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NOTE

White Perch in Small North Carolina Reservoirs:
What Explains Variation in Population Structure?

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Abstract

White Perch Morone americana have been introduced into many inland systems throughout the United States. To determine factors affecting White Perch abundance and size structure, we compared White Perch growth, timing of maturity, and trophic level; the abundance of a predator (Largemouth Bass Micropterus salmoides); the abundance of an ecologically significant mid-level omnivore (Gizzard Shad Dorosoma cepedianum); prey availability (chironomid and zooplankton abundances); and environmental variables (specific conductivity, Secchi depth, dissolved oxygen concentration, and temperature) among four reservoirs (two with high White Perch abundance and two with low abundance). White Perch size structure was closely tied to abundance, with truncated size structure as abundance increased. Among the other variables we tested, only Largemouth Bass abundance had a significant (negative) relationship with White Perch abundance. White Perch size structure appeared to be highly density dependent, and variables that commonly explain variation in abundance of introduced fishes did not explain differences in the four White Perch populations we studied. Further study of the competitive and predatory interactions of White Perch and Largemouth Bass over ontogeny could shed light on the mechanism(s) potentially shaping population structure of the two species where they coexist.

Introductions of White Perch Morone americana have been problematic for freshwater fisheries managers (Madenjian et al. 2000; Gosch et al. 2010b). Since the 1950s, White Perch have been intentionally and unintentionally moved from their native range (i.e., North America’s Atlantic coastal estuaries and river tributaries) into freshwater lakes, inland rivers, and reservoirs of the eastern and central United States (Scott and Christie 1963; Irons et al. 2002; Harris 2006). Studies in these areas have indicated detrimental impacts of White Perch on native species, primarily through competition (Schaeffer and Margraf 1986; Parrish and Margraf 1990, 1994; Prout et al. 1990) and egg predation (Schaeffer and Margraf 1987; Hartman and Margraf 1992; Madenjian et al. 2000).

White Perch are excellent competitors and invaders due to a variety of life history traits, including their tolerance of a wide range of environmental conditions (Stanley and Danie 1983; Johnson and Evans 1990); their role as omnivorous, opportunistic feeders (Reid 1972; Hines 1981; Schaeffer and Margraf 1987; Hurley 1992; St-Hilaire et al. 2002; Couture and Watzing 2008; Gosch et al. 2010b); and their reproductive flexibility, with maturation occurring as early as age 1 or as late as age 3 (Sheri and Power 1968; Bur 1986; Schaeffer and Margraf 1986; Chizinski 2007; Feiner et al. 2012). Population dynamics of native and introduced White Perch have been linked to a variety of abiotic and biotic factors (Feiner et al. 2012, 2013), such as turbidity, dissolved solids, and specific conductivity (Hawes and Parrish 2003; North and Houde 2003) as well as predation (Ward and Neumann 1998; Margulies 1990; Vrtiska et al. 2003; Gosch et al. 2010a) and competition with other fish (Hawes and Parrish 2003). Finally, intraspecific competition in high-density White Perch populations may structure those populations through density-dependent mechanisms (Sheri and Power 1972; Schaeffer and Margraf 1986; Hurley 1992).

White Perch are widely distributed in North Carolina reservoirs, and the population abundance and size structure of White Perch vary among these systems (Wong 2002; Feiner et al. 2012). Anecdotal evidence indicates that White Perch attain a large adult body size in some reservoirs. Conversely, White Perch can also dominate reservoir fish communities via high densities of small individuals. A better understanding of the mechanisms driving this variability in abundance and size structure is important for managing present and future introductions of White Perch. Thus, we quantified the abundance and size structure of White Perch in four small North Carolina reservoirs.

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and compared life history traits among the populations. We examined a variety of potential mechanisms to explain the disparity in White Perch body size across these reservoirs, including growth, timing of maturity, trophic interactions, lake productivity, and predator and prey abundances. By including many abiotic and biotic variables in our analysis, our intention was to take a comprehensive approach to understanding White Perch population dynamics in small southeastern reservoirs.

**METHODS**

**Study reservoirs.**—We sampled four reservoirs ranging from 137 to 662 ha in North Carolina’s Piedmont region, including two systems with relatively high-abundance White Perch populations and two systems with low-abundance populations based on unpublished data from the North Carolina Wildlife Resources Commission (NCWRC). Lake Holt (Granville County; 137 ha) and Lake Reece (Randolph County; 242 ha) had low-abundance White Perch populations; Oak Hollow Lake (Guilford County; 280 ha) and Lake Townsend (Guilford County; 662 ha) had high-abundance White Perch populations. All four systems were municipal reservoirs with water levels that were stable throughout the study period. White Perch have been established in each of the reservoirs for over 30 years, but their origin is unknown except in Lake Townsend, where they were introduced intentionally by an angler during the 1970s (S. Davis, City of Greensboro, personal communication).

