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Zebra mussels affect benthic predator foraging success and habitat choice on soft sediments

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Abstract The introduction of zebra mussels (Dreissena spp.) to North America has resulted in dramatic changes to the complexity of benthic habitats. Changes in habitat complexity may have profound effects on predator-prey interactions in aquatic communities. Increased habitat complexity may affect prey and predator dynamics by reducing encounter rates and foraging success. Zebra mussels form thick contiguous colonies on both hard and soft substrates. While the colonization of substrata by zebra mussels has generally resulted in an increase in both the abundance and diversity of benthic invertebrate communities, it is not well known how these changes affect the foraging efficiencies of predators that prey on benthic invertebrates. We examined the effect of zebra mussels on the foraging success of four benthic predators with diverse prey-detection modalities that commonly forage in soft substrates: slimy sculpin (Cottus cognatus), brown bullhead (Ameirus nebulosus), log perch (Percina caprodes), and crayfish (Orconectes propinquus). We conducted laboratory experiments to assess the impact of zebra mussels on the foraging success of predators using a variety of prey species. We also examined habitat use by each predator over different time periods. Zebra mussel colonization of soft sediments significantly reduced the foraging efficiencies of all predators. However, the effect was dependent upon prey type. All four predators spent more time in zebra mussel habitat than in either gravel or bare sand. The overall effect of zebra mussels on benthic-

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feeding fishes is likely to involve a trade-off between the advantages of increased density of some prey types balanced against the reduction in foraging success resulting from potential refugia offered in the complex habitat created by zebra mussels.

Introduction

Habitat complexity is an important structuring component of biological communities and may stabilize predator-prey interactions (Murdoch and Oaten [1975;](#page-6-0) Caswell [1978](#page-5-0); Holt [1984](#page-6-0); Taylor [1984;](#page-6-0) McNair [1986\)](#page-6-0). Changes in habitat complexity may affect encounter rates between predators and prey, and predator efficiency (Briand and Cohen [1987](#page-5-0); Whitehead and Walde [1992\)](#page-6-0). Complex habitats may provide prey refuges by reducing predator efficiency in portions of the habitat (Huffaker [1958\)](#page-6-0). Habitat complexity interacts with predator foraging modes and encounter rates so that the number of prey consumed generally decreases with structural complexity (Savino and Stein [1982](#page-6-0), [1989](#page-6-0); Eklöv [1997](#page-5-0); Eklöv and Diehl [1994](#page-5-0); Persson and Eklöv [1995](#page-6-0)). In a number of fish-prey systems, per capita prey-capture rates declined with habitat complexity as search and pursuit times increased (Glass [1971;](#page-6-0) Ware [1973](#page-6-0); Vince et al. [1976;](#page-6-0) Stein [1977;](#page-6-0) Stoner [1982](#page-6-0)).

In freshwater systems, habitat complexity is most commonly associated with littoral vegetation, large-particle substrates, or depth-gradient diversity. Each has profound effects on species abundance and diversity of prey (Werner et al. [1977](#page-6-0); Eadie and Keast [1984\)](#page-5-0). Different forms of habitat complexity provide prey refuges by reducing predation efficiency (Mittelbach [1988](#page-6-0); Werner and Hall [1988\)](#page-6-0). For example, the foraging efficiency of pike increased at low vegetation densities and decreased at high vegetation densities (Savino and Stein [1989\)](#page-6-0).

Zebra mussels have been associated with system-wide changes over virtually all trophic levels. The introduction of zebra and quagga mussels (Dreissena polymorpha and D. bugensis) to North America has significantly altered water clarity (Mellina et al. [1995\)](#page-6-0), chlorophyll *a* levels (Caraco et al. [1997\)](#page-5-0), zooplankton communities (Jack and Thorp [2000](#page-6-0)), and benthic community structure (Ricciardi et al. [1997\)](#page-6-0). In addition to enriching the benthic food web (e.g., Stewart et al. [1998](#page-6-0)), zebra mussels benefit benthic macro-invertebrates by creating structural complexity on previously simple surfaces. Increased complexity provides benthic invertebrates protection from disturbances such as increased wave action (Bially and MacIssac [2000](#page-5-0)) and more importantly, potential refugia from predation (Mayer et al. [2001;](#page-6-0) Cobb and Watzin [2002](#page-5-0); Kolar et al. [2002\)](#page-6-0).

