INFLUENCE OF FEEDING HABITS ON ORGANOCHLORINE CONTAMINANT ACCUMULATION IN WATERFOWL ON THE GREAT LAKES

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Abstract. Zebra mussels (Dreissena polymorpha) are an important component of benthic communities in the Great Lakes and are exploited by a host of predators, including waterfowl. In this study, we analyze diet content and stable isotope and organochlorine contaminant patterns in Lesser Scaup (Aythya affinis), Greater Scaup (Aythya marila), Bufflehead (Bucephala albeola), Redhead (Aythya americana), Canvasback (Aythya valisineria), and Mallard (Anas platyrhynchos) collected from three sites (Fighting Island, western Lake Erie, Big Creek) in the lower Great Lakes. Lesser and Greater Scaup from Fighting Island were classified as either zebra mussel (≥67% of diet) or macrophyte (≥85% of diet) consumers. Bufflehead, Canvasback, Mallard, and Redhead consumed mainly (≥89%) macrophyte at Fighting Island. Zebra mussel was the principal food of Lesser Scaup (>99%), Greater Scaup (97%), and Bufflehead (72%) in western Lake Erie. Stable isotope analysis revealed enrichment of δ¹⁵N in Lesser Scaup (≥2.24%), Greater Scaup (≥1.28%), and Bufflehead (≥0.63%) that exploited mussels relative to conspecifics with macrophyte diets and relative to mussel prey.

Representative contaminants of low (hexachlorobenzene [HCB]), moderate (PCB [polychlorinated biphenyl] 153), and high (PCB 180) hydrophobicity were examined in waterfowl. Lipid-normalized concentrations of PCBs 153 and 180 were significantly higher in scaup and Bufflehead that consumed Dreissena than in individuals that ate mainly macrophytes. Among taxa that consumed primarily Dreissena, concentrations of PCBs 153 and 180 were significantly higher in individuals from Lake Erie than in those Fighting Island. Principal components analysis revealed broad differences in contaminant patterns of waterfowl based principally on diet.

Results from this study illustrate that Dreissena has become a primary food source of some waterfowl in the lower Great Lakes and serves as an effective conduit for transfer of persistent organic contaminants to higher trophic levels.

Key words: bioaccumulation; Dreissena; ducks; ecotoxicology; Great Lakes; Lake Erie; organochlorine contaminants; pesticides; polychlorinated biphenyls; principal components analysis; stable isotopes; waterfowl; zebra mussel.

INTRODUCTION

The Laurentian Great Lakes have been subject to species introductions since the onset of European settlement. More than 135 nonindigenous species have become established in the basin (see Mills et al. 1993). The zebra mussel, Dreissena polymorpha Pallas, established in the Great Lakes in 1985, and has since expanded its distribution to most major rivers and some inland lakes in temperate eastern North America. Zebra mussels are dominant contributors to benthic invertebrate abundance and biomass in many regions of the Great Lakes, including Saginaw Bay, Lake St. Clair, western Lake Erie, and the Detroit River (see Nalepa and Schloesser 1993). Well-established effects of Dreissena include enhanced water transparency, reduced phytoplankton concentration, and increases in densities of many benthic invertebrates (reviewed in MacIsaac 1996). Much less is known regarding the mussel’s predator-prey relationships, and its effects on contaminant dynamics in invaded systems.

In the Great Lakes, Dreissena is consumed by a host of predators including crayfish and fishes (reviewed in MacIsaac 1996). Waterfowl are the predators most likely to have significant impacts on Dreissena populations owing to rapid numerical responses and strong feeding preferences of some species for bivalve prey. The number and duration of staging Greater Scaup (Aythya marila), Lesser Scaup (A. affinis), and Common Goldeneye (Bucephala clangula) increased dramatically subsequent to Dreissena establishment in the western basin of Lake Erie (Wormington and Leach 1992). Waterfowl
predators quickly exploit recently established *Dreissena* populations. For example, Lesser Scaup preyed on *Dreissena* during the first winter following mussel establishment at a power plant on Lake Michigan (Mitchell and Carlson 1993). Predation by Greater Scaup, Lesser Scaup, Common Goldeneye, and Bufflehead (*B. albeola*) may strongly affect zebra mussel population biomass (Wormington and Leach 1992, Custer and Custer 1996). For example, ducks reduced *Dreissena* biomass by 57% at a nearshore habitat on western Lake Erie (Hamilton et al. 1994).