**Fish collection.**—Fish collection took place in April, August, and October 2010 and April, June, and July 2011. We collected White Perch, Largemouth Bass *Micropterus salmoides*, and Gizzard Shad *Dorosoma cepedianum*. Largemouth Bass were chosen because they are the most common top predator in all of our study systems; Gizzard Shad were selected because they can significantly impact fish communities and may be an important source of prey for piscivores (e.g., Stein et al. 1995). In each reservoir, we collected fish during one 24-h period of each sampling month using nighttime electrofishing and gill-netting at four randomly selected sites distributed throughout the reservoir. Sampling took place at the same sites on each sampling date. Electrofishing began at least 0.5 h after sunset and was conducted in 1,200-s (about 200-m) shoreline transects at each site at an amperage output of 3.8–4.8 A. Gill nets were set perpendicular to the shoreline in at least 2.5 m of water 1 h before sunset and were soaked for 6 h. At two of the four gill-net sampling sites, we set a 2-×-50-m net with 25.4-mm bar mesh. At the other two sites, we set a 2-×-100-m net consisting of two 50-m panels (50.8- and 63.5-mm bar mesh). In the summer months, when much of the lower water column was hypoxic in each reservoir, we suspended gill nets at a depth that ensured placement in habitat with a dissolved oxygen (DO) concentration of at least 3 mg/L.

On each sampling date, we kept a maximum of 20 White Perch per gear and site from each of four size categories (<120, 120–179, 180–229, and >230 mm TL) and placed them on ice for laboratory processing. Length categories for White Perch were based on *a priori* information from previous collections conducted by the NCWRC. In June and July 2011, we also kept White Perch and Largemouth Bass for stable isotope analysis (SIA) and stomach content analysis (Table 1). Largemouth Bass from two size categories (<200 and ≥200 mm TL) were sampled from each study system using nighttime electrofishing and gill-netting. In the summer months, when much of the lower water column was hypoxic in each reservoir, we suspended gill nets at a depth that ensured placement in habitat with a dissolved oxygen (DO) concentration of at least 3 mg/L.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Lake Holt</th>
<th>Lake Reece</th>
<th>Oak Hollow Lake</th>
<th>Lake Townsend</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Small WHP</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Trophic level</td>
<td>3.6 (0.04)</td>
<td>3.7 (0.06)</td>
<td>4.1 (0.08)</td>
<td></td>
</tr>
<tr>
<td>TL (mm)</td>
<td>125 (9)</td>
<td>145 (2)</td>
<td>115 (12)</td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>13</td>
<td>6</td>
<td>10</td>
<td></td>
</tr>
<tr>
<td><strong>Large WHP</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Trophic level</td>
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<td>4.1 (0.02)</td>
<td>3.9 (0.07)</td>
<td>4.0 (0.10)</td>
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<td>TL (mm)</td>
<td>182 (4)</td>
<td>207 (5)</td>
<td>155 (5)</td>
<td>177 (7)</td>
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<tr>
<td>n</td>
<td>4</td>
<td>11</td>
<td>3</td>
<td>5</td>
</tr>
<tr>
<td><strong>Small LMB</strong></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Trophic level</td>
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<td>4.2 (0.06)</td>
<td>3.8 (0.06)</td>
<td>4.4 (0.32)</td>
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<tr>
<td>TL (mm)</td>
<td>123 (13)</td>
<td>99 (10)</td>
<td>105 (11)</td>
<td>123 (12)</td>
</tr>
<tr>
<td>n</td>
<td>12</td>
<td>15</td>
<td>10</td>
<td>3</td>
</tr>
<tr>
<td><strong>Large LMB</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Trophic level</td>
<td>4.2 (0.04)</td>
<td>4.3 (0.03)</td>
<td>4.0 (0.09)</td>
<td>4.6 (0.11)</td>
</tr>
<tr>
<td>TL (mm)</td>
<td>420 (17)</td>
<td>386 (7)</td>
<td>413 (13)</td>
<td>434 (22)</td>
</tr>
<tr>
<td>n</td>
<td>6</td>
<td>10</td>
<td>8</td>
<td>8</td>
</tr>
</tbody>
</table>
on stock length (Gabelhouse 1984) were retained for analyses. For each gear at each site, we weighed (g wet weight), measured (mm TL), and released up to 20 additional White Perch and Largemouth Bass. Gizzard Shad were counted and weighed in aggregate. All other collected fish were counted and released.

