Recruitment of striped bass in Chesapeake Bay: spatial and temporal environmental variability and availability of zooplankton prey

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ABSTRACT: Causes of recruitment variability in young-of-the-year (YOY) striped bass Morone saxatilis from Chesapeake Bay were investigated based on (1) surveys from 2001 to 2003 to document spatio-temporal variability in abundance of larval striped bass, zooplankton prey, and feeding success of larvae; (2) a synthetic analysis (1996, 1998, 1999, 2001 to 2003) to describe how environmental factors and prey affect recruitment success; and (3) a 10 yr analysis (1993 to 2002) of inter-annual differences in spatial and temporal patterns of copepods and cladocera eaten by striped bass larvae. Striped bass YOY recruitment levels varied >11-fold in the 6 years examined. In those years, mean daily freshwater flows from the Susquehanna River to the bay in March and April varied >2-fold and controlled distribution and apparent survival of striped bass larvae. Strong recruitments of YOY striped bass were associated with matches in space and time of larval striped bass and high concentrations of zooplankton prey, especially the copepod Eurytemora affinis and cladoceran Bosmina longirostris. The strongest year classes (1996, 2003) were produced in years of high freshwater flow, characterized by a high abundance of feeding-stage larvae and a spatio-temporal match between peak abundance of larvae and zooplankton prey. Enhanced feeding opportunities were most pronounced in high freshwater-flow years (1996, 1998, 2003), when larvae and zooplankton prey were strongly associated with, and apparently retained near, the estuarine turbidity maximum. First-feeding larvae fed more successfully in a high-flow year (2003; prey incidence 91%) than in a drier year (2001; prey incidence 35%). A regression model that may have forecasting potential was developed to describe recruitment of YOY striped bass for the years from 1985 to 2006. The model includes spring freshwater flow and air temperatures to predict age-0 striped bass recruitment strength (R² = 0.65). Flow and temperature control environmental and hydrographic conditions that strongly influence spatio-temporal overlap of larval striped bass and zooplankton. The model provided accurate recruitment forecasts for 2007 and 2009, but was less successful in 2008, a year of exceptionally low recruitment.

KEY WORDS: Chesapeake Bay · Striped bass · Recruitment variability · Larval fish · Zooplankton · Trophodynamics · Biophysical interactions

INTRODUCTION

Research on recruitment variability has emphasized the connection between larval fish survival and prey availability since Hjort (1914) proposed a critical period at the onset of larval feeding as the determinant of year-class strength. The match–mismatch hypothesis refined the critical period hypothesis by incorporating the degree of temporal overlap between the spring zooplankton bloom and larval production as a critical determinant of recruitment strength (Cushing 1990). Together, these hypotheses have motivated research on recruitment variability in fishes for decades. Implicit in both hypotheses is the recognition that higher prey concentrations lead to enhanced encounter rates between larvae and prey,
leading to higher larval growth rates and increased larval survival (Houde 2008).

Spatial variability in the prey available to larvae, while not explicitly formalized in critical period and match–mismatch hypotheses, is recognized as a determinant of growth and survival. Feeding conditions of larval fish across mesoscale (>1 to 100 km) gradients in prey concentrations are often attributable to prevailing circulation patterns, frontal features, and interacting spatial dynamics of predators and prey (Munk 2007). A spatial mismatch with negative consequences for growth and survival may occur if peak concentrations of larvae and prey do not coincide (Chick & Van Den Avyle 1999a). Despite supportable hypotheses and tests to link year-class strength of fishes and prey availability, challenges result from contrasts between the spatial and temporal scales over which fish larvae perceive and react to prey and the scales at which they are surveyed (Pepin 2004).

For striped bass <i>Morone saxatilis</i>, abundance of age-0 juveniles (i.e. young-of-the-year [YOY] recruitment level) is strongly associated with the abundance of larval survivors at from 8 to 10 mm length (Uphoff 1989, Rutherford et al. 1997), indicating that dynamics in the larval stage coarsely govern recruitment success in Chesapeake Bay. In a synthesis of larval mortality and growth conducted on 5 annual cohorts of striped bass larvae (Houde 1996, 1997), dynamics of cohort production in the bay were variable and sensitive to environmental conditions. Cohorts that accumulated biomass while in the smallest size classes had a higher potential for strong YOY recruitment.

Links between larval striped bass growth, survival, and prey availability have been investigated in field (Setzler-Hamilton et al. 1987, Rutherford & Houde 1995, Limburg et al. 1999), laboratory (Eldridge et al. 1982, Tsai 1991), and modeling (Logan 1985, Cowan et al. 1993) research. Growth and survival were positively associated with zooplankton when compared across locations in Lake Marion, South Carolina (Chick & Van Den Avyle 1999a). In the Hudson River, striped bass larvae co-occurring with a bloom of the cladoceran <i>Bosmina frenyi</i> showed higher survival rates than larvae temporally mismatched with prey (Limburg et al. 1999). There is evidence indicating that prey availability is important in shaping year-class success in Chesapeake Bay striped bass, although most is correlative, with years of high prey abundance often coinciding with high recruitment (Rutherford et al. 1997). An individual-based modeling study on egg-, larva-, and juvenile-stage striped bass in the Potomac River tributary of Chesapeake Bay supported the hypothesis that variability in prey availability to larvae could generate 10-fold recruitment variability (Cowan et al. 1993).