*Dreissena* has been used extensively as a sensitive biomonitor of organochlorine contaminants (e.g., Duursma et al. 1984, Bruner et al. 1994, Morrison et al. 1995, Roe and MacIsaac 1997). Briefer and Hunter (1993) reported significantly higher accumulations of PCB congeners in *Dreissena* than in native *Lampsilis* clams in the Great Lakes. Despite the mussel's significant bioaccumulation ability, information regarding biomagnification in *Dreissena* predators is lacking.

Like *Dreissena*, uptake of PCBs occurs very rapidly in waterfowl, and it may have important ecotoxicological implications. Gebauer and Weseloh (1993) reported that mallard (*Anas platyrhynchos*) accumulated PCBs to 5300 times initial values within 10 d of natural exposure. In a European study, caged Tufted Ducks (*Aythya fuligula*) fed *Dreissena* contaminated with an array of organic contaminants laid fewer eggs, abandoned nests more often, and had higher embryo and chick mortality rates than ducks fed less polluted mussels (de Kock and Bowmer 1993). Thus, consumption of contaminated *Dreissena* has the potential to adversely affect reproductive success of waterfowl in North America.

Seasonal and spatial variation in feeding habits of waterfowl may encumber assessments of dietary sources of contaminants because traditional diet analysis (i.e., gut content) reflects only the most recent feeding history. This problem can be resolved by using stable isotopes of N and C to characterize trophic relationships of complex food webs (Hobson 1990, Hamilton et al. 1992, Cabana and Rasmussen 1994, Kidd et al. 1995). The nitrogen isotopic ratio ($\delta^{15}N$) is strongly indicative of the trophic position of an organism, whereas the carbon isotopic ratio ($\delta^{13}C$) can be useful in defining food sources (e.g., Kirilik et al. 1995, Riera and Richard 1996). The nitrogen and carbon isotopic ratios increase by $\approx 3.4$ and $1\%$, respectively, per trophic level, owing to a combination of biological, chemical, and physical processes that cause differential assimilation of isotopes (Fry and Quinones 1994, Keough et al. 1996). Recent studies have also illustrated that $\delta^{15}N$ values are correlated with contaminant burden in a variety of freshwater taxa (Kirilik et al. 1995, Kidd et al. 1995).

The purpose of this study was to assess variation in contaminant burden in waterfowl in relation to diet, particularly with respect to individuals that exploit zebra mussels. We complement conventional diet analysis with that of stable isotopes to determine the importance of zebra mussels as food and contaminant sources. We also assess relative contaminant exposure of waterfowl at three different locations in the lower Detroit River–western Lake Erie ecosystem.

**METHODS**

**Study sites and waterfowl collection**

Waterfowl were collected from three sites: Fighting Island in the lower Detroit River, Big Creek Marsh, and in western Lake Erie between Middle Sister and Hen Islands and the Canadian mainland (Fig. 1). Wetlands in the lower Detroit River are frequented by migrating and overwintering waterfowl (Prince et al. 1992). Shallow areas downstream of Fighting Island support extensive beds of macrophytes, i.e., *Vallisneria americana*, *Elodea canadensis*, and *Potamogeton* spp. (Davis and Erwin 1982). Gastropods and *Dreissena* occur on macrophytes and on mud flats in the Fighting Island area (Davis and Erwin 1982; E. Mazak, personal observation). Hen Island is located 8 km west of Pelee Island in western Lake Erie, while Middle Sister Island is located 15.7 km from the Canadian mainland in the central region of the basin. Mean depth in the basin is 7.3 m, though deeper passages exist between islands and the mainland. Commercial fishermen deploy gill...
nets between the islands and the mainland throughout the ice-free season.