Prey collection.—We collected zooplankton and benthic invertebrates from each reservoir in August and October 2010 and April–July 2011 to assess prey availability. On each sample date, we collected zooplankton at three sites by using a 1.5-m conical net with a 0.5-m-diameter opening and 250-μm mesh. We sampled zooplankton from the littoral zone using a vertical tow from twice the Secchi depth to the surface; samples were preserved with Lugol’s iodine solution. We sampled benthic invertebrates by using a petite Ponar dredge at four sites throughout the littoral zone of each reservoir; an area of 0.023 m² was sampled at each site, and sediment was rinsed through a 1,000-μm-mesh wash bucket. Benthos samples were preserved in ethanol and stained using rose Bengal. Additional benthos samples were taken in August and October 2011 and were frozen (not preserved with ethanol) for baseline SIA (see below). Frozen benthos samples were later thawed, and chironomid larvae were separated from the sediment and refrozen until processed for SIA.

Abiotic measurements.—We measured DO (mg/L) and temperature (°C) on each sampling date using a Quanta Hydrolab (Hydrolab Corp., Loveland, Colorado) at the deepest accessible part of each reservoir. We measured specific conductivity (mS/cm) 1 m below the water’s surface and 1 m above the reservoir bottom and then averaged the two values for comparisons. Temperature and DO concentrations were measured at 1-m intervals from the surface to the bottom; epilimnetic measurements (top 4 m of the water column, not including the surface) of temperature and DO were averaged and compared across systems. Secchi depth (m), which was used as a proxy for turbidity and productivity within each reservoir, was measured with a Secchi disk. All measurements were taken at least 2 h prior to sunset on each sampling date.

Laboratory methods.—Sampled fish were frozen at −20°C until laboratory processing. We used macroscopic visual inspection of the gonads to determine sex and stage of maturity for all White Perch collected in the April samples. Stage of maturity was determined using a four-point scale (1 = immature; 2 = developing; 3 = mature; 4 = spent) adapted from Núñez and Duponchelle (2009). Fish with a score of 3 or 4 were considered mature; in contrast to Núñez and Duponchelle (2009), we did not further subdivide scores for female fish. Dorsal muscle tissue (~1 g) was removed from White Perch and Largemouth Bass sampled in June and July 2011 and was frozen for SIA (Table 1). Sagittal otoliths were removed, sectioned, and aged independently by two readers. If the two readers disagreed, the otolith was examined by a third reader; in all cases, the age determined by the third reader was in agreement with the age estimated by one of the first two readers, and that age was assigned to the fish.

Benthic invertebrates and zooplankton were identified to family and enumerated by using a dissecting microscope (zooplankton were subsampled; at least 5% of the sample volume was examined). Chironomid larvae were the most common benthic invertebrate (present in >90% of benthos samples) and are also known to be an important component of White Perch diets (Gosch 2010b; Feiner et al. 2012). Therefore, we used mean densities of chironomid larvae (number/m²) in data analysis as a measure of available benthic prey. For each sampling date, we averaged the total density of all zooplankton (number/m³) collected from each site.

Diet and stable isotope analysis.—Frozen muscle tissue and chironomid larvae were sent to the Stable Isotope Laboratory at Cornell University, where the samples were dried with a Viris Freezemobile 25SL freeze dryer, homogenized with a Spex Cer- tiPrep 6750 freezer/mill, weighed to the nearest 1 mg with a Sartorius MCS microbalance, and analyzed for nitrogen stable isotope ratios (δ¹⁵N) with a Finnigan MAT Delta Plus mass spectrometer. Stable isotope values were compared with the atmospheric nitrogen standard. Measured δ¹⁵N values were standardized relative to the baseline for each reservoir and were used to calculate trophic level (Vander Zanden and Rasmussen 2001), with benthic invertebrates assumed to have a trophic level of 2 (Vander Zanden and Rasmussen 2001; Post 2002). For both Largemouth Bass and White Perch, δ¹⁵N values measured in June and July were not significantly different ($F_1, >50 < 0.07$, $P > 0.08$), so for each species the SIA results from June and July were grouped for analysis. Grouping of SIA samples taken within a month of each other is acceptable because the turnover rate of stable isotopes in fish muscle tissue is more than 30 d (Buchheister and Latour 2010; Weidel et al. 2011).