A relationship between meteorological conditions and Chesapeake Bay striped bass recruitment strength has been recognized for decades (Merriman 1941). In the 1970s, it was postulated that there is a positive association between cold conditions in the spring and recruitment of anadromous fishes (Mihursky et al. 1981). Other research on recruitment variability in Chesapeake Bay striped bass evaluated temperature and precipitation as factors affecting the survival of weekly egg and larval cohorts (Ulanowicz & Polgar 1980, Rutherford & Houde 1995, Secor & Houde 1995, McGovern & Olney 1996). Subsequently, it was demonstrated that frequencies of favorable or unfavorable climate patterns control spring temperatures and precipitation, as well as the abundance and types of prey available to anadromous fish larvae and juveniles in Chesapeake Bay, including striped bass (Wood 2000, Kimmel & Roman 2004). North & Houde (2003) reported higher concentrations of fish larvae and zooplankton prey in the estuarine turbidity maximum (ETM), a convergence zone at the head of many coastal plain estuaries (Schubel 1968), in a wet year when the ETM feature was enhanced.

Our research was conducted in the upper Chesapeake Bay, in the region of the salt front and ETM (Sanford et al. 2001). In Chesapeake Bay and the St. Lawrence River, there is evidence of a link between the survival of anadromous fish larvae, e.g. striped bass <i>Morone saxatilis</i> and rainbow smelt <i>Osmerus mordax</i>, and the dynamics and structure of the ETM and salt front (Dodson et al. 1989, Sirois & Dodson 2000a,b, North & Houde 2003, 2006). In Chesapeake Bay, the ETM is a favorable nursery for striped bass and larvae of the congeneric white perch <i>Morone americana</i> (North & Houde 2001, Shoji et al. 2005), possibly because high concentrations of the copepod <i>Eurytemora affinis</i>, a common prey of these larvae, are found in and near the ETM (Boynton et al. 1997, North & Houde 2001, Roman et al. 2001).

Mechanisms that support strong striped bass recruitment in years characterized by wet, cool, spring conditions are still poorly understood. Here, we report on climatic and biophysical controls of recruitment variability, emphasizing temporal and spatial variability in processes controlling the inter-annual prey availability to striped bass larvae. Specifically, we analyzed the distribution of striped bass larvae in the upper Chesapeake Bay relative to environmental gradients and the spatial and temporal variability in zooplankton concentrations in years of contrasting hydrological conditions. Our analysis included new data on the distribution and feeding success of striped bass larvae for the years from 2001 to 2003; a retrospective analysis of larval and zooplankton data from 1996, 1998, and 1999; and a synthetic analysis on annual variability in the
Martino & Houde: Recruitment of striped bass abundance and timing of zooplankton prey of striped bass larvae for the years from 1993 to 2002. Additionally, we developed a simple statistical model driven by freshwater flow and temperature that describes the level of YOY striped bass recruitment.

Our 4 objectives were to demonstrate how hydro-climate variability (1) controls the distribution of striped bass larvae in the upper Chesapeake Bay; (2) facilitates spatio-temporal overlap of larvae with mesozooplankton prey; (3) enhances or inhibits larval feeding success; and (4) contributes to variability in larval growth, survival, and year-class strength. The research addresses the broad question: ‘Does availability of mesozooplankton prey control striped bass recruitment in the upper Chesapeake Bay?’

MATERIALS AND METHODS

Research surveys. Survey cruises in the upper Chesapeake Bay were conducted during April and May of 1996, 1998, 1999, and 2001 to 2003 (Fig. 1). Prior to ichthyoplankton sampling in each cruise, the location of the ETM and salt front were determined from a CTD survey by plotting salinity and turbidity profiles at 10 km intervals. The ETM’s location and center were defined as the geographic range (typically 5 to 10 km) and its mid-point location, respectively, of maximum turbidity. Sampling stations were selected to map ichthyoplankton abundance up-estuary, within, and down-estuary of the ETM and salt front (Fig. 1). A CTD cast to profile temperature, salinity, and turbidity preceded ichthyoplankton sampling at each station. Station depths ranged from 11 to 13 m.

Ichthyoplankton was sampled with an opening-closing, 1 m² Tucker trawl, with 280 µm mesh, fished at 3 discrete depths in 2001 and 2002 and 2 depths in 2003, to include the entire water column (Table 1). The volume of water filtered by a 2 min tow in each depth stratum was, on average, 126 m³. Samples were preserved in 95% ethanol. In the laboratory, striped bass

<table>
<thead>
<tr>
<th>Year</th>
<th>No. of samples</th>
<th>Eggs (µg/m³)</th>
<th>Yolk-sac larvae (no. m⁻³)</th>
<th>Feeding-stage larvae (no. m⁻³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2001</td>
<td>130</td>
<td>0.84 (1.38)</td>
<td>0.05 (0.09)</td>
<td>0.02 (0.05)</td>
</tr>
<tr>
<td>2002</td>
<td>126</td>
<td>0.68 (2.23)</td>
<td>0.03 (0.06)</td>
<td>1.50 × 10⁻³ (0.01)</td>
</tr>
<tr>
<td>2003</td>
<td>65</td>
<td>0.05 (0.11)</td>
<td>0.90 (1.67)</td>
<td>3.70 (6.93)</td>
</tr>
</tbody>
</table>

Table 1. *Morone saxatilis*. Number of Tucker-trawl samples, and mean (SD) concentrations (no. m⁻³) of striped bass eggs, yolk-sac larvae, and feeding-stage larvae in the upper Chesapeake Bay in 2001, 2002, and 2003.

Fig. 1. Research area and ichthyoplankton and zooplankton sampling stations in Chesapeake Bay. Tucker-trawl sampling stations during May cruises are shown explicitly on the map for 2001 to 2003; station locations in other years were similar. (+) Mesozooplankton monitoring stations of the Chesapeake Bay Program (www.chesapeakebay.net) that were used in the present study (NB Stn 4.3C is located further down-estuary at 38.56°N, 76.43°W). Ellipse demarcates the general location of the estuarine turbidity maximum (ETM) for the years 2001 to 2003.
*Morone saxatilis* eggs and larvae were identified, enumerated, and expressed as number per cubic meter. Mean concentrations of larvae among years were compared using the non-parametric Kruskal-Wallis test, with pairwise multiple comparisons testing for significant ($p < 0.05$) differences with appropriate p-value adjustments.