Seventy-one waterfowl were collected and analyzed for diet contents and organochlorine contaminants. Fighting Island and Big Creek waterfowl were donated by hunters. Lake ducks that had drowned in gill nets were furnished by commercial fishermen. Fighting Island taxa collected included Mallard, Canvasback (Aythya valisineria), Redhead (Aythya americana), Buffalohead, and Greater and Lesser Scaup. All ducks were collected between autumn 1993 and winter 1994. Hunteed ducks were at identified feeding sites when killed, while drowned individuals were found at considerable depth (up to 7 m) in areas supporting mussels and often had Dreissena in their mouth and esophagus.

Duck mass was determined to the nearest 25 g using a spring balance (Ohaus/Canadawide Scientific, Ottawa, Ontario, Canada) length was measured (±0.25 cm), the digestive tract was removed, and liver and/or wing tissues were excised and wrapped in hexane-rinsed aluminum foil and frozen pending organochlorine contaminant and stable isotope analyses. Species, age, and sex of waterfowl were determined according to Carney (1992).

The esophagus, proventriculus, and gizzard of each duck were analyzed for diet contents. Decomposition of food items, particularly in the gizzard, limited taxonomic resolution of diet contents. Diet items were categorized into four main fractions: Dreissena, macrophyte (mainly Elodea, Vallisneria, and Potamogeton spp.), snail, or amphipod (Gammarus fasciatus). Right valve and septa lengths of whole mussels obtained from duck digestive tracts were measured to the nearest 0.01 mm using an ocular micrometer and dissecting microscope (Hamilton 1992). Right valve length (VL) was estimated from right septa length (SL) for mussel shell fragments recovered from waterfowl gizzards as \( VL = 0.29 + 8.32SL \) \( r^2 = 0.92 \). Biomass of diet contents was determined to the nearest 0.5 mg with an electronic balance (A&D, Model FX-200, Milpitas, California, USA) after samples were air dried at 20°C for at least 72 h.

Samples of waterfowl diet items were collected from field sites for contaminant and stable isotope analysis. Macrophytes from Fighting Island were collected in 1.5 m of water during August 1994, while snails and Dreissena were obtained by benthic dredge (mesh size 2 mm) from 1.5 to 2.5 m depth during December 1994. Dreissena, macrophytes, and gastropod samples from Middle Sister Island were collected during September 1994 in 2–3 m of water.

Contaminant and isotope analysis

Contaminant concentrations were measured directly in livers of 62 ducks, of which 11 individuals were also assayed for concentrations in wing tissues. Lipid-normalized liver concentrations were regressed on lipid-normalized wing concentrations for these 11 individuals. Lipid-adjusted liver concentrations were calculated for nine additional ducks for which only lipid-adjusted wing values were available. Lipid-adjusted wing and liver values were highly correlated, with the coefficient of determination \( r^2 \) ranging between 0.72 for pentachlorobenzene (QCB) and PCB 28 and 0.99 for PCBs 153 and 194. The number of individuals analyzed for contaminants ranged between 3 and 10 for each group of species classified by site and diet (see Table 1).

Sample preparation and gas chromatography procedures for organochlorine contaminant analyses of waterfowl wing and/or liver tissue, Dreissena soft tissues, and macrophyte aboveground tissues follow Lazar et al. (1992). A small portion of each animal sample was used for gravimetric determination of lipid content (see Lazar et al. 1992). Contaminant analyses were conducted on a Hewlett Packard-5890 Series II Gas Chromatography/N-Electron Capture Detector instrument (Hewlett-Packard, Mississauga, Ontario, Canada). Canadian Wildlife Service Standards (CWS–RSM 8229–Herring Gull egg pool) were run as references every 6–8 samples. Sixty-five contaminants encompassing a broad array of pesticides, chlorinated hydrocarbons, and PCBs were assayed in waterfowl and prey items. Twenty-six chemicals were deleted from analysis because of coelution problems or because the majority of samples contained nondetectable (<0.05 μg/kg wet mass) levels of contaminant. If more than half of the samples had detectable levels of a particular contaminant, nondetectable values were replaced by random numbers between 0.00 and 0.05 μg/kg wet mass generated using Quattro Pro. All animal wet mass chemical concentrations were lipid normalized; mean lipid content in liver and wing tissue varied between 3.3 and 6.5%, and was typically slightly higher in ducks from Lake Erie than from either of the other sites (see Table 1). Wet mass contaminant levels in macrophytes were normalized based on organic carbon content. PCB identification numbers and associated octanol–water partition coefficients (\( K_{ow} \); a measure of hydrophobicity) are based on the IUPAC classification system (Shiu and Mackay 1986; Table 1).