Conversely, due to the fine time scale of stomach content sampling, we analyzed the diet data from June and July separately. Prey were grouped into five categories: benthic invertebrates, fish, suspended invertebrates, zooplankton, and other. Each prey group’s contribution to the diet of each size-group within each species was quantified using the index of relative importance, an integrated measure of diet that incorporates prey weight, number, and frequency of occurrence (Cortés 1997). Diet overlap was then determined among size-classes and species by using Schoener’s overlap index ($D$; Schoener 1968); Schoener’s $D$ values greater than 0.6 were considered to indicate ecologically significant diet overlap (Wallace 1981). The number of stomach content samples from White Perch caught in June was insufficient for comparison; therefore, Schoener’s $D$ was only used to compare July diets.

Data analysis.—We used ANOVA to compare White Perch and Largemouth Bass CPUE, individual size, and trophic level among reservoirs. We divided White Perch and Largemouth Bass into two size categories for analysis based on the onset of piscivory (Mittelbach and Persson 1998). We calculated the mean CPUEs of White Perch, Largemouth Bass, and Gizzard Shad by summing the total number of fish from each species caught with both gears at each site and then averaging among
sites sampled on each date. We estimated White Perch growth by fitting length and age data from each lake to the von Bertalanffy growth equation (von Bertalanffy 1938). We used the Kruskal–Wallis extension of the Wilcoxon–Mann–Whitney test, based on the normal (Z) distribution to compare White Perch length frequency distributions across populations. Using linear regression, we tested six variables to assess whether there was a relationship with White Perch CPUE: Largemouth Bass CPUE, Gizzard Shad CPUE, mean density of chironomid larvae, mean total zooplankton density, mean Secchi depth, and mean specific conductivity. Fish CPUE was log_{10} transformed for linear regression analysis. All data analyses were carried out using the Statistical Analysis System version 9.2 (SAS Institute, Cary, North Carolina), with \( \alpha \) set at 0.05.

**RESULTS**

**White Perch Abundance, Size Structure, and Age Structure**

White Perch displayed a gradient of abundance from low to high across the four reservoirs, and CPUE was statistically different among reservoirs \( (F_3, 16 = 18.79, P < 0.0001) \). Size structure differed among the four White Perch populations and was strongly related to White Perch abundance (Figure 1; Table 2). High-abundance White Perch populations had a truncated size structure and smaller maximum size than low-abundance populations \( (Z_1 = 439.2, P < 0.0001) \). Length frequency distributions were not significantly different between the reservoirs with high White Perch abundance \( (Z_1 = -1.37, P = 0.1723) \) but were statistically different between the reservoirs with low White Perch abundance \( (Z_1 = 5.86, P < 0.0001) \). Mean TL of White Perch was significantly lower in high-abundance populations than in low-abundance populations \( (F_1, 3 = 4.41, P < 0.0001) \). Age frequency distributions were generally similar among White Perch populations; maximum age ranged from 9 to 14 years, and there were no missing year-classes in any population from ages 0 to 9.

**White Perch Growth and Maturity**

Growth of White Perch differed among populations; asymptotic maximum length \( (L_\infty) \) estimates from the von Bertalanffy growth function were 60–100 mm larger for the two low-abundance populations (Lakes Holt and Reece) than for the two high-abundance populations (Oak Hollow Lake and Lake Townsend). The growth curves for Lake Townsend and Oak Hollow Lake overlapped, whereas growth estimates were significantly different between Lakes Reece and Holt. White Perch in

![FIGURE 1. Length frequency distributions of White Perch sampled from four North Carolina reservoirs in 2010 and 2011 (n = number of fish sampled). Lakes Reece and Holt have low-abundance White Perch populations; Lake Townsend and Oak Hollow Lake have high-abundance White Perch populations.](image-url)
Lake Holt grew faster and to a larger maximum size than White Perch from Lake Reece. Contrary to differences in growth rate, the timing of maturity was fairly consistent among study populations; at least 78% of males and females from all populations were sexually mature by age 2.

**White Perch Abundance in relation to Biotic and Abiotic Variables**

White Perch CPUE was inversely related to Largemouth Bass CPUE (Table 2). White Perch CPUE was not significantly related to Gizzard Shad CPUE, invertebrate prey densities (chironomids and zooplankton), or abiotic variables (Secchi depth and specific conductivity; Table 2). Temperature and DO measurements were similar among systems except Lake Holt, which had a lower mean temperature than the other reservoirs (Table 2). Only Oak Hollow Lake, which is aerated, did not exhibit a summer oxycline.