Zebra mussel colonies on soft sediments may reduce fish foraging success by blocking access to the benthos or by reducing the ability of fish to find and capture prey. Soft-sediment macroinvertebrates including burrowing mayflies (Ephemeroptera: Ephemeridae: Hexagenia spp.) and Chironomus (Diptera: Chironomidae) are important dietary components of fish such as suckers (Catostomidae), catfishes (Ictaluridae), cyprinids (Cyprinidae), lake sturgeon (Acipenser fulvescens), and commercially important fishes such as walleye (Stizostedion vitreum) and yellow perch (Perca flavescens; Scott and Crossman [1973](#page-6-0); Beamish et al. [1998](#page-5-0)).

The impact of zebra mussels on fish foraging will likely be related to foraging mode and prey type. Predators such as sculpins (Cottus spp.) that utilize lateral line detection of infaunal prey will likely be impeded by zebra mussels on soft substrates and may switch to visually detected crawling and swimming prey. In contrast, surface "pickers" such as logperch (Percina caprodes) may benefit from increased invertebrate densities in zebra mussel colonies. Zebra mussels may protect burrowing invertebrate prey, whereas free-swimming and crawling invertebrates may find refugia within zebra mussel colonies. Therefore, in order to understand the impact of zebra mussels on foraging efficiencies of benthic fish species, it is necessary to examine multiple predators with different foraging modes and their interaction with different prey types.

We examined the effect of zebra mussels on foraging efficiency of four benthic predators; slimy sculpins, logperch, brown bullheads, and crayfish (Orconectes propinquus). We conducted laboratory experiments to assess foraging success with multiple prey types and different levels of habitat complexity. We also examined the habitat preference of each predator in order to determine how zebra mussels may affect fish and crayfish distributions in addition to foraging success. We discuss our results in relation to foraging modalities, prey type, and habitat complexity.

Materials and methods

Organisms

All predator species were wild-caught in the Lake Champlain drainage and used once for foraging experiments and once for habitat choice before being returned to the wild. The predator species were acclimatized in the laboratory for a minimum of 2 days, placed in experiments, and released within 2 weeks. The predators were fed a diet of frozen blood worms and maintained without food for 24 h before the foraging experiments. Bullheads were collected in minnow traps in the Winooski Reservoirs on Gill Brook in Winooski Township, Vermont, and by electrofishing in Alder Creek in Essex Township, Vermont (total length $\bar{x}=97$ mm, range 61– 140 mm). The crayfish were minnow-trapped and hand picked in the shallows of Lake Champlain (average carapace length from tip of rostrum to the mid-dorsal posterior margin of the carapace=28 mm; range 19– 41 mm). We collected log perch in minnow traps and by snorkeling and hand netting in Lake Champlain (TL $\bar{x}=84$ mm; range $65-120$ mm) The sculpins were electrofished in Lake Champlain and Brown's River in Underhill Vermont (TL \bar{x} = 66 mm; range 48–90 mm).

We reared chironomids in bulk cultures in the laboratory (length range 4–8 mm). Isopods were purchased from Connecticut Valley Biological Supply Company (length range 4–11 mm). Amphipods were collected from springs in Pennsylvania (length range 4–8 mm). We collected zebra mussels from Lake Champlain and washed them free of macroinvertebrates and sediment in a coarse sieve with dechlorinated water prior to their use in the experiments. We used a mixture of live mussels and the shells of dead individuals as is typically encountered in benthic mussel colonies. We added the mussels to the experiments individually and in small clusters. The live mussels laid down byssal threads such that most individual mussels were attached to conspecifics and dead shells before the experiments began.

Foraging success

To test whether zebra mussels affect the foraging success of benthic predators, we conducted a set of separate laboratory experiments with three prey species and four predator species. Each predator-prey combination was evaluated separately for a total of 12 replicated experiments. The number of replicates per experiment depended on predator availability and varied between three and nine (Fig. [1](#page-2-0)). The selected prey represented three modes of habitat use: burrowing (chironomids—*Chironomus* sp.), crawling (isopods—Asellus communis), and swimming (amphipods—Gammarus minus). The predator species use distinctly different modes of foraging and habitat use; lateral line system, visual, and strictly benthic (sculpins), barbels and mostly benthic (bullhead); visual and partly pelagic (logperch); chemotactic, strictly benthic and burrowing (crayfish).