Stable carbon and nitrogen isotope analyses were conducted on wing tissue from 33 waterfowl and 9 macrophyte, snail, and Dreissena diet samples collected from the field. Waterfowl tissues were trimmed to remove visible lipid, and cut into 5-mm square cubes. Dreissena and snails collected from Lake Erie were shocked, and the soft tissues diced. Samples (1 g) were placed in a glass-stoppered flask and swirled in 100 mL of acetone suspension for 15 min. Samples were rinsed with 50 mL dichloromethane for 1 min, dried for 48-72 h at 60°C, and powdered with mortar and pestle. Plant material from Fighting Island was dried at 60°C, then powdered in a Wiley mill. Samples were analyzed in an automated elemental analyzer (Carlo Erba NA 1500) attached to a VG Optima au-
TABLE 1. Concentration (mean ± 1 SE) of representative low-$K_{ow}$ (hexachlorophenol [HCB], PCB 28), mid-$K_{ow}$ (PCBs 149, 153), and high-$K_{ow}$ (PCBs 180, 194) organochlorine contaminants, and total PCB concentration, in waterfowl and primary prey. Mass and length are mean values.

<table>
<thead>
<tr>
<th>Species</th>
<th>Site</th>
<th>Sample size</th>
<th>Male (%)</th>
<th>Adult (%)</th>
<th>Mass (g)</th>
<th>Length (cm)</th>
<th>Diet</th>
<th>Concentration (µg/kg lipid)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mallard</td>
<td>BC</td>
<td>4</td>
<td>50</td>
<td>75</td>
<td>1238</td>
<td>59</td>
<td>5</td>
<td>HCB 4 ± 1, PCB 28 4 ± 1, PCB 149 14 ± 9</td>
</tr>
<tr>
<td>Mallard</td>
<td>FL</td>
<td>3</td>
<td>67</td>
<td>100</td>
<td>1067</td>
<td>53</td>
<td>m</td>
<td>4 ± 1, 2 ± 0, 13 ± 5</td>
</tr>
<tr>
<td>Canvasback</td>
<td>FL</td>
<td>6</td>
<td>50</td>
<td>67</td>
<td>1193</td>
<td>54</td>
<td>m</td>
<td>23 ± 15, 3 ± 0, 17 ± 10</td>
</tr>
<tr>
<td>Redhead</td>
<td>FL</td>
<td>3</td>
<td>33</td>
<td>67</td>
<td>1085</td>
<td>50 mm</td>
<td>3</td>
<td>5 ± 2, 3 ± 1, 9 ± 4</td>
</tr>
<tr>
<td>Bufflehead</td>
<td>LE</td>
<td>5</td>
<td>75</td>
<td>0</td>
<td>498</td>
<td>35</td>
<td>zm</td>
<td>37 ± 5, 9 ± 3, 32 ± 6</td>
</tr>
<tr>
<td>Lesser Scaup</td>
<td>FI</td>
<td>4</td>
<td>50</td>
<td>0</td>
<td>479</td>
<td>37 zm</td>
<td>zm</td>
<td>30 ± 5, 8 ± 2, 154 ± 10</td>
</tr>
<tr>
<td>Lesser Scaup</td>
<td>LE</td>
<td>3</td>
<td>100</td>
<td>33</td>
<td>883</td>
<td>42 zm</td>
<td>zm</td>
<td>30 ± 3, 2 ± 1, 9 ± 4</td>
</tr>
<tr>
<td>Greater Scaup</td>
<td>LE</td>
<td>8</td>
<td>63</td>
<td>75</td>
<td>985</td>
<td>44 zm</td>
<td>zm</td>
<td>23 ± 2, 4 ± 1, 114 ± 20</td>
</tr>
<tr>
<td>Greater Scaup</td>
<td>FL</td>
<td>4</td>
<td>50</td>
<td>50</td>
<td>1028</td>
<td>43 m</td>
<td>m</td>
<td>63 ± 46, 2 ± 0, 14 ± 2</td>
</tr>
<tr>
<td>Greater Scaup</td>
<td>FL</td>
<td>3</td>
<td>33</td>
<td>33</td>
<td>1000</td>
<td>40 zm</td>
<td>zm</td>
<td>36 ± 23, 1 ± 0, 7 ± 5</td>
</tr>
<tr>
<td>Greater Scaup</td>
<td>LE</td>
<td>5</td>
<td>40</td>
<td>40</td>
<td>1270</td>
<td>48 zm</td>
<td>zm</td>
<td>42 ± 10, 6 ± 1, 147 ± 13</td>
</tr>
<tr>
<td>Macrophyte³</td>
<td>FI</td>
<td>6</td>
<td>&lt;1</td>
<td>0</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>&lt;1 ± 0</td>
</tr>
<tr>
<td>Zebra mussel</td>
<td>FL</td>
<td>2</td>
<td>78</td>
<td>16</td>
<td>4 ± 0</td>
<td>190 ± 37</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Zebra mussel</td>
<td>LE</td>
<td>5</td>
<td>4 ± 1</td>
<td>4 ± 1</td>
<td>193 ± 7</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note: Site key: BC = Big Creek, FI = Fighting Island, LE = Lake Erie. Diet refers to major prey type found in the digestive tract. Diet key: m = macrophyte, zm = Dreissena. ND = not detectable.

† Values in parentheses are log $K_{ow}$.
‡ Macrophyte (Elodea canadensis, Myriophyllum spicatum) data from R. Russell (unpublished data) and are adjusted for organic carbon content. All other contaminant data are lipid adjusted.

Statistical analysis

Examination of Fighting Island Greater and Lesser Scaup revealed that individuals consumed either mussels or macrophytes as the primary diet component. These individuals were thus classified as mussel consumers or macrophyte consumers. In order to justify this classification scheme, a two-way ANOVA was conducted on the proportion of Dreissena in diets of all Lesser and Greater Scaup from Fighting Island, with duck species and diet as main effects. Similarly, a two-way ANOVA with species and collection site as main effects was conducted on the proportion of Dreissena in diet of Greater and Lesser Scaup mussel consumers from Fighting Island and Lake Erie. Differences in the proportion of Dreissena in diets of Lake Erie and Fighting Island bufflehead were explored using Student’s $t$ test.

Differences in utilization of macrophytes by Fighting Island waterfowl (Canvasback, Redhead, Mallard, and Greater and Lesser scap) that consumed mainly macrophytes were explored using one-way ANOVA. All proportion diet data were arcsine square-root transformed prior to analyses. All transformed data were homoscedastic (Cochran’s test, $\alpha = 0.01$).

Size distributions of mussels consumed by Buflehead, Lesser Scaup, and Greater Scaup from Lake Erie were compared using Kolmogorov-Smirnov tests at an adjusted significance level ($\alpha$) of 0.02.

All statistical analyses of variation in contaminant concentrations utilized log($x + 1$)-transformed, lipid-normalized data, and were limited to three compounds (HCB, PCB 153, and PCB 180). These chemicals were selected as representatives of low-, mid- and high-$K_{ow}$ compounds and because none of the compounds are metabolized. Variation in contaminant concentrations of these compounds in Lesser and Greater Scaup from Fighting Island that consumed plant or mussel diets was assessed using two-way multivariate analysis of variance (MANOVA) with species and diet main effects. Species and site effects were analyzed for these compounds using two-way MANOVA on Greater and Lesser Scaup from Fighting Island and Lake Erie that consumed mussel diets. Species differences in concen-
Table 1. Continued.

<table>
<thead>
<tr>
<th>Concentration (µg/kg lipid)</th>
<th>PCB 153 (6.92)</th>
<th>PCB 180 (7.36)</th>
<th>PCB 194 (7.80)</th>
<th>Total PCB</th>
</tr>
</thead>
<tbody>
<tr>
<td>30 ± 6</td>
<td>14 ± 5</td>
<td>3 ± 0</td>
<td>347 ± 149</td>
<td></td>
</tr>
<tr>
<td>29 ± 14</td>
<td>19 ± 8</td>
<td>3 ± 2</td>
<td>330 ± 172</td>
<td></td>
</tr>
<tr>
<td>67 ± 42</td>
<td>43 ± 8</td>
<td>10 ± 7</td>
<td>433 ± 238</td>
<td></td>
</tr>
<tr>
<td>16 ± 5</td>
<td>7 ± 2</td>
<td>1 ± 0</td>
<td>179 ± 75</td>
<td></td>
</tr>
<tr>
<td>73 ± 19</td>
<td>55 ± 12</td>
<td>9 ± 2</td>
<td>693 ± 154</td>
<td></td>
</tr>
<tr>
<td>462 ± 93</td>
<td>446 ± 93</td>
<td>75 ± 15</td>
<td>3722 ± 496</td>
<td></td>
</tr>
<tr>
<td>29 ± 20</td>
<td>17 ± 15</td>
<td>4 ± 4</td>
<td>300 ± 190</td>
<td></td>
</tr>
<tr>
<td>131 ± 55</td>
<td>113 ± 35</td>
<td>16 ± 7</td>
<td>914 ± 317</td>
<td></td>
</tr>
<tr>
<td>298 ± 77</td>
<td>340 ± 64</td>
<td>63 ± 11</td>
<td>2734 ± 483</td>
<td></td>
</tr>
<tr>
<td>16 ± 3</td>
<td>18 ± 5</td>
<td>2 ± 0</td>
<td>185 ± 21</td>
<td></td>
</tr>
<tr>
<td>200 ± 182</td>
<td>187 ± 173</td>
<td>33 ± 29</td>
<td>1462 ± 1191</td>
<td></td>
</tr>
<tr>
<td>457 ± 165</td>
<td>477 ± 135</td>
<td>79 ± 25</td>
<td>4464 ± 1363</td>
<td></td>
</tr>
<tr>
<td>&lt;1 ± 0</td>
<td>&lt;1 ± 0</td>
<td>&lt;1 ± 0</td>
<td>25 ± 2</td>
<td></td>
</tr>
<tr>
<td>203 ± 35</td>
<td>99 ± 3</td>
<td>20 ± 3</td>
<td>3375 ± 580</td>
<td></td>
</tr>
<tr>
<td>224 ± 13</td>
<td>169 ± 10</td>
<td>29 ± 2</td>
<td>2359 ± 97</td>
<td></td>
</tr>
</tbody>
</table>

Site-based variation in contaminant concentrations was also analyzed for Buf®head from Lake Erie and Fighting Island, and, separately, for Mallard from Big Creek and Fighting Island, using Bonferroni-adjusted (α = 0.015) Student’s t tests. Principal components analysis (PCA) was performed on log(x + 1)-transformed, lipid-adjusted chemical concentrations using nine compounds. In addition to the chemicals listed above, two other representative low- (QCB, PCB 28), mid- (PCBs 149, 118), and high- (PCBs 174, 194) K<sub>ow</sub> compounds were utilized in the model. The model employed a correlation matrix and varimax rotation (SYSTAT 1992). Two-way MANOVA was conducted on the first three principal component factor scores for scaup from Fighting Island to assess species and diet main effects. Similarly, factor scores for the first three principal components for scaup from Fighting Island and Lake Erie were analyzed using two-way MANOVA to assess species and location main effects. Variation in species factor scores for all waterfowl from Fighting Island that consumed mainly macrophytes was examined using one-way MANOVA. Variation in PCA factor scores of Buf®head and Mallard from different sites was analyzed separately using t tests with Bonferroni adjustments (α = 0.015).

Carbon and nitrogen isotope patterns in Greater and Lesser Scaup from Fighting Island were subject to a two-way MANOVA to assess effects of species and diet. MANOVA was also used to test for species and site differences in N and C isotope ratios for Fighting Island and Lake Erie scaup that consumed mainly Dreissena. One-way ANOVAs followed by Dunnett’s one-sided means tests were run separately for 13C and 15N stable isotopes to determine whether isotope en-

RESULTS

Physical examination revealed no gross external or internal abnormalities in any waterfowl in the study. Mallard from Big Creek and Greater Scaup from Lake Erie were the largest and heaviest waterfowl, respectively, while Buf®head were the smallest and lightest ducks (Table 1). Lesser and Greater Scaup from Lake Erie were slightly larger and heavier than conspecifies from Fighting Island (Table 1).

Lesser and Greater Scaup from Lake Erie consumed Dreissena almost exclusively (>97%; Fig. 2). By contrast, individuals from Fighting Island could be classified into those that ate either mainly Dreissena or mainly macrophytes. Scaup classified as mussel consumers ate significantly more Dreissena than those classified as macrophyte consumers (F = 225, df = 1, 17, P < 0.001); species (F = 3.8, df = 1, 17, P = 0.069) and species × diet (F = 1.4, df = 1, 17, P = 0.241) differences were not evident.

Diet of scaup from Lake Erie also contained more Dreissena than those classified as mussel consumers.
Concentrations of most mid- and high-
low compounds varied by diet
(F = 3.9, df = 3, 15, P = 0.030, MANOVA), though
differences were limited to mid- and high-
K\text{ow} compounds (PCB 153: F = 8.9, df = 1, 17, P = 0.008) and high-
K\text{ow} compounds (PCB 180: F = 12.8, df = 1, 17, P = 0.002) congeners. No species or species \times diet interactions were detected among
Fighting Island scaup (P = 0.606 and 0.524, respectively; MANOVA).

Average total PCB concentration in Lake Erie
lesser scaup and greater scaup were 3.5 and 2.7 times higher, respectively, than in conspecifics from Fighting Island that consumed mainly
Dreissena (Table 1). While contaminant concentrations were significantly higher in waterfowl from Lake Erie (F = 6.9; df = 3, 17, P = 0.003, MANOVA), these differences were also limited to mid- and high-
K\text{ow} compounds (P \leq 0.004, ANO-
VAs). No species or species \times site interaction differences in concentrations of representative chemicals were detected among scaup from Lake Erie and Fighting Island (MANOVA, P > 0.500).

Site differences in chemical concentrations were also apparent in Bubblehead. Concentrations of most mid- and high-
K\text{ow} contaminants were between 2 and 10 times higher in individuals from Lake Erie relative to those from Fighting Island. For example, bubblehead from Lake Erie had significantly higher concentrations of PCB 153 (t = 5.8, df = 10, P = 0.002) and PCB 180 (t = 5.0, df = 10, P = 0.0005), though these site differences were confounded by diet (Fig. 2; Table 1). Variation in low-
K\text{ow} congeners was similar among sites (e.g., HCB: t = 1.2, df = 10, P = 0.28).

Mallard from Fighting Island and Big Creek did not vary with respect to concentrations of any of the compounds tested (t tests, P > 0.50).

Principal components (PC) analysis was conducted on contaminant data in waterfowl using nine compounds of varying
K\text{ow}. PC1 accounted for 41\% of original data variability, and was determined primarily by mid- and high-
K\text{ow} congeners (PCBs 180, 153, 194, and 118). PC2 and PC3 accounted for an additional 13\% and 12\% of data variability and were, respectively, determined by the low-
K\text{ow} compounds QCB and PCB 28.

Waterfowl separated clearly on PC1 based on diet (Fig. 4). In general, waterfowl that consumed mussels had much higher PC1 scores than those that ate macrophyte diets (Fig. 4). For example, scores of Lesser and Greater Scaup from Fighting Island varied by diet (F = 4.9, df = 3, 15, P = 0.014, MANOVA), though differences were limited to PC axis 1 only (F = 15, df = 1, 17, P = 0.0012, ANOVA). Waterfowl with mussel
scores were insigniﬁcant (t < 0.06), though the latter result likely reﬂects small sample sizes (Fig. 5; Table 1).

Stable isotope ratios of N and C in Island scaup did not vary by species (F = 0.4, df = 2, 7, P > 0.10, MANOVA) or by diet (F = 4.2, df = 2, 7, P = 0.06), though the latter result likely reﬂects small sample sizes (Fig. 5; Tabachnick and Fidell 1996). As evidence, univariate analysis of the latter result was signiﬁcant for nitrogen (F = 8.9, df = 1, 8, P < 0.02), indicating trophic enrichment in mussel predators. Moreover, ratios of δ15N and δ13C isotopes in scaup that consumed mussels did not vary by species (F = 0.52, df = 2, 7, P > 0.10) or by location (F = 0.36, df = 2, 7, P > 0.10, MANOVA).

All Lake Erie waterfowl that consumed mussels had isotope ratios that were very similar with respect to both δ15N and δ13C (Fig. 5). δ15N ratios were enriched by ≥2.3‰ in mussel predators relative to Dreissena, though differences among groups were not signiﬁcant (F = 1.17, df = 3, 8, P = 0.37, ANOVA).

All Fighting Island ducks that consumed macrophytes were signiﬁcantly depleted with respect to δ15N relative to macrophytes (Dunnett’s test, P < 0.01; Fig. 5). In addition, Redhead, Canvasback, and Bufflehead were enriched in δ15N by 2–4‰ relative to macrophytes (Dunnett’s test, P < 0.01), while values for Mallard and Lesser and Greater Scaup were similar to those of macrophytes (Fig. 5).

**Discussion**

Detroit River sediments are highly polluted with organochlorine compounds. River inputs serve as the primary source (73%) of organic contaminants to western Lake Erie (Carter and Hites 1992). Koslowski et al. (1994) documented signiﬁcant accumulations of a wide array of organochlorine compounds in the western Lake Erie food web, and argued that trophic interactions play an important role in contaminant exposure. Results from this study establish that recent diet is a good predictor of contaminant exposure in lower Great Lakes waterfowl, and that these patterns are generally corroborated by stable isotope analyses. Ducks that consumed Dreissena typically displayed elevated contaminant concentrations and were enriched with respect to δ15N relative to co-occurring individuals that ate little or no Dreissena (Fig. 5; Table 1).

Uptake of PCBs occurs very rapidly in waterfowl (Gebauer and Weseleh 1993). Mallard exposed to dieldrin achieve steady-state concentrations in muscle, liver, and fat within 8 d (Nebeker et al. 1994). The rapidity of uptake indicates that migrating waterfowl are susceptible to short-term contaminant exposure, and body burdens reﬂect recent exposure to contami-
nants. High contaminant levels in ducks with mussel diet almost certainly reflect local exposure. Every duck examined for organic contaminants that had a diet of zebra mussels was collected between late November and mid-December or during mid-April, at least 3 wk after the ducks had arrived on western Lake Erie in mass migrations (G. Ives, personal communication). This time differential should have been sufficient to permit significant accumulation of organochlorine contaminants. Conversely, it is unlikely that the ducks became contaminated via exploitation of another food source prior to migrating to Lake Erie. The ducks arrive on Lake Erie after migrating south from Alaskan breeding grounds through Saskatchewan, Manitoba, and the upper Great Lakes (Barclay and Zingo 1993). Hatching-year scaup from Saskatchewan contain extremely low levels of organochlorine compounds, indicating that they become contaminated after migrating through the prairies (E. Mazak, unpublished data). The lower Detroit River–western Lake Erie corridor is the most contaminated region visited by Greater Scaup on the flyway that takes them from Alaska through the Lake Erie