Distributions of striped bass eggs and larvae were analyzed by pooling ichthyoplankton data from all surveys in each year to evaluate average annual distribution patterns. This analysis was motivated by knowledge of the strong positive relationship between the mean freshwater discharge in spring months and the year-class strength (Houde & Secor 1996, North & Houde 2001), which suggested that the effects of inter-annual variability in flow surpassed the effects of within-year, seasonal flow variability. Except for 1996, when sampling of eggs was insufficient, egg and larval concentrations were averaged in 5 km bins and standardized as $z$-scores (residual divided by standard deviation) to delineate stage-specific centers of abundance.

**Zooplankton spatio-temporal distribution and overlap with larvae.** Inter-annual and spatial variability in distributions of zooplankton were analyzed in the upper bay. Most analyses were on 2 dominant prey of striped bass larvae, the calanoid copepod *Eurytemora affinis* and the cladoceran *Bosmina longirostris* (Beaven & Mihursky 1980, Limburg et al. 1997, Campfield 2005). Additionally, spatial (1996, 1998, 1999, 2001 to 2003) and temporal (1993 to 2002) patterns in the combined concentrations of these taxa were analyzed and compared to minimum prey thresholds for striped bass larvae. Minimum prey thresholds were defined as prey concentrations below which sharp declines in larval striped bass survival and growth occurred in 3 laboratory studies. One study tested *Artemia* nauplii as prey (Eldridge et al. 1981), one tested *Eurytemora affinis* nauplii and other life stages (Tsai 1991), and one used a natural assemblage of zooplankton (Chick & Van Den Avyle 1999b). Local aggregations of zooplankton prey due to small-scale physical processes at spatial scales below the resolution of survey data analyzed here (1 to 10 km) may enhance feeding success (MacKenzie et al. 1990). The prey thresholds we adopted are coarse, but conservative, minimum prey levels above which most larvae are expected to feed successfully.

Inter-annual trends in spring zooplankton along the channel axis in the upper bay were evaluated from samples taken at 4 monitoring stations by the Chesapeake Bay Program (CBP; CB-Program 2002) (Table 2). These stations were sampled approximately once in April and March and twice in May in all years. The CBP zooplankton monitoring was terminated in 2002; consequently, zooplankton analysis for 2003 was based on abundance estimated from high-frequency acoustics sampling (see below). Three of the four CBP stations (Fig. 1) were located within the primary striped bass nursery, and one (CB4.3C) was located 10 to 15 km down-estuary. The CBP zooplankton was sampled in duplicate, stepped-oblique tows, with paired 20 cm bongo nets (202 µm mesh). *Eurytemora affinis* copepodes and adults, immature and adult *Bosmina longirostris*, and the sum of these prey types were standardized to the number of individuals per cubic meter of water sampled.

In 2003, a Tracor acoustic profiling system (TAPS) provided estimates of zooplankton abundance and distribution. The TAPS was mounted on the CTD. TAPS records depth and acoustical volume back-scattering strength (Sv) at 6 transducer frequencies (265, 420, 700, 1100, 1850, and 3000 kHz) (Holliday & Pieper 1995). We analyzed TAPS zooplankton between 0.5 and 1.4 mm, or the equivalent spherical radii (ESR) of from 0.06 to 0.25 mm, which includes the size range of zooplankton eaten by striped bass larvae. TAPS pro-

<table>
<thead>
<tr>
<th>Data source</th>
<th>Temporal period</th>
<th>Spatial coverage</th>
<th>Sampling method</th>
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<tbody>
<tr>
<td><strong>Spatial analyses</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chesapeake Bay Program</td>
<td>April and May 1996, 1998, 1999, 2001, 2002, May 2003</td>
<td>Stns CB2.1, CB2.2, CB3.3C, CB4.3C (see Fig. 1). 50 km at 5–10 km intervals (see Fig. 1)</td>
<td>Bongo net oblique tows (202 µm mesh). Multifrequency acoustic backscatter, TAPS</td>
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<tr>
<td>NSF BITMAX project</td>
<td></td>
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<td></td>
</tr>
<tr>
<td><strong>Temporal analyses</strong></td>
<td>All stations sampled once monthly March–May 1993–2002, Stns CB2.1 and CB2.2 sampled twice each May 1993–2001</td>
<td>Stns CB2.1, CB2.2, CB3.3C (see Fig. 1)</td>
<td>Bongo net oblique tows (202 µm mesh)</td>
</tr>
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</table>

Table 2. Data sources used for analyses of spatial and temporal mesozooplankton variability. The time periods for spatial analyses were selected to coincide with available ichthyoplankton survey data. Time periods for the temporal analyses represent the longest consecutive periods during which there were no major changes in sampling in the upper bay. Chesapeake Bay Program data are available at www.chesapeakebay.net/. TAPS: Tracor acoustic profiling system.
vided estimates of the total prey concentration in the water column (usually ~13 m depth). The estimates of total prey in the designated size range mostly represented *Eurytemora affinis* and *Bosmina longirostris*, based on their numerical dominance in plankton tows from other years.

To evaluate inter-annual and spatial variability in *Eurytemora affinis* and *Bosmina longirostris*, concentration data were fit with a loess smoothing function (span = 0.7, degree = 1). We report mean zooplankton concentrations, based on the loess fits and standard errors of the fits. The loess fits helped to define trends and patterns, as well as their conspicuous peaks in some years.