All experiments were conducted in 8-l circular microcosms (27 cm diameter, 14 cm depth). Each microcosm was filled with clean, sifted, washed sand to a depth of 2.5 cm. The microcosms were filled with continuously aerated de-chlorinated water. We established a full factorial design with percent zebra mussel coverage and predator presence/absence as the factors in the design and number of prey items remaining after each experiment as

Fig. 1 Consumption of prey by four benthic predators in enclosures with three levels of zebra mussel coverage of substrate (0, 50, and 95%). The proportion of prey (chironomids, isopods, and amphipods) remaining after 48 (chironomids) or 24 h averaged across replicates are plotted versus the presence and absence of four predators: sculpins, logperch, bullheads, and crayfish. Initial number of prey were 70 for chironomids and 30 for isopods and amphipods for all predators. n number of replicate experiments with each predator

the response variable. This design resulted in the following six treatments: (1) bare substrate; (2) bare substrate and predator; (3) 50% zebra mussel coverage; (4) 50% zebra mussel coverage plus predator; (5) 95% zebra mussel coverage; and (6) 95% zebra mussel coverage plus predator. Percent cover of zebra mussels was estimated visually. This design allowed us to measure the main effects and interactions of fish and zebra mussels on the number of prey remaining.

A statistically significant predator effect would simply confirm that the predator in question consumed the prey item offered under the experimental conditions. A significant effect of zebra mussels would clearly imply that zebra mussels were somehow affecting the prey densities. A significant zebra mussel by predator interaction would confirm that the presence of zebra mussels modified the foraging success of the predators.

The duration of the trial and the number of prey added to each microcosm depended on the prey species. We added 70 chironomids (effective density 1,223 individuals per m²) to each microcosm 24 h prior to the addition of predators to allow the chironomids time to burrow into the substrate. Because amphipods and isopods are substantially larger than chironomids, fewer individuals will likely be consumed to achieve similar predator satiation; we added 30 (effective density 524 individuals per m^2) of either of these species to the microcosms at least 2 h prior to the addition of predators. Predators were allowed to forage on chironomids for 48 h and on isopods or amphipods for 24 h. Upon termination of the experiments,

the predators were removed, measured to the nearest 0.1 mm (standard length for fish, carapace length for crayfish) and weighed (wet weight) to the nearest 0.1 g. Remaining prey were removed from the zebra mussels and sand and counted using sieves and forceps.

Habitat choice

We examined fish preference for habitat type in experimental laboratory microcosms. Bare sand, gravel, and live zebra mussels were arranged in three equal-sized segments in each of six circular 43.2-l tubs (61 cm diameter). Identical depths of all three substrates were used. In contrast to the foraging experiments, the mussels were placed directly on the microcosm floor, as was the gravel. The experimental setups were allowed to sit for 24 h prior to addition of the fish or crayfish. We examined habitat preference of slimy sculpins $(n=12)$, brown bullheads $(n=8)$, logperch $(n=11)$, and crayfish $(n=12)$. Individuals were placed in the tubs and allowed to settle for 4–6 h prior to the start of the experiment. Each tub contained a single individual. We recorded the location of each individual at 10-min intervals. The observer was careful to minimize potential disturbance of the predator, and the predators rarely moved when the observer recorded their position. Observations were conducted during three time periods; dawn (5:30–9:30 a.m.), midday (10:30 a.m.–2:30 p.m.), and dusk (6:30–10:30 p.m.), for a total of 72 observations per individual predator.

Statistical analyses

The foraging experiments were analyzed using a factorial ANOVA with forager presence/absence and zebra mussel density (absent, 50% coverage, and 95% coverage) as the two factors in the analysis. In order to satisfy the assumptions of ANOVA, the response variables (number of prey remaining) were arcsine transformed prior to analysis. We used a Tukey's post hoc analysis to ask if the number of prey differed among the three mussel density treatments. Because the question of interest in these experiments was the effect of zebra mussels on fish foraging, non-predator treatments were not compared. Simple inspection suggested that differences in prey recovery were trivial regardless of zebra mussel presence or density.