Largemouth Bass CPUE was higher ($F_{1,3} = 8.06, P = 0.013$) in reservoirs with low White Perch abundance than in reservoirs with high White Perch abundance (Table 2). Reservoirs with low-abundance White Perch populations had a higher proportion of small Largemouth Bass than reservoirs with high-abundance White Perch populations (Figure 2). Mean TL of Largemouth Bass was higher ($F_{1,426} = 14.23, P = 0.0002$) in Oak Hollow Lake and Lake Townsend than in Lakes Holt and Reece, primarily due to differences in the abundance of small Largemouth Bass in the latter two lakes. Largemouth Bass maximum TL was similar among reservoirs (Figure 2).

**Diets and Stable Isotope Analysis**

There were significant among-reservoir differences in the trophic level of each species and size-group, but differences were not correlated with the abundance of White Perch (Table 1). Trophic level of small White Perch was higher in Lake Reece than in Lake Holt ($F_{1,19} = 8.41, P = 0.0009$; Table 1) but was similar among the other reservoirs. Small Largemouth Bass from Oak Hollow Lake had a lower trophic level than small Largemouth Bass from the other three reservoirs ($F_{1,36} = 6.02, P = 0.0020$). Large Largemouth Bass from Oak Hollow Lake also had a lower trophic level than large Largemouth Bass from the other reservoirs but to a lesser degree, and large Largemouth Bass from Lake Townsend had the highest trophic level ($F_{1,28} = 10.08, P = 0.0001$; Table 1).

We documented only one instance of Largemouth Bass predation on White Perch (in Lake Reece; June 2011) and one instance of White Perch predation on Largemouth Bass (also in Lake Reece; August 2010). We detected significant diet overlap between small Largemouth Bass and both size categories of White Perch in all comparisons that had sufficient sample sizes (Table 3). Large Largemouth Bass exhibited low diet overlap with both size-classes of White Perch in the three reservoirs for which sample size allowed comparisons (Table 3).

**DISCUSSION**

White Perch size structure in our study populations was strongly linked to White Perch abundance; this result suggests the importance of density-dependent mechanisms, consistent with previous investigations (Busch et al. 1977; Zuerlein 1981; Gosch et al. 2010a). Although we can attribute the differences in size structure to intraspecific abundance, the factors affecting White Perch abundance in our study systems remain unclear. Despite differences in growth, White Perch from high-abundance and low-abundance populations occupied similar trophic niches. Furthermore, although early timing of maturity has been linked to decreases in size structure for some White Perch populations (Chizinski 2007; Feiner et al. 2012), maturity timing did not vary among the White Perch populations we studied. In addition, abiotic variables have been useful predictors of White

### Table 2

<table>
<thead>
<tr>
<th>Variable</th>
<th>Lake Holt</th>
<th>Lake Reece</th>
<th>Oak Hollow Lake</th>
<th>Lake Townsend</th>
<th>$P$</th>
<th>$R^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>White Perch CPUE</td>
<td>1.9 (1.6)</td>
<td>23.3 (5.7)</td>
<td>44.1 (20.4)</td>
<td>127.2 (52.4)</td>
<td></td>
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</tr>
<tr>
<td>Largemouth Bass CPUE</td>
<td>22.2 (7.9)</td>
<td>6.8 (2.4)</td>
<td>2.6 (0.9)</td>
<td>2.4 (0.9)</td>
<td>0.032</td>
<td>0.94</td>
</tr>
<tr>
<td>Gizzard Shad CPUE</td>
<td>0.0 (0.0)</td>
<td>26.7 (12.5)</td>
<td>17.6 (8.0)</td>
<td>34.2 (20.2)</td>
<td>0.071</td>
<td>0.87</td>
</tr>
<tr>
<td>Chironomid density (number/m³)</td>
<td>54.6 (17.3)</td>
<td>97.9 (47.6)</td>
<td>78.2 (13.5)</td>
<td>59.6 (18.1)</td>
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<tr>
<td>Zooplankton density (number/m³)</td>
<td>9.7 × 10⁴</td>
<td>2.1 × 10⁵</td>
<td>1.8 × 10⁵</td>
<td>1.8 × 10⁵</td>
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<td></td>
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<td>Secchi depth (m)</td>
<td>1.7 (0.2)</td>
<td>1.1 (0.2)</td>
<td>1.0 (0.1)</td>
<td>0.9 (0.1)</td>
<td>0.196</td>
<td>0.65</td>
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<tr>
<td>Specific conductivity (mS/cm)</td>
<td>0.06 (0.02)</td>
<td>0.10 (0.01)</td>
<td>0.11 (0.01)</td>
<td>0.10 (0.01)</td>
<td>0.228</td>
<td>0.60</td>
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<td>Temperature (°C)</td>
<td>15.5 (1.2)</td>
<td>21.1 (1.3)</td>
<td>23.5 (1.4)</td>
<td>20.6 (1.0)</td>
<td>0.194</td>
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<td>DO (mg/L)</td>
<td>8.0 (2.0)</td>
<td>7.2 (0.4)</td>
<td>7.7 (0.5)</td>
<td>8.7 (0.4)</td>
<td>0.708</td>
<td>0.85</td>
</tr>
</tbody>
</table>
Perch abundance in previous studies (Hawes and Parrish 2003; North and Houde 2003), but we found no relationship between White Perch abundance and Secchi depth, specific conductivity, temperature, or DO concentration.