Loess fits (span = 0.7, degree = 1) of larval striped bass concentrations by distance down-estuary (km) were compared for overlap with zooplankton. For each year, the degree of spatial overlap was evaluated by comparing locations of maximum concentrations of zooplankton and larvae between years. Additionally, mean annual concentrations of zooplankton were calculated for (1) the entire upper bay, (2) the area where striped bass larvae occurred, and (3) the area of maximum larval concentrations. Correlation statistics were derived between mean zooplankton concentration in each of the 3 designated areas during the period of peak production of larval-stage striped bass (April to May) and an age-0 striped bass recruitment index (mean number of YOY striped bass per seine haul in the Maryland Department of Natural Resources seine survey) (Durell & Weedon 2009) to evaluate the relationship between zooplankton prey availability and recruitment success.

The timing of the spring zooplankton peak, based on CBP zooplankton monitoring, and its possible relationship to striped bass recruitment were also evaluated. For this analysis, mean concentrations of *Eurytemora affinis* and *Bosmina longirostris*, and their summed mean concentration in the striped bass nursery area (Table 3), were examined by ordinal day (from March to June) to evaluate inter-annual differences in seasonality.

### Diet analyses.

Diet analysis was conducted on striped bass larvae from collections taken between 2001 and 2003. Standard lengths (SL) of larvae that ranged from 4.0 to 10.5 mm were measured to the nearest 0.1 mm. Gut contents (n = 193) were removed under a dissecting microscope, identified, and measured with an ocular micrometer. In samples with 5 or fewer larvae, gut contents of all larvae were analyzed. In samples with >5 larvae, from 5 to 10 individuals were selected randomly for stomach analysis.

Location and environmental effects on feeding success of larvae were evaluated. Binomial and Poisson regression were applied to test for differences in the percentage of larvae with prey in their gut and the mean number of prey items per gut, respectively. Analysis of covariance (ANCOVA) was applied to compare feeding success among years using larval length as the covariate. At 7.5 mm SL, the yolk sac is fully absorbed and teeth are easily visible, indicating obligate external feeding by this size and ontogenetic stage (Mansueti 1958). Accordingly, larvae were separated into 2 size classes, <7.5 and ≥7.5 mm. For each length class, regression analyses were used to evaluate the effects of salinity, turbidity, and distance down-estuary on mean number of prey items per gut. Significance levels were adjusted for multiple comparisons using Bonferroni’s correction, by multiplying the accepted level of significance (p < 0.05) by 1/n, where n is the number of comparisons.

### Recruitment model.

A multiple regression model was developed to describe and forecast recruitment of age-0 juvenile striped bass. Spring freshwater flow and temperature were included in the model, because

<table>
<thead>
<tr>
<th>Year</th>
<th>All upper bay stations sampled</th>
<th>Stations of overlapping occurrence</th>
<th>Stations of peak larval abundance</th>
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<tbody>
<tr>
<td><strong>Low recruitment</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1999</td>
<td>1672</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>2002</td>
<td>531</td>
<td>606</td>
<td>1935</td>
</tr>
<tr>
<td><strong>Average recruitment</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1998</td>
<td>1299</td>
<td>1090</td>
<td>1802</td>
</tr>
<tr>
<td>2001</td>
<td>7422</td>
<td>2296</td>
<td>275</td>
</tr>
<tr>
<td><strong>Strong recruitment</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1996</td>
<td>7416</td>
<td>14 404</td>
<td>269 263</td>
</tr>
<tr>
<td>2003</td>
<td>22 873</td>
<td>15 877</td>
<td>286 160</td>
</tr>
</tbody>
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spring freshwater flow controls the abundance of zoo-
plankton prey (Kimmel et al. 2006) and temperature is
an important controller of zooplankton post-embryonic
development (Devreker et al. 2006) and striped bass
spawning, growth, and larval survival (Rutherford &
Houde 1995, Secor & Houde 1995). The model was fit
to the recruitment time series for the years from 1985 to
2006 and then run to forecast recruitments for the
years from 2007 to 2009. We evaluated model perfor-
mance with and without zooplankton concentration as
an independent variable for the years from 1985 to
2002 when mesozooplankton monitoring data were
available (CB-Program 2002). Akaike’s information
criterion (Akaike 1974) was used to evaluate good-
ness-of-fit between models with and without zoo-
plankton included.

RESULTS

Abundance of larvae and freshwater flow

Freshwater discharge into the upper Chesapeake
Bay during spring (March and April) varied >2-fold
during the 6 years included in this research (Fig. 2); with highest mean flows in 1996, 1998, and 2003; lowest
flows in 1999 and 2002; and moderate flows in 2001. Concentrations of feeding-stage striped bass Morone saxatilis larvae were significantly different among years (Kruskal-Wallis, df = 5, p < 2.2 × 10^{-16}). Concentrations of larvae were significantly (p < 0.05) higher in 2003 (mean = 3.7 m^{-3}), 1996 (mean = 2.5 m^{-3}), and 1998 (mean = 0.1 m^{-3}), when hydrological conditions were very wet, and when recruitment levels were exceptionally high (1996 and 2003) and moderate (1998). Concentrations of feeding-stage larvae were significantly (p < 0.05) lower in 1999 (mean = 0 m^{-3}) and 2002 (mean = 0.0015), 2 very dry years that experienced poor YOY recruitment.