Habitat choice experiments were analyzed using G-tests on each individual fish or crayfish for each daily time period. The number of observations in each substrate type was compared to the distribution that would be expected if a fish or crayfish was unbiased in its habitat choice, i.e., it was found in each habitat type for an equal number of observation periods. Data from each individual fish and crayfish were independent from data for other conspecifics; therefore, for each species at a given time period we summed the G -test values and the df to achieve a statistic for all replicates combined.

Results

Foraging success

In general, zebra mussels inhibited the ability of predators to forage successfully. However, the degree to which predators were inhibited by zebra mussels was influenced by prey type and percent coverage of zebra mussels. In most cases predators consumed far more prey in the absence of zebra mussels than in treatments containing zebra mussels. Foraging success also varied among predator species.

Predators tended to consume fewer chironomids over 48 h in treatments containing zebra mussels than in the absence of zebra mussels (Fig. [1\)](#page-2-0). Both zebra mussel coverage and predator presence/absence had significant effects on the number of chironomids remaining after the termination of the experiments in all cases $(P<0.05$, Table 1). There was also an interaction effect between predator species and percent zebra mussel coverage for all predators with the exception of crayfish such that fewer chironomids were consumed in the presence of zebra mussels than in their absence $(P<0.05$, Table 1). Sculpins consumed more chironomids in the treatment lacking zebra mussels than in either the 50 or 95% zebra mussel coverage treatments (Tukey's, $P<0.05$). Logperch and bullheads consumed significantly more chironomids in the treatment lacking zebra mussels than in the 95% zebra mussel coverage treatment; however, there was no difference between the bare and 50% treatments (Tukey's, P<0.05). In contrast to fish-foraging results, zebra mussels had no significant effect on the consumption of chironomids by crayfish. Crayfish foraged successfully regardless of percent zebra mussel coverage (Fig. [1\)](#page-2-0).

On average, the consumption of isopods declined as percent zebra mussel coverage increased (Fig. [1](#page-2-0)). There was a predator effect for all predator species $(P<0.05$,

Table 1 Results of factorial ANOVAs on the effects of zebra mussels on foraging success of each of four predators. Zebra mussel density (0, 50, or 95% coverage), and predator presence or absence were the two factors in each analysis. A significant effect of zebra mussels indicates that the number of prey items recovered from the experimental microcosm after the foraging period differed among zebra mussel treatments. A significant predator effect simply confirms that the predators consumed the prey in question. A zebra mussel by predator interaction effect indicates that foraging success was influenced by the presence of zebra mussels

Table 1). There was an effect of zebra mussel coverage on isopod consumption for sculpins, bullheads, and crayfish $(P<0.05$, Table 1), but not for logperch. Sculpins consumed more isopods in the 0% zebra mussel coverage treatment than in the 95% treatments (Tukey's, $P<0.05$). Bullheads and crayfish consumed more isopods in the absence of zebra mussels than in either the 50 or 95% zebra mussel coverage treatments (Tukey's, $P<0.05$). Regardless of zebra mussel density, sculpins and logperch consumed fewer isopods than did either of the other predators. Sculpins and logperch consumed more chironomids and amphipods than they did isopods. This pattern was very consistent among individual replicates.

All predators consumed more amphipods in the absence of zebra mussels than in either treatment containing zebra mussels (Fig. [1](#page-2-0)). In all cases there was a significant effect of predator on the number of amphipods remaining $(P<0.05$, Table 1). There was an effect of percent zebra mussel coverage on amphipod consumption for bullhead and crayfish $(P<0.05$, Table 1); both species consumed more amphipods in bare treatments than in either the 50 or 95% zebra mussel coverage treatment (Tukey's, P<0.05). There was no effect of percent zebra mussel coverage on amphipod consumption for sculpins or logperch.