Other organisms in the reservoirs had mixed or nonexistent relationships with White Perch abundance. Studies have linked prey availability to population dynamics in fish (e.g., Deelder 1951; Jansen and MacKay 1992), but in the present study White Perch abundances were not related to invertebrate densities. Gizzard Shad abundance had a marginally significant positive relationship with White Perch abundance, providing weak evidence that the abundances of the two species may be related. High abundances of Gizzard Shad may alleviate predation pressure on White Perch if piscivorous fish select Gizzard Shad over White Perch (Aday et al. 2003; Gosch et al. 2010a).

White Perch abundance has been directly linked to predation (Margulies 1990; Hartman and Margraf 1992; Vrtiska et al. 2003). Gosch and Pope (2011) found that predation may increase stunting in White Perch populations if large predators remove older, larger (rather than younger, smaller) individuals from the population, thereby decreasing the overall size structure. Contrary to this pattern, Largemouth Bass in the present study did not consume many White Perch. Our study focused on Largemouth Bass because they were the most abundant predator in all four study systems, but other piscivores may also interact with White Perch (Schaeffer and Margraf 1987; Prout et al.

### TABLE 3. Schoener’s overlap index ($D$) values based on percent index of relative importance and trophic level of small White Perch (WHP; <150 mm TL), large WHP ($\geq$150 mm TL), small Largemouth Bass (LMB; <200 mm TL), and large LMB ($\geq$200 mm TL). Schoener’s $D$-values greater than 0.6 (in bold italics) were considered to indicate significant diet overlap. A dash (–) signifies that the sample sizes were insufficient ($n < 5$) for analysis.

<table>
<thead>
<tr>
<th>Lake</th>
<th>WHP size</th>
<th>Small LMB</th>
<th>Large LMB</th>
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<tbody>
<tr>
<td>Holt</td>
<td>Small WHP</td>
<td>–</td>
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</tr>
<tr>
<td></td>
<td>Large WHP</td>
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<td>–</td>
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<tr>
<td>Reece</td>
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</tbody>
</table>
Our ability to draw inferences about trophic interactions was limited by a lack of historical data and the coarse time scale at which we collected data on prey availability. Without data describing the state of Largemouth Bass populations before the introduction of White Perch, it is difficult to interpret whether Largemouth Bass abundances have shaped White Perch abundances (or vice versa). It is also possible that more intensive sampling effort (e.g., bimonthly invertebrate sampling throughout the year) could permit the detection of important seasonal differences in prey abundance that affect White Perch abundance at key ontogenetic stages. Furthermore, data from a greater number of reservoirs would have strengthened our ability to draw conclusions. Despite these limitations, we were able to detect a strong density-dependent response in White Perch growth and an interesting relationship between White Perch and Largemouth Bass abundances.

Overall, density-dependent mechanisms drove differences in size structure among the White Perch populations we studied, and our results suggest that White Perch density is a function of Largemouth Bass density. Although we evaluated a suite of common factors that are widely assumed to drive fish abundance, the factors ultimately affecting White Perch abundance in these small reservoirs remain unclear. More research into the competitive and predatory relationships between Largemouth Bass and White Perch, especially at the juvenile stage, may provide insight into the mechanisms that determine White Perch abundance and thus size structure.

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