capture for larva \( k \) on prey item \( i \) in a given day was calculated in the model with an empirical function based on the Chesson’s \( \alpha \) metric and derived from our size-based selectivity experiments (Fig. 1). The probability that the next prey item attacked and captured by larva \( k \) will be of prey type \( i \) (\( Q_{i,k} \)) has two components: (i) the selectivity of the larva defined empirically in our experiments and expressed as \( \alpha_{i,k} \) and (ii) the relative abundance \( p_i \) of prey type \( i \) on a given model day.

The best-fit function between \( \alpha_{i,k} \) and larval size (5–35 mm TL) was found separately for each prey type \( i \) based on a comparison of the least-squared fit to the data for a series of common functions. Function types tested were linear, logarithmic, exponential, second-order polynomial, generalized logistic, and double Weibull.

Chesson’s \( \alpha \) is generally insensitive to changes in prey relative abundance; however, selectivity will always be near zero for any prey type when its relative abundance falls below a threshold value (Chesson 1983). When relative density for a particular prey item falls below this threshold, negative selectivity for that prey item will be indicated because larvae did not encounter this rare prey type, not because of prey type avoidance. Therefore, a threshold was defined as the relative abundance below which selectivity was always negative because of an absence of that prey item in larval guts; trials for which relative abundance of any prey item was less than the threshold were not used to fit the larval size–selectivity relationship for that prey item.

The best function for each prey type was based on a coefficient of multiple determination (\( R^2 \)) adjusted for the number of parameters in the model (Neter et al. 1990). The best-fit function for each prey type \( i \) was then used in the IBM to predict \( \alpha_{i,k} \) for prey type \( i \). Data used for calculations of \( \alpha_{i,k} \) for larvae larger than 25 mm TL included diet and field zoop-
Fig. 1. Flow chart summarizing the individual-based model for foraging and growth used for the numerical experiments. Starvation threshold was set at 53% of previous maximum dry mass for all model simulations.

**Foraging submodel**

- **Larva hatched**
  - Hatch date
  - Initial length (mm)
  - Initial dry mass (μg)

**Handling time (s)**

\[
HT_{i,k} = e^{-0.264 \cdot 10^{7.0151(PL_i/l)}}
\]

- **PL_i** = length of prey type i (mm)
- **l** = length of individual larva (mm)

**Probability of attack and capture**

\[
Q_{i,k} = \frac{a_{i,k} \cdot p_i}{\sum_{j=1}^{m} a_{k,j} \cdot p_j}
\]

- **a_{i,k}** = selectivity of larva k for prey type i generated from selectivity experiments
- **p_i** = Relative abundance of prey type i

**Daily consumption (μg·day\(^{-1}\))**

\[
C_k = \frac{\sum_i (B_{i,k} \cdot m_i)}{1 + \sum_i E_{i,k} \cdot Q_{i,k} \cdot HT_{i,k}} \cdot 43,200
\]

- **B_{i,k}** = stochastic estimate of number of prey type i consumed by larval k based on binomial calculation with **E_{i,k}** as total trials and **Q_{i,k}** as success probability.
- **m_i** = individual mass of prey type i

**Bioenergetics submodel**

- **Daily growth (μg·day\(^{-1}\))**
  \[
  G = (C_k \cdot AE_k) - R_k - [C_k \cdot (U + SDA)]
  \]

- **AE_k** = assimilation efficiency based on mass of larva k
- **R_k** = daily costs of respiration for larva k (μg·day\(^{-1}\))
- **U** = daily costs of excretion
- **SDA** = daily costs of specific dynamic action

- **Temperature Larval dry mass (μg)**

**Reached starvation threshold?**

- Yes → Larva dead
- No →

**Update larval mass (μg·day\(^{-1}\))**

**Update larval length (mm·day\(^{-1}\))**
plankton data from the literature (Bulkley et al. 1976; Mills et al. 1984). Diet selectivity indices were calculated for yellow perch in these studies, but in cases where the index was not Chesson’s α, we converted the selectivity index to Chesson’s α. In the model, \( Q_{i,k} \) is calculated each day for each larva \( k \) and each available prey item \( i \).