Habitat choice

All four predators were observed more frequently on complex substrates (gravel and zebra mussels) than on sand (Fig. [2](#page-4-0)). For each species, the number of times a predator was observed in each of the three habitat types was nonrandom $(G=438.2)$ for bullhead, $G=207.8$ for sculpin, $G=296.8$ for logperch, $G=255.4$ for crayfish, $d \ge 16$, P<0.001 for all species). All predators except sculpin were observed more frequently on zebra mussels than on gravel or sand. Sculpin were observed less often on zebra mussels as the day progressed (Fig. [2\)](#page-4-0). Both

sculpin and crayfish were frequently observed burrowed among zebra mussels. We did not observe any of the predators actively feeding on zebra mussels during the experiments.

Discussion

The effects of habitat structure on predator-prey interactions has been very well studied in a wide range of ecological systems (e.g. Huffaker [1958;](#page-6-0) Persson and Eklöv [1995;](#page-6-0) Dudgeon [1996](#page-5-0); Lombardo [1997](#page-6-0)). In general, increased habitat complexity has been shown to influence the ability of predators to forage successfully (Crowder and Cooper [1982;](#page-5-0) Gilinsky [1984;](#page-5-0) Gotceitas and Colgan [1989](#page-6-0); Deihl [1992\)](#page-5-0). Zebra mussels increase habitat complexity, and, in an elegantly designed experiment, Botts et al. [\(1996](#page-5-0)) convincingly demonstrated that the positive effects of zebra mussels on benthic macroinvertebrate communities are mediated through increased habitat complexity as well as food web enrichment.

As zebra mussels colonize soft sediments, they have the potential to affect the foraging success of a range of species. In particular, juveniles of some species (e.g., yellow perch, walleye, sturgeon) utilize soft sediment zoobenthos for food during critical growth periods (Ritchie and Colby [1988](#page-6-0); Scott and Crossman [1973](#page-6-0); Beamish et al. [1998\)](#page-5-0). Negative effects on already threatened or endangered species such as the lake sturgeon

Fig. 2 Habitat use of four benthic species, a sculpins, b logperch, c bullhead, and d crayfish, presented as the percentage of time spent in zebra mussels, gravel, and bare sand during three time periods: dawn, midday, and dusk. Stacked bars represent the average time spent in each habitat for each time period. $n=12$, 11, 8, and 11, for sculpins, logperch, bullheads, and crayfish, respectively

may have important implications for conservation and reintroduction programs. Zebra mussels may impede the ability of these species to obtain burrowing prey. Whether this potential negative effect of zebra mussels on fish foraging is offset by zebra-mussel facilitation of the macrozoobenthos would depend on the prey specificity of the fish species in question.

We have demonstrated that zebra mussels impede the foraging of bullheads, sculpins, and logperch on burrowing prey (Chironomus). Because crayfish burrow under the mussel colonies, they consume Chironomus regardless of zebra mussel coverage. Crawling prey (isopods) were not readily consumed by logperch, regardless of treatment, but all other foragers had reduced foraging success on isopods in the presence of zebra mussels. Amphipods (swimming prey) were consumed by all predators, but there was no effect of zebra mussels on the number consumed by sculpins or log perch. There was a marked reduction in the number of amphipods consumed by bullheads and crayfish in the presence of zebra mussels. It is possible that because crayfish and bullheads are less visual than the other predators, the amphipods stood a better chance of evading capture in the complex environment provided by the zebra mussels.

Other investigators have used a range of techniques to assess the impacts of zebra mussels on fish foraging and habitat selection. Mayer et al. ([2001\)](#page-6-0) used videotaped trials and counted yellow perch foraging attacks and captures of amphipods in aquariums with and without zebra mussels, at three different light levels. In addition to videotaping trials of yellow perch foraging on amphipods with zebra mussels and bare sand in the same tanks, Cobb and Watzin [\(2002](#page-5-0)) ran foraging trials with and without zebra mussels followed by stomach analysis to quantify the number of amphipods and chironomids consumed on sandy and hard substrates. The video techniques used by Mayer et al. ([2001\)](#page-6-0) and Cobb and Watzin ([2002\)](#page-5-0) are advantageous because failed foraging attempts can be distinguished from successful captures. González and Downing's ([1999\)](#page-6-0) experiments are most directly comparable to our study in that they ran timed foraging trials with yellow perch or bluegill in aquariums with and without zebra mussels on rocky substrates. By adding a known number of amphipods to each aquarium and recovering the remaining prey at the end of the foraging trial, González and Downing ([1999\)](#page-6-0) had a measure of foraging success that was similar to ours.