Diet and feeding success

Striped bass larvae fed almost exclusively on the
cladoceran Bosmina longirostris and the calanoid copepod Eurytemora affinis. All other prey taxa, including the cladocera Daphnia spp., the calanoid copepod Acartia tonsa, and the cyclopoid copepod Oithona sp., were uncommon and occurred in <5% of larval guts. Feeding success was higher in 2003 (2.0 prey larva^{-1}) than in 2001 (0.8 prey larva^{-1}), based on
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In 2001, abiotic factors that we measured had no detectable effect on feeding success of small (<7.5 mm) or large (≥7.5 mm) striped bass larvae. However, in 2003, feeding success of small and large larvae was related to abiotic factors. For small larvae in 2003, feeding on the cladoceran *Bosmina longirostris* was higher up-estuary (p = 6.5 × 10^{-5}; df = 32) and at lower turbidities (p = 9.2 × 10^{-3}; df = 32). Similarly, feeding on *B. longirostris* by large larvae in 2003 was higher up-estuary (p = 2.9 × 10^{-7}; df = 40), at lower salinities (p = 0.01; df = 40), and at lower turbidities (p = 0.02; df = 40). Large larvae in 2003 fed more successfully on the copepod *Eurytemora affinis* at down-estuary stations (p = 5.3 × 10^{-4}; df = 40), at higher salinities (p = 0.01; df = 40).

Overlap of striped bass larvae and zooplankton prey: relationship to YOY recruitment

The distribution of zooplankton serving as prey for larval striped bass is controlled by hydrological conditions. Maximum concentrations of the combined, dominant, upper bay mesozooplankters *Eurytemora affinis* and *Bosmina longirostris* were located further down-estuary (p < 0.01) in years of high freshwater discharge. In dry years, concentrations of *E. affinis* were higher at up-estuary locations (e.g. 1999 and 2002), while in years of high freshwater discharge (1996 and 1998) *E. affinis* peaked down-estuary, at or

![Fig. 3. Morone saxatilis. Inter-annual variability in the average distribution of striped bass eggs and larvae along the main axis of the upper Chesapeake Bay during April and May in 2 dry years (1999 and 2002), an average freshwater discharge year (2001), and 2 wet years (1998 and 2003). ETM mid-point is indicated by a triangle on the x-axis](image)

both number of prey per larva (p = 2.2 × 10^{-12}; df = 189) and prey incidence in guts (p = 4.6 × 10^{-6}; df = 189). In 2002, there were too few (n = 2) feeding-stage larvae available to analyze feeding success.

The most notable difference in diet between years was the greater importance of *Bosmina longirostris* in 2003, the year of high freshwater discharge. The incidence of *B. longirostris* in the guts of larvae increased from <22% in 2001 (n = 102) to 50% in 2003 (n = 89). The number of *B. longirostris* consumed per larva in 2003 was also higher than in 2001 (p = 4.7 × 10^{-11}; df = 189). Feeding on *Eurytemora affinis* did not differ between years.

For the combined years, the number of prey in guts (p = 9.1 × 10^{-9}; df = 189) and the percentage of larvae with prey present in their guts (p = 2.8 × 10^{-6}; df = 189) were directly related to larval size. For individual years, the relationship was significant in 2001 (p = 3.6 × 10^{-8}; df = 100), but not in 2003, when numbers of prey per gut were variable, but relatively high for larvae of all lengths (Fig. 4). In 2001, results indicated that small (<7.5 mm), first-feeding larvae fed less successfully than larger larvae, and, importantly, first-feeding larvae in 2001 fed less successfully than in 2003.

![Fig. 4. Morone saxatilis. Relationship between feeding success and length of striped bass larvae in the upper Chesapeake Bay based on the number of prey items in larval guts in (a) 2001 and (b) 2003. The regression line for the significant relationship in 2001 is shown; the relationship in 2003 was not significant. Horizontal lines in each plot indicate the mean feeding success, i.e. number of prey per larva, for each year](image)
below the salt front and ETM (Fig. 5). B. longirostris occurred in highest concentrations up-estuary of the salt front and ETM in all years. Still, in years of moderate and high freshwater discharge, its distribution shifted down-estuary, coinciding to a greater degree with the ETM (Fig. 5).

The effect of hydrological variability on zooplankton distribution and abundance apparently served to control availability of prey for larvae (Table 3). During the dry years of 1999 and 2002, combined concentrations of Eurytemora affinis and Bosmina longirostris were almost always below the threshold levels required for production of striped bass larvae (ca. 50 000 prey m\(^{-3}\)) at all locations in the upper bay (Fig. 6). In 1999 and 2002, recruitment index values of YOY striped bass in the Maryland Department of Natural Resources seine survey were 1.9 and 1.4 ind. haul\(^{-1}\), respectively, and were well below the long-term mean of 5.5 ind. haul\(^{-1}\).

In 1998 and 2001, when freshwater discharges during March and April were high and near average, respectively, zooplankton concentrations were at moderate levels (Fig. 6) and YOY recruitments of striped bass were moderate (4.3 and 7.2 ind. haul\(^{-1}\), respectively). Zooplankton concentrations reached the minimal prey threshold of 50 000 prey m\(^{-3}\) at some upper bay locations in 1998 and 2001, but most striped bass larvae were located up-estuary of the sites with peak prey concentrations and did not experience favorable feeding conditions (Fig. 6).

In 1996 and 2003, 2 wet, high-discharge years, zooplankton concentrations were notably higher throughout the upper bay (Table 3, Fig. 6). YOY striped bass recruitment indices were also high (15.0 and 11.9 ind. haul\(^{-1}\), respectively). In 1996 and 2003, zooplankton concentrations were far above the threshold prey level (50 000 prey m\(^{-3}\)) at locations supporting the highest concentrations of striped bass larvae and often were above the threshold level at all locations where larvae occurred (Table 3, Fig. 6). Near the ETM, zooplankton concentrations exceeded 100 000 prey m\(^{-3}\). Most importantly, there was high spatial overlap of zooplankton and striped bass larvae in 1996 and 2003 (Fig. 6).

Considering only locations where larval striped bass occurred, the correlation between spring zooplankton concentration and YOY recruitment indices for July through September was significant (p < 0.05, \(r^2 > 0.74\)). Considering all sites sampled for striped bass larvae, correlations between mean spring zooplankton concentrations and YOY striped bass recruitment indices were positive, but not significant (Table 4).