The number of each prey type consumed in a given day was then converted to mass (\( \mu g \) dry mass) by multiplying the number of each prey type eaten in a day by the mean mass of prey type \( i \). Mean mass for each prey type was derived from a prey type specific, length–mass relationship and mean length for each prey type taken from zooplankton collected in Lake Michigan (Table 1). Zooplankton relative abundance in the model was based on the average seasonal pattern observed during the yellow perch larval period over 2 years for each prey type taken from the field data for either Green Bay or Lake Michigan. Zooplankton density was manipulated as a part of the numerical experiments (described below).

### Individual-based bioenergetics model

Total mass consumed by larva \( k \) each day was converted to individual daily growth (\( \mu g \) dry mass) in a bioenergetics submodel adapted from two models: one developed previously for larval yellow perch (Hanson 1997) and a general larval bioenergetics model developed specifically for an IBM approach (Letcher et al. 1996). The combined model uses a mass-balance approach to predict daily growth (\( G, \mu g\text{day}^{-1} \)) from predicted daily consumption (\( C, \mu g\text{day}^{-1} \)) from the foraging submodel (Fig. 1). Larval mass was updated daily based on predicted growth. During a model run, a larva could not lose length but could lose mass based on the difference between total daily consumption predicted by the foraging submodel and calculated daily metabolic costs. If a larva lost more than 53% of its previous maximum mass at any point in a model run, then that larva starved to death. If larval mass increased beyond the previous maximum value for larva \( k \), length was updated based on a length–mass conversion (Table 2). Full details regarding the bioenergetics model used are available in Fulford (2003).

Input to the bioenergetics submodel was daily mean water temperature and current dry mass of larva \( k \). Daily mean temperature data were taken from a variety of sources depending on the system being modeled. Temperature data for nearshore Lake Michigan were taken from hourly temperature records collected 1 m below the surface at a site located 1.2 km from shore due east of Texas Rock near Milwaukee, Wisconsin (42°59’60”N, 87°50’38”W), from June to August 1999–2001 (R. Fulford, unpublished data). Temperature data for Green Bay were taken from temperature data collected hourly at the surface of the intake canal for the Pulliam power plant in Green Bay, Wisconsin, and from weekly temperature measurements taken at the surface 0.5 km from shore near Little Tail Point on the southwestern side of Green Bay. Little Tail Point is located 15 km north of the City of Green Bay. The area around Little Tail Point is a known spawning area for yellow perch (B. Belorder, unpublished data).

Each model simulation began with 1000 larvae, and initial size was randomly assigned to each larva from a normal distribution (mean = 5.7 mm, SD = 0.3 mm). Initial mass was calculated from randomly assigned length according to a larval yellow perch specific mass–length formula (Table 2). Model output was larval size-specific diet data, individual and cohort mean growth rate over the period from hatch to 45 days, distribution of size at age for the entire cohort, and the proportion of larvae that starved to death during each model run.

### Numerical experiments

We conducted two numerical experiments with the foraging and growth IBM to address two questions. First, we wanted to establish that the model would accurately predict diet for larvae exposed to different prey assemblages. To accomplish this, the functions relating \( \alpha_{ik} \) to larval TL were fit to a subset of the data from selectivity trials (\( n = 45 \) trials).
als). We then used the model to predict the diet of larvae exposed to the prey assemblage in the remaining set of reference data (n = 10 trials). The reference data were for larvae between 7 and 15 mm TL and were broken into two sets: larvae exposed to a zooplankton community from nearshore Lake Michigan and larvae exposed to a zooplankton community from Lake Nagawicka. These predictions were then compared with a χ² analysis to observed diet for larvae used in the reference trials.

Second, we asked how larval growth rate, size at age, and likelihood of starvation are affected by differences in zooplankton community composition between Green Bay and Lake Michigan. The model was run with two separate prey assemblages consisting of zooplankton composition observed in nearshore Lake Michigan in 2000–2001 or in Green Bay in 1998–1999. Simulations were conducted at 50, 100, 150, 200, and 250 prey·L⁻¹ for each zooplankton assemblage. A broad range of zooplankton densities was used to explore more generally relevant patterns in growth and survival as a function of prey density than observed in just the 2 years for which we had data. We also wanted to explore more directly how the interacting effects of prey density and composition on larval growth and starvation mortality differ between nearshore Lake Michigan and Green Bay. The range of zooplankton density used in this experiment approximates the full range of daily and site-specific mean zooplankton densities observed in both nearshore Lake Michigan and Green Bay (Fulford 2003; J. Dettmers, unpublished data). Three replicate model runs were conducted for each combination of zooplankton density and zooplankton assemblage to account for variability due to stochastic functions in the model.