Despite the diversity of techniques employed in previous studies, a strong pattern is emerging. With one exception (yellow perch; González and Downing [1999\)](#page-6-0) zebra mussels reduced the foraging success of fish on macroinvertebrates on hard substrates. In addition, Cobb and Watzin [\(2002](#page-5-0)) found that yellow perch were proportionately more successful when foraging on sand versus foraging on zebra mussel colonies. Yellow perch also made more foraging attempts in zebra mussel colonies and fewer attempts in sand than would have been expected based on the area of each substrate type available.

González and Downing ([1999\)](#page-6-0) found no effect of zebra mussels on yellow perch foraging success and suggested that this was because the perch were more adept at capturing macroinvertebrates and thus less affected by zebra mussels. Another possibility is that the larger perch used by González and Downing ([1999\)](#page-6-0) had lower affinity for macroinvertebrates. Cobb and Watzin (2002) found in one of their trials that larger perch consumed fewer amphipods, and indeed ontogenetic changes in diet have previously been documented in yellow perch such that their diets shift from zooplankton, to macroinvertebrates, to piscovory as they grow larger (e.g., Wu and Culver [1992](#page-6-0)). González and Downing [\(1999](#page-6-0)) used perch with an average total body length of 130 mm, at the upper extreme of the range of the fish used by Cobb and Watzin (2002; range 59–131 mm; hard substrate, full coverage trial). Similarly, Mayer et al. ([2001\)](#page-6-0) used small yellow perch (range 65–126 mm) and found that the presence of zebra mussels reduced foraging success.

Our results compliment the previously existing studies by examining a range of foraging styles and prey uses of the habitat. The absence of an effect of zebra mussels on sculpins and log perch foraging on amphipods suggests that these predators may benefit from the increases in nonburrowing amphipod abundance reported following zebra mussel colonization. In addition, because zebra mussels have different effects on predation depending on the prey species considered, predators may switch prey species in response to zebra mussel colonization. Thus, even though the number of a given prey species consumed may change, there may not be a net total effect of zebra mussels on total biomass consumed by a given predator.

Predator utilization of zebra mussel beds as habitat may result in significant local redistribution of fish, potentially altering access to food and vulnerability to predation. Predators typically associated with rocky substrates may utilize zebra mussels on soft sediments. With the exception of the sculpins in two out of three time periods, the predators in our experiments spent the majority of their time on zebra mussels. Cobb and Watzin (2002) found that yellow perch made a disproportionate number of foraging attempts on zebra mussel colonies relative to surrounding bare substrates, but this response was in part because prey density in the zebra mussel colonies was 300 times greater than in the surrounding substrate. Our results are free from the effects of prey density and demonstrate that there is a strong tendency for benthic predator species to spend more time on zebra mussels than on sand or gravel.

Zebra mussels increase surface area, and add threedimensional structure to the surfaces to which they attach. The zebra mussel druses (isolated clumps) and layers that form on sandy and silty sediments produce a nook-andcranny effect unlike anything previously existing on soft or bedrock substrates in newly invaded lakes. Our results convincingly demonstrate that zebra mussels reduce the foraging success of a range of benthic predators, including the first such demonstration for an invertebrate. By testing a range of prey types, we have demonstrated that prey position in the habitat (burrowing, crawling, or swimming)

interacts with the foraging modes of the predators (tactile, visual, lateral line, strictly benthic, burrowing) to produce a range of potential interactions among zebra mussels, macroinvertebrates, and predators. Zebra mussels clearly inhibit predation on burrowing and crawling prey by benthic foraging fish. In contrast, swimming/crawling prey such as amphipods are susceptible to predation even in the presence of zebra mussels, except by bullheads. All predators used in these experiments spent more time among zebra mussels as compared to bare sand or gravel. Clearly, increased habitat complexity created by zebra mussel colonization of substrates has an effect on habitat use. Whether this usage shift is influenced by reduced predation from larger predators remains to be determined.

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