There was substantial temporal variability in occurrence of dominant zooplankton, with probable consequences for synchrony between production of striped bass larvae and prey. The seasonality of Eurytemora affinis differed strongly among years (Fig. 7). Its concentrations usually peaked before May and declined thereafter. Bosmina longirostris concentrations and seasonality differed less among years. A B. longirostris bloom commenced between late April and early May, and it usually peaked in early June or at dates after our surveys had been terminated.

Feeding conditions for striped bass larvae, judged by temporal overlap with zooplankton, vary inter-annually
Peak concentrations of larvae occur from late April through mid-late May (ordinal days 110 to 140) in the upper Chesapeake Bay. There is a strong relationship between the YOY recruitment index and the date of peak zooplankton concentration (p < 0.01, r² = 0.72) (Fig. 8). Aggregate zooplankton concentrations peaked within the 110 to 140 d interval in 4 years (1993, 1994, 1996, and 2001) during the period from 1993 to 2002 (Fig. 8). In most years, zooplankton concentrations peaked earlier, before the third week of April, and then quickly declined. In those years, age-0 recruitment levels of striped bass were generally low (Fig. 8). The notable late-season peaks in spring zooplankton concentrations in 1993 and 1996 were associated with the highest recruitment levels. The degree of temporal overlap and synchrony between combined zooplankton concentrations and striped bass larvae were driven mostly by variability in the seasonality of *Eurytemora affinis*. The overlap was enhanced when *E. affinis* peaked during May rather than in March or April, or when high concentrations of *E. affinis* persisted in May (Fig. 7).

**Recruitment model**

Based on evaluating a suite of abiotic and biotic variables, a multiple regression model for the years from 1985 to 2006 was developed that successfully described recruitment levels of YOY striped bass at 4 mo of age. The evaluation of potential independent variables indicated that only spring freshwater flow (p = 1.4 × 10⁻⁴) and spring temperature (p = 0.01) were significant in the model. The model described recruitment variability reasonably well (R² = 0.65). It also successfully forecasted YOY recruitment levels for 2007 and 2009, but was not successful for 2008 (Fig. 9).
Environment, prey availability, and recruitment

In Chesapeake Bay, recruitment of striped bass *Morone saxatilis* is controlled by both direct and indirect effects of hydrological variability on egg and larval survival. Indirect effects, driven by trophodynamics, distinguish years when recruitments are moderate (e.g. 1998 and 2001) and strong (e.g. 1996 and 2003). Direct effects (e.g. reduced retention of eggs and larvae, possible osmotic stress) and indirect trophodynamic effects are implicated during dry years (e.g. 1999 and 2002) when recruitments are poor, and when abundances of feeding-stage larvae are very low.

Direct effects of hydrological variability in 2 dry years (1999 and 2002) resulted in poor survival of pre-feeding larvae and low abundances of feeding-stage larvae, most of which were located >15 km up-estuary of the salt front and ETM. The low numbers of feeding-stage larvae at the salt front and ETM in these years could have been attributable to an up-estuary shift in location of adult spawning in dry years. Alternatively,
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reduced capacity for retention and increased salinities near the ETM and salt front in dry years are a probable cause of down-estuary loss of eggs and larvae and, thus, of poor survival. This mechanism was proposed to explain the paucity of feeding-stage larvae in 1999 (North & Houde 2001).

Our analysis highlighted how indirect effects of trophodynamics, including spatial and temporal matches and mismatches in occurrences of striped bass larvae and their zooplankton prey, are governed by temperature and hydrological variability before and during the season of egg and larval production. Enhanced spatial-temporal availability of zooplankton prey promotes larval feeding success and also faster growth (Martino 2008) and is associated with higher larval-stage survival. Results reported here indicate that (1) feeding success of striped bass larvae is enhanced in wet, strong recruitment years; (2) larvae and zooplankton prey occur further down-estuary and are more associated with the ETM and salt front in wet, strong recruitment years; and (3) the strongest recruitments occur when spring-season peaks in zooplankton prey occur in May rather than earlier in the year.

Both temporal and spatial matches of prey availability and striped bass larvae are necessary to produce the strongest recruitments. In 1996 and 2003, larvae were at their highest concentrations in the ETM region, where zooplankton prey exceeded 250 000 prey m$^{-3}$, levels far above the reported threshold prey levels for striped bass larvae that range from 50 000 to 100 000 prey m$^{-3}$ (Eldridge et al. 1981, Chesney 1989, Tsai 1991). In contrast, in poor recruitment years, such as 1999 and 2002, mean prey concentrations were approximately 2 orders of magnitude lower (average: <2000 prey m$^{-3}$) throughout the upper bay. In years of low or moderate freshwater flow that have strong temporal overlap between production of striped bass larvae and prey, suitable prey levels can occur at some locations in the upper bay, but generally not where most larvae occur. For example, in 2001, there was a spatial mismatch between striped bass larvae and prey (Fig. 6) that may explain the unexpectedly low recruitment despite a strong temporal match (Fig. 8). In contrast, during the high freshwater-flow year of 1998, larvae and zooplankton tended to co-occur in the ETM, but there was little temporal overlap between peaks in zooplankton concentrations and larval abundance. Similarly, Rutherford et al. (1997) suggested that high and low recruitment years in 1989 and 1988, respectively, had partly resulted from temporal matches and mismatches of prey with striped bass larvae in the upper bay. Although prey matches and mismatches were recognized, Rutherford et al. (1997) believed that the direct effects of temperature on larval growth and survival were of greater importance in those years.

Past research in Chesapeake Bay had identified temperature as a primary factor governing the growth of striped bass larvae (Rutherford & Houde 1995, Secor &

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**Fig. 9. Morone saxatilis.** Observed, predicted, and forecasted recruitment of age-0 juvenile striped bass in the upper Chesapeake Bay. The solid line is age-0 abundance from the Maryland Department of Natural Resources recruitment seine survey (www.dnr.state.md.us/fisheries/juvinventory/index.html). The model to predict recruitment includes Susquehanna River discharge for March through May from the US Geological Survey and air temperatures for March through May from Baltimore–Washington International Airport. The dashed line is modeled age-0 abundance for 1985 through 2006 and forecasted abundance for the 3 most recent years (2007 to 2009).
Houde 1995). While temperature was important in the period from 2001 to 2003, it is probable that interannual variability in feeding success and growth were largely driven by interannual variability in prey availability. A regression relationship describing larval striped bass growth with respect to temperature (Rutherford & Houde 1995) predicted higher growth in 2001 (0.26 mm d\(^{-1}\) at 17.5°C) than in 2003 (0.22 mm d\(^{-1}\) at 15.5°C). However, observed growth rates of small larvae (<12 d old) did not differ in these 2 years, and the growth of larger larvae was faster in 2003 (Martino 2008), demonstrating the benefits of enhanced prey availability in this cool year.

**Spatial and inter-annual differences in diet composition**

Larvae fed more successfully in 2003 than in 2001 and 2002. In 2003, hydrological conditions were very wet and peak concentrations of larvae were within and just up-estuary of the ETM and salt front where *Bosmina longirostris* and *Eurytemora affinis* were abundant. Higher concentrations and strong seasonal variability in the abundance of *E. affinis* suggest that it is a more critical prey resource than *B. longirostris* for striped bass larvae in the upper Chesapeake Bay.

Concentrations of *Bosmina longirostris* rarely exceeded 8000 ind. m\(^{-3}\) in the upper Chesapeake Bay. However, this cladoceran enhances feeding success by larvae where it overlaps spatially during strong recruitment years. In 2003, when recruitment was high, 50% of striped bass larvae had eaten *B. longirostris* compared to 21% in 2001 when recruitment was lower. Feeding on *B. longirostris* was enhanced in freshwater, up-estuary of the ETM and salt front. In contrast, feeding on *Eurytemora affinis* was enhanced further down-estuary at higher salinities. Campfield (2004) also reported higher consumption of *B. longirostris* by striped bass larvae up-river of the salt front and ETM in the Patuxent River, while larvae down-river consumed mostly *E. affinis*. In the Hudson River, *Bosmina freyi* was the dominant food of larvae near the freshwater–salt water interface, where concentrations exceeded 100 000 prey m\(^{-3}\) (Limburg et al. 1997).

**Seasonal timing of zooplankton and recruitment strength**

In the upper Chesapeake Bay, zooplankton concentrations peak most often in March and April. But, most striped bass larvae hatch from late April to early May (Rutherford & Houde 1995) when temperatures rise above 12°C (Secor & Houde 1995), indicating potential for frequent mismatches between prey availability and larval striped bass production. The strongly positive relationship (p < 0.01, r = +0.85) between the day of peak zooplankton concentration and YOY recruitment strength (Fig. 8) indicates that matches occur in years when zooplankton production, especially *Eurytemora affinis*, peaks in May, as observed in 1993 and 1996. Increased precipitation and freshwater flows during late winter and spring are associated with increased abundances of *E. affinis* (Wood 2000, Kimmel & Roman 2004). While *E. affinis* usually peaks in March or April in Chesapeake Bay (Kimmel & Roman 2004), peak concentrations occur or extend into May in years when a climate pattern termed the ‘Ohio Valley High’ (cool and wet) persists through March (Wood 2000) and match the occurrences of peak abundance in striped bass larvae.

The other important prey, *Bosmina longirostris*, peaks more consistently, usually in May and overlaps peak production of striped bass larvae. This temporal consistency in overlapping may ensure that some larvae encounter sufficient prey in all years. In the Hudson River, *B. longirostris* also blooms predictably in late May to mid-June, when its concentration surges from 100 to 100 000 ind. m\(^{-3}\) and striped bass larvae exhibit positive selectivity for this important prey (Limburg et al. 1997). In the Patuxent River, *B. longirostris* is second only to *Eurytemora affinis* in importance as prey, although it is not positively selected by striped bass larvae (Campfield 2004).

**The ETM and recruitment success**

It was hypothesized that freshwater flow controls recruitment by its effect on the overlap of temperature/salinity zones preferred by larvae and the elevated secondary productivity in the ETM (North & Houde 2006). Our results are generally consistent with this hypothesis, although we found that distributions of larval striped bass are not closely coupled to the ETM in all years. In wet years, down-estuary displacement of eggs and yolk-sac larvae enhances the degree of overlap between feeding-stage larvae and the ETM. In the wet year of 2003, larvae tended to occur >20 km further down-estuary and were more associated with the ETM than in the dry year of 2002. Recruitment of YOY juveniles was nearly 9-fold higher in 2003 than in 2002.

The ETM with its associated salt front is a compressed frontal feature in estuaries and, as such, is expected to aggregate prey and potentially enhance feeding opportunities and growth of fish larvae. Frontal features, in general, enhance prey availability, feeding success, or growth of marine and estuarine fish.
larvae (Grimes & Kingsford 1996). There are many examples of such enhancements, e.g. larval cod Gadus morhua (Munk 2007), Japanese sand lance Ammodites personatus (Nagano et al. 2001), dab Limanda limanda (Lee et al. 2007), sprat Sprattus sprattus (Munk 1993), and yellowfin tuna Thunnus albacares (Lang et al. 1994) in tidal, thermohaline, and river plume fronts.

The ETM in Chesapeake Bay appears to support retention and high production of zooplankton eaten by striped bass larvae (Roman et al. 2001, North & Houde 2006). High freshwater flows increase estuarine gravitational circulation (Hetland & Geyer 2004) and may favor higher zooplankton production in addition to retention of particles, including detritus, zooplankton, and larval fish at the salt front and ETM. During wet years, when gravitational circulation is enhanced, Eurytemora affinis abundance increases in Chesapeake Bay (Kimmel et al. 2006) and its center of abundance shifts down-estuary with the ETM. In the 2 wettest years in our analyses (1996 and 2003), there were notable aggregations of zooplankton and striped bass larvae in the ETM and salt front (Fig. 5). In 2003, our diet analysis indicated a high percentage (91%) of successful, first-feeding striped bass larvae. In contrast, under average (2001) or dry (1999 and 2002) hydrological conditions, concentrations of larvae in the ETM were lower, suggesting that the ETM was less important as a nursery habitat. In 2001, only 35% of first-feeding striped bass larvae had fed successfully.

Dynamics and properties of ETMs and salt fronts probably play a role in controlling recruitment of striped bass in other sub-estuaries and tributaries of Chesapeake Bay. Releases of chemically marked striped bass larvae in the tidal Patuxent River at and up-river of the salt front yielded up-river recaptures, but releases below the salt front resulted in no recaptures, suggesting that the ETM/salt front supported retention (Secor et al. 1995). In the tidal Nanticoke River in 1992 and 1993, Houde & Secor (1996) reported that most naturally produced and recaptured hatchery-source, striped bass larvae occurred immediately up-river of the ETM/salt front. Consistent with our analysis of upper bay striped bass in 1993, recruitment in the Nanticoke was >2-fold higher in 1993, when high freshwater flow displaced the salt front down-river, resulting in a larger nursery volume up-river of the front than in a contrasting dry year, 1992.

**Recruitment model**

We developed a relatively simple, predictive model that describes and forecasts striped bass YOY recruitments, based on temperature and hydrological conditions in the upper Chesapeake Bay. The model was successful in describing age-0 recruitment of striped bass ($R^2 = 0.65$), and it successfully forecasted age-0 recruitment in 2 of 3 recent years (Fig. 9). The very poor age-0 recruitment observed in 2008 was substantially lower than that we forecasted, suggesting that important factors were unaccounted for in this year.

We did not include adult stock in our model because spawner biomass alone explains little of the inter-annual variability in striped bass recruitment. A Ricker spawner–recruit model that we fit to upper bay adult biomass data (Maryland DNR gillnet monitoring) for the years from 1985 to 2007 explained only 2% of the variability in YOY recruitment compared to a much higher percentage (46%) explained by a Ricker model including both freshwater flow and adult biomass. In an earlier analysis that included data for the years from 1987 to 1999, Ricker models without and with freshwater flow explained 3 and 44%, respectively, of the variability in recruitment (North & Houde 2003).

Our 2-variable regression model successfully described and predicted recruitment and was helpful in explaining the process. Spring freshwater flow and temperature largely control the seasonal timing of production and abundance of mesozooplankton prey, as well as the degree of spatial overlap between striped bass larvae and prey. The importance of freshwater flow in controlling survival of striped bass larvae in Chesapeake Bay (Houde & Secor 1996, McGovern & Olney 1996, Wood 2000, North & Houde 2001) and elsewhere (Turner & Chadwick 1972, Kimmerer et al. 2001) had been recognized previously. Down-estuary displacement of the salt front and a concomitant increase in volume of freshwater and low-salinity habitat up-estuary of the front have been proposed to explain the positive effects of freshwater on striped bass larvae in the Nanticoke River (Secor & Houde 1996) and in the San Francisco Bay estuary (Kimmerer et al. 2009).

Zooplankton is not included explicitly in our recruitment model, because data on abundance and seasonality of zooplankton were no longer available after 2002 (CB-Program 2002). In the upper Chesapeake Bay, abundance and distribution of the most important prey (Eurytemora affinis) of striped bass larvae, are controlled largely by the level of freshwater flow (Kimmel & Roman 2004, Lloyd 2006). Temperature controls the timing of striped bass spawning and larval production (Secor & Houde 1995) and plays a role in the production of E. affinis (Devreker et al. 2006). Including the mean zooplankton concentration for the period from April to May in the years from 1985 to 2002 as an independent variable in our model slightly improved the model fit (Akaike’s information criterion = 97.2 to 94.6) for the 18 yr recruitment time series when zoo-
plankton data were available. Presumably, a zooplankton variable would improve the model's predictive and forecasting capability if zooplankton data were presently available.

**Management implications**

Our findings, potentially, will find application in management planning for the striped bass fishery. The recruitment model has the capability of forecasting YOY juvenile abundances. The model is underpinned by an enhanced understanding of the mechanisms by which hydrological conditions affect recruitment and is supported by new information on the spatial and temporal processes that control year-class strength of YOY striped bass in Chesapeake Bay.

Disentangling the effects of environmental variability from the effects of adult biomass and demographics on recruitment may contribute to effective management of the striped bass resource. Uncertainty in the stock–recruitment relationships used in striped bass stock assessment models (ASMFC 2008) could be reduced by accounting for environmental effects. Striped bass have the characteristics of a fish stock that could benefit from recruitment-process information in stock assessments, i.e. a strong environment–recruitment relationship and periodic dominance in the fishery by strong year classes (De Oliveira & Butterworth 2005). Ecosystem-based approaches to the management of striped bass and other fisheries are under development for Chesapeake Bay (EBFMCB 2010), drawing more attention to the urgency of understanding how environmental variability influences pre-recruit dynamics. At present, environmental indicators are not included in management planning or reference points for Chesapeake Bay striped bass. Our analysis of spatio-temporal variability in hydrological factors and zooplankton prey availability, and their effects on early life survival, could potentially contribute an important impetus to ecosystem-based plans for the management of striped bass.

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