Results

Selectivity experiments

The three zooplankton assemblages used in selectivity trials (nearshore Lake Michigan, Lake Nagawicka, and mixed) differed significantly in composition (Wilks’ λ F₁₂,₉₄₁ = 12.34, p < 0.0001). The Lake Michigan treatment was dominated (>70%) by rotifers, while the Lake Nagawicka assemblage was composed largely of Daphnia spp. and cyclopoid copepods. The mixed assemblage was the most balanced among prey types but remained high in rotifers (>40%). Mean initial zooplankton density in the trials tanks was 382 prey·L⁻¹ and ranged from 112 to 782 prey·L⁻¹. The higher than expected mean and variance in zooplankton density in the tanks was due largely to variance in rotifer density, because rotifers can vary in number without having much effect on total mass of a zooplankton sample. However, high zooplankton density in the tanks should not bias our measurements of selectivity as long as density is high enough to generate larval feeding activity.

Values of α for rotifers indicated strong positive selection by 5.5 mm larvae, but dropped quickly to near zero, indicating negative selection at all larger larval sizes (Fig. 2a). Values of α for copepod nauplii were also highest for larvae at 5.5 mm TL, but selection was neutral rather than positive. Selection for nauplii was generally neutral up to 15 mm TL, and α values showed no trend with size (Fig. 2a).

Selection differed for cyclopoid and calanoid copepods (Fig. 2b). Values of α for cyclopoid copepods were initially neutral but rose sharply for larvae at 8 mm TL, indicating a period of strong positive selection by early-feeding larvae. Larval selection for cyclopoid copepods was neutral for larvae larger than 12 mm TL. Data from the literature used in the model also suggest that selectivity for cyclopoid copepods by yellow perch larvae at 26.5 mm TL is neutral (Bulkley et al. 1976).

Selection for calanoid copepods was negative for yellow perch larvae at 5.5 mm TL. Values of α rose with larval size beginning around 8 mm TL and were generally positive for larvae larger than 12 mm TL (Fig. 2b). Data from the literature used in the model suggest that selection for calanoid copepods by larvae at 26.5 and 35 mm TL is consistently positive (Bulkley et al. 1976; Mills et al. 1984).

Selectivity patterns for cladocerans all showed similar patterns with slightly different shapes (Fig. 2c). Selectivity for small cladocerans was negative for all larvae less than 12 mm TL, became neutral for larvae between 12 and
Table 3. Summary of the model fit between Chesson’s $\alpha$ and larval total length ($L$, mm) for each prey type based on experimental data.

<table>
<thead>
<tr>
<th>Prey Item</th>
<th>Model type</th>
<th>Parameters</th>
<th>$R^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rotifers</td>
<td>Power: $\alpha = a L^b$</td>
<td>$a = 193.499$</td>
<td>0.89</td>
</tr>
<tr>
<td></td>
<td></td>
<td>$b = -7.64$</td>
<td></td>
</tr>
<tr>
<td>Copepods</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nauplii</td>
<td>No trend</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cyclopid</td>
<td>Polynomial: $\alpha = a_1 L^2 + a_2 L + a_3$</td>
<td>$a_1 = -0.042$</td>
<td>0.67</td>
</tr>
<tr>
<td></td>
<td></td>
<td>$a_2 = 0.75$</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>$a_3 = -2.60$</td>
<td></td>
</tr>
<tr>
<td>Calanoid</td>
<td>Log-linear: $\alpha = a \ln(L) + b$</td>
<td>$a = 0.272$</td>
<td>0.68</td>
</tr>
<tr>
<td></td>
<td></td>
<td>$b = -0.3834$</td>
<td></td>
</tr>
<tr>
<td>Small cladocers</td>
<td>Logistic: $\alpha = \frac{a_1}{1 + \left( \frac{a_1}{a_2 - 1} \right) e^{-a_3 a_4 L}}$</td>
<td>$a_1 = 0.40$</td>
<td>0.65</td>
</tr>
<tr>
<td></td>
<td></td>
<td>$a_2 = 0.09$</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>$a_3 = 13$</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>$a_4 = 0.031$</td>
<td></td>
</tr>
<tr>
<td>Daphnia spp.</td>
<td>Logistic: $\alpha = \frac{a_1}{1 + \left( \frac{a_1}{a_2 - 1} \right) e^{-a_3 a_4 L}}$</td>
<td>$a_1 = 0.445$</td>
<td>0.86</td>
</tr>
<tr>
<td></td>
<td></td>
<td>$a_2 = 0.092$</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>$a_3 = 20$</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>$a_4 = 0.029$</td>
<td></td>
</tr>
</tbody>
</table>

Note: Larval size range was 5.5–35 mm total length for all functions, and data for larvae >25 mm total length includes data from the literature.

Parameterizing the IBM

Based on the selectivity data, the relative abundance minimum threshold for inclusion in the function was set at 0.03%. When relative abundance of a particular prey item in a trial was below 0.03% of the assemblage, data from that trial were not used to fit the larval size – selectivity relationship for that prey item.

No single function fit the data relating Chesson’s $\alpha$ to larval TL best for all six prey types (Table 3). Selectivity for rotifers was best described by a power function (Fig. 3a). The model fit was highest for rotifers among all prey types. We observed no relationship between larval selectivity for copepod nauplii and larval size; selectivity for nauplii in the model was described with the mean value of $\alpha$ for nauplii across all larval sizes (0.07; Fig. 3b). Selectivity for cyclopoid copepods was best described by a second-order polynomial function with a minimum $\alpha$ level set at $m^{-1}$ (Fig. 3c). This function allowed for the steep rise in $\alpha$ for larvae around 8 mm TL and predicted neutral selection for larvae larger than 12 mm TL. Selectivity for calanoid copepods was best described by a logarithmic regression, with a maximum $\alpha$ value of 0.4 for larvae larger than 15 mm TL (Fig. 3d). Selectivity for both small cladocerans and Daphnia spp. was best described by a generalized logistic function (Figs. 3e, 3f). Values of $\alpha$ for both cladoceran groups were low initially and rose with increasing larval size.

Zooplankton community composition

The nearshore Lake Michigan zooplankton assemblage was dominated numerically by rotifers (80%; Fig. 4a). Prey densities across sites in nearshore Lake Michigan in 2000–2001 were between 50 and 200 prey·L$^{-1}$, with a 2-year mean density of 75 prey·L$^{-1}$. In contrast, the prey assemblage of Green Bay was more diverse. Cladocerans and copepods were nearly equal in abundance, and no prey type represented more than 50% of the total assemblage at any time over the larval period (Fig. 4b). Prey densities in 1998–1999 in Green Bay were between 50 and 300 prey·L$^{-1}$, with a 2-year mean density of 124 prey·L$^{-1}$ over both years.

Numerical experiments

In experiment one, model predictions of diet for larvae feeding on a nearshore Lake Michigan or Lake Nagawicka zooplankton assemblage did not differ significantly from observed diet ($\chi^2 = 1.23$, degrees of freedom (df) = 5, $p > 0.1$), which suggests the model performed well (Fig. 5). Diets for larvae exposed to the nearshore Lake Michigan assemblage were dominated by calanoid and cyclopoid copepods, with rotifers and cladocerans being a minor dietary component. In
contrast, diets of larvae exposed to the Lake Nagawicka assemblage were composed mostly of *Daphnia* spp., with calanoid copepods a distant second in relative dietary abundance. There were noticeable differences between what was available to and what was eaten by larvae, particularly for the nearshore Lake Michigan treatment, where larvae avoided rotifers despite their high relative abundance. However, available prey composition did not differ significantly from either observed diet or model predictions ($\chi^2 = 1.3$, df = 5, $p > 0.1$).

In experiment two, larvae exposed to the nearshore Lake Michigan assemblage in the model fed mainly on rotifers and copepod nauplii initially and made a transition to mainly larger calanoid copepods as they grew, mainly at the expense of rotifers (Fig. 6a). Larvae exposed to the Green Bay assemblage in the model fed largely on cyclopoid copepods initially, avoided the smaller rotifers, and shifted gradually to a diet split between small cladocerans and *Daphnia* spp. as they increased in size. The transition to small cladocerans and *Daphnia* spp. was predicted to occur around 20 mm TL, and these two items dominated larval diet in Green Bay for the remainder of the model run (Fig. 6b).

These observed differences in diet between larvae exposed to different prey assemblages translated into differences in both growth and survivorship in the model. Larvae allowed to forage on the Green Bay assemblage had a higher predicted mean growth rate compared with nearshore Lake Michigan larvae at all prey densities except the lowest (50 prey·L$^{-1}$). Growth rates for larvae exposed to the nearshore Lake Michigan assemblage did not rise substantially until prey density rose above 100 prey·L$^{-1}$ (Fig. 7). In contrast, larvae exposed to the Green Bay assemblage starved at prey densities fewer than 150 prey·L$^{-1}$ (Fig. 8). Starvation rates for larvae exposed to the nearshore Lake Michigan assemblage declined linearly with increasing prey density over the entire density range (50–250 prey·L$^{-1}$). In contrast, larvae exposed to the Green Bay assemblage had a high starvation rate at the lowest density, but it dropped rapidly to around 20% or lower for prey densities at or above 150 prey·L$^{-1}$ (Fig. 8). These differences in growth and survivorship were reflected in larval size distributions at the end of the model runs. Larval survivors that fed on the Green Bay prey assemblage (Figs. 9b, 9d) had a wider size range and were generally larger than larval survivors that fed on the nearshore Lake Michigan assemblage (Figs. 9a, 9c) at the same prey density. At a prey density of 250 prey·L$^{-1}$, both distributions were bimodal, but the major mode in the length distribution for larvae exposed to the Green Bay assemblage (mode centered on 34 mm TL) included most of the larval survivors. The minor mode in the length distribution of the Green Bay assemblage at ~20 mm TL overlapped the major mode in the length distribution of the nearshore Lake Michigan assemblage (at ~18 mm TL). The minor mode in the length distribution of larval survivors exposed to the nearshore Lake Michigan assemblage (7 mm TL) was composed of larvae demonstrating little or no growth over the model period. In contrast, the minor mode of larvae exposed to the Green Bay assemblage was composed of larvae that grew substantially.

The size distribution of larval survivors exposed to both prey assemblages at a lower mean density (100 prey·L$^{-1}$) had similar length ranges as observed at the higher prey density, but the peak in larval size dropped 5 and 13 mm for larval exposed to the nearshore Lake Michigan and Green Bay assemblages, respectively (Figs. 9a, 9b). In the case of larvae exposed to the nearshore Lake Michigan assemblage, the
distribution was no longer bimodal, as observed in this distribution at the higher prey density.

Discussion

Diet shifts and community composition

Larval yellow perch appear to make three distinct transitions in selection throughout ontogeny: from rotifers to small copepods to large copepods and cladocerans. However, the first and last transitions seem most dependent on prey community composition.

Larval yellow perch initially showed positive selection for rotifers in our selectivity experiments. This early preference for rotifers has been observed in field data from other oligotrophic systems (Siefert 1972). Surprisingly, a preference for rotifers appears to lower predictions of larval survival. Larvae that switched to small copepods at a smaller size in the model had a distinct growth advantage over larvae that fed largely on rotifers, but this energetic conclusion does not appear to translate to a high preference for copepods in early-stage larvae. Low or negative growth for yellow perch larvae that have been fed rotifers has been observed previously in laboratory experiments (Graeb et al. 2004).

It could be that first-feeding larval yellow perch select rotifers simply because they are much more abundant and easy to catch and that this initial positive selection is dependent on limited foraging ability rather than true preference. The full recruitment of visual acuity occurs after first feeding in larval yellow perch (Wahl et al. 1993); first-feeding larvae are likely to select prey based on a more immediate definition of optimality (i.e., avoiding starvation). Model results do predict that larvae exposed to the Green Bay prey community (i.e., low rotifer abundance) will switch to small copepods earlier than larvae exposed to a nearshore Lake Michigan prey community (i.e., high rotifer abundance), and early-stage larvae feeding on copepods have a major growth advantage over larvae that are not. Such selective choices demonstrate the difference between the long- and short-term optimality of foraging decisions. In the short term, it is clearly optimal to eat rotifers rather than nothing at all. However, in the long term, larvae that shift to higher value prey will grow faster and are more likely to survive the larval stage. Model predictions suggest that larvae that fail to make this early transition may be trading a death from starvation for a death by size-selective processes (i.e., predation) and are no more likely to contribute to the year class than larvae that eat nothing at all.

The final transition in larval diet prior to the beginning of the juvenile stage involves a community-dependent trade-off between larger copepods and cladocerans. Larval perch were predicted to select for cladocerans at intermediate larval sizes even when cladocerans comprised only 1% of the prey assemblage previously in laboratory experiments (Graeb et al. 2004).

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the model shifted their selection to larger calanoid copepods. Larval yellow perch larger than 16 mm TL showed positive selection for calanoid copepods in all assemblages, but the importance of calanoids tended to vary based on the relative abundance of cladocerans.

Laboratory experiments have demonstrated that late-stage larval perch positively select for calanoid copepods and incur a growth advantage by doing so (Confer and Lake 1987). Nevertheless, positive selection for cladocerans has been consistently observed for yellow perch larvae in field data from other systems, including Lake Mendota (Schael et al. 1991) and Oneida Lake (Hansen and Wahl 1981; Mills et al. 1984), but the timing of this transition with respect to larval size varies greatly. Variance in the observed onset of positive selection for Daphnia spp. may be due in part to variance in the composition of the Daphnia complex between systems. Daphnia spp. collected in Green Bay, Lake Nagawicka, and nearshore Lake Michigan were mainly medium-bodied Daphnia spp. (0.62–1.17 mm). In contrast, Oneida Lake is dominated by Daphnia pulex, which has a size range of 0.7–2.1 mm (Mills et al. 1984). Positive preference for Daphnia spp. was evident at a smaller larval size in our data compared with similar data from other systems; this difference is most likely due to observed differences in the size of the Daphnia complex. Such differences are an important reason to understand both size and taxonomic selectivity for larval perch.

**Predicting larval diet shifts**

Our empirical approach to predicting larval diet is well suited to understanding larval responses to changing prey community composition. Traditional approaches to modeling larval foraging have used optimization rules, such as maximization of the benefit–cost ratio (Eggers 1977). This approach has two distinct disadvantages for comparing feeding selectivity between different prey assemblages. First, optimization rules are frequently based on inherent qualities of prey, and decisions regarding prey type \(i\) do not directly take into account the relative abundance of all available prey types. As a result, changes in diet composition in optimality models often occur abruptly when prey density crosses a dis-
crete threshold. Such changes result in prey types being included or not included in the diet in a “knife-edge” fashion not representative of the real world (Stephens and Krebs 1986). Such optimization-based models are not well suited to predicting gradual shifts in selection, such as shifts caused by subtle changes in prey relative abundance.

A second disadvantage of optimization rules is that they require an assumption regarding what criteria would be of value to the larva to rank prey types. This criterion is usually size, as this is the most numerically and biologically tenable metric available (Eggers 1977). Prey frequently differ in other significant ways, such as behavior and morphology (Kerfoot et al. 1980), which are more difficult to quantify.

Our empirical approach to predicting prey selection for larval yellow perch allows for selection without assumptions regarding optimal criteria. That is, size is considered as a criterion for selection just as it is in an optimality model, but it is bundled with other less mathematically tenable characteristics of prey that may also be important. This approach exploits a selectivity metric (Chesson’s $\alpha$) that describes changes in larval preference, is mathematically tenable, and is resistant to most changes in the prey assemblage. There is a minimum relative abundance greater than zero, below which selectivity data are affected, but it was extremely low in this study (0.03%) and is only likely to be important for extremely simple prey communities (Confer and Moore 1987). By utilizing empirical data, we were able to more accurately describe how selection for a particular prey item may change both as a function of larval size and as larvae are exposed to different prey assemblages.

Two potential sources of bias exist in this empirical approach to measuring larval selectivity. First, a significant amount of variation among prey may exist within the taxonomic prey types. We have tried to define prey types in our model so that this bias is minimized. It does appear that variation among species of Daphnia may be important. The onset of positive selection for Daphnia spp. occurred relatively early in our selectivity trials compared with observations of yellow perch in smaller systems, likely because of differences in the size of the dominant Daphnia species between systems like Oneida Lake and Lake Michigan. Differences within the Daphnia species complex would need to be addressed to increase the generality of model results. These differences strengthen the argument that comparisons of selectivity data for larval yellow perch between systems warrant greater scrutiny.

A second potential source of bias in this analysis is differential digestion of hard- vs. soft-bodied zooplankton in larval guts (Sutela and Huusko 2000). Smaller, softer-bodied prey such as rotifers would be less likely to be present in the gut, so selection for these prey types might be underestimated. By limiting the foraging experiments to half an hour, we likely minimized the impact of this bias in comparison with analysis of field-collected larvae, for which total digestion time is unknown. Our results generally agree with recent field-based results on larval yellow perch diet in Green Bay (Bremigan et al. 2003); however, digestive bias may also be a reason why rotifers are not often cited as an important item in larval guts from field data.

Data from the literature suggest that larvae of comparable size in different systems can make very different foraging decisions. For instance, larval perch in Clear Lake, Iowa, fed primarily on copepods up to a larval size of 25 mm TL (Bulkley et al. 1976), but larvae of similar size in Oneida Lake, New York, showed a strong positive selection for Daphnia spp. (Mills et al. 1984). Similar studies on larval and juvenile European perch (Perca fluviatilis) have shown variance in selectivity for cladocerans vs. copepods among systems ranging from small lakes to coastal embayments, which suggests that variable selection based on prey community composition is a trait these two species share (Persson and Greenberg 1990; Mehner 2000). Such data provide evidence, as we do here, that it may be important to account for prey community composition, as well as for

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inherent preferences, when using selectivity criteria to predict diet of larval yellow perch. Moreover, the importance of community composition to a comparison of larval diet between foraging habitats will only increase as differences in community composition increase, which is more likely with increasing differences in geography and size between systems. Our empirical selectivity model is a strong tool for quantifying these differences so that we can effectively compare different yellow perch populations.

Implications of diet shifts across systems

Different foraging decisions made by larval perch in the model when exposed to nearshore Lake Michigan or Green Bay prey assemblages resulted in large differences in growth and survival. Model results suggest that larval perch in Green Bay grow faster than those in nearshore Lake Michigan at comparable prey densities. The resulting size distribution of survivors indicates that prey density affects how many individuals achieve the maximum observed growth rate, while prey community composition controls the magnitude of the maximum observed growth rate.

The bimodality in the size distributions of larval survivors predicted by the model appears to be the result of an interaction between zooplankton community composition and timing of individual hatch. Larvae that hatched into the model at times when small copepods were abundant selected for copepods at smaller larval sizes than larvae hatched when small copepods were rare relative to other available prey. This diet shift resulted in a higher initial larval growth rate, and this early growth advantage was still detectable in the bimodal size distribution of larvae 30 days after peak hatch. Post et al. (1997) observed a bimodality in the size structure of young perch cohorts in Lake St George, Ontario, in some years but not in others. They attributed this within-cohort size difference to density-dependent competition, but this splitting of a cohort into two size groups is also consistent with an early growth advantage for the part of the cohort that may be related to earlier onset of optimal feeding due to temporal differences in prey community composition.

Our model simulations did not include predation, and the character distributions of survivors would differ if predation mortality had been included. However, the higher predicted growth rate for larvae in Green Bay is also likely to result in lower vulnerability to predation. Based on differences in the size distribution of survivors, larval yellow perch growth rates in Green Bay are predicted to be more variable among individuals, resulting in a higher maximum growth rate among individuals. High growth variability among individuals means that larvae in the upper portion of the growth rate distribution are likely to reach a refuge from size-dependent predation faster; this may increase overall survival (Rice et al. 1993).

Larvae in nearshore Lake Michigan are also predicted to be less responsive to changes in prey abundance than larvae in Green Bay. Starvation rate was over 80% and mean growth rate was less than 0.1 mm·day⁻¹ for larvae in both Green Bay and nearshore Lake Michigan at the lowest prey density modeled. Starvation rate decreased and growth rate increased slowly with increasing prey density for nearshore Lake Michigan in the model, and growth rates did not approach growth rates for yellow perch we observed in the field (0.12–0.57 mm·day⁻¹) until prey densities exceeded 150 prey·L⁻¹. In contrast, both growth rate and starvation rate of larvae exposed to a prey assemblage from Green Bay improved more rapidly as prey density increased.

Many of the larvae predicted to starve in our model would likely be eaten before they starved to death in the natural world. The significance of our results is that in years when other forms of mortality such as predation are low in Green Bay, larval survival will likely be high, but variability in these other forms of mortality will have a less consistent effect on larval recruitment in nearshore Lake Michigan. Larval survival in nearshore Lake Michigan is primarily dependent on larvae finding high-density patches of prey to grow. Historical examinations of zooplankton density in Lake Michigan suggest this is less likely now than it was in the late 1980s when yellow perch recruitment was high (Dettmers et al. 2003). A strong initial dependence on foraging success means that in years when larval overlap with optimal prey is low, year-class strength will likely be low, but in years when this overlap is high, year-class strength could be high or low, depending on the importance of other forms of mortality. Consequently, good larval survival and a subsequent strong year class will occur less often in Lake Michigan than in Green Bay, increasing the dependence of longer-term population stability for yellow perch in Lake Michigan on extrinsic factors affecting larval survival.

These conclusions are based on prey community differences at a large scale (i.e., between systems). We did not attempt to take into account small-scale factors such as changes in foraging efficiency due to water clarity or turbulence. Furthermore, because we are comparing the zooplankton assemblages of Green Bay in 1998–1999 to nearshore Lake Michigan 2000–2001, we cannot make predictions about the year class in Green Bay being better than that in Lake Michigan in any particular year. Nonetheless, by studying the effect of community composition across multiple years and a range of zooplankton densities, it is possible to address the larger questions of whether differences in prey composition between systems are important to differences in larval survival and whether a single model of foraging behavior is sufficient to describe larval diet across a range of prey communities. Our model results suggest this is the case.

We did have to make several simplifying assumptions within the IBM framework. Most of these are part of the general IBM framework and have been well addressed previously (Letcher et al. 1996; Fulford 2003). In particular, larval foraging activity was assumed to be evenly spread across the day, and no feeding was assumed to occur at night. These assumptions are reasonable based on observations of yellow perch larval activity in tank systems (R. Fulford, unpublished data). Further, both predator and prey were assumed to be evenly distributed in space, and while this assumption may not always be true in nature, it is a reasonable simplification for comparisons of larval growth across systems.

It is important to have an understanding of how different factors affecting growth and survival of yellow perch populations may differ in importance between systems prior to making comparisons between those systems. Factors found to be important to growth and survival of larval yellow perch in smaller systems include density dependence (Post and McQueen 1988; Sanderson et al. 1999), predation (Campbell
1998; Mayer et al. 2000), interspecific competition (Roseman et al. 1996), and overwinter mortality (Post and Evans 1989). Owing to the size and depth of Lake Michigan, density-dependent effects are going to be ameliorated by larval dispersion, and long-term survey data indicate that year-class strength is highly correlated with juvenile abundance in the fall (Clapp and Makauskas 2002), which suggests that overwinter mortality is not an important factor. The importance of predation and competitive interactions with other species such as alewife remains open. Nevertheless, our results suggest that the interaction of prey community composition in Lake Michigan with larval yellow perch selectivity patterns results in a strong relationship between prey availability and larval survival not observed in smaller systems.

Yellow perch life history seems optimized for smaller, more productive systems where survival during the pelagic phase is more consistent between years. For populations of yellow perch in systems such as Green Bay or Oneida Lake that exhibit higher larval survival (Mayer et al. 2000), population regulation is more dependent on factors regulated by perch density, such as overwinter mortality or predation during the juvenile stage (Mayer et al. 2001). In Lake Michigan, where annual recruitment appears dependent on density-independent factors such as spatial and temporal overlap with prey, recruitment patterns observed in smaller systems would have less value for predicting annual recruitment. In such cases, the individual-based modeling approach is a valuable and flexible tool for a comparative assessment of factors important to larval survival.

For larval yellow perch, the importance of diet choice constrained by limited foraging ability (e.g., first feeding) and limited availability of optimal prey items (e.g., Lake Michigan) can be both strong and important for describing patterns in annual recruitment success. Under these conditions, it is vital to describe changes in larval prey community composition in both time and space in Lake Michigan to better characterize how foraging-mediated recruitment patterns may be important to predicting population recovery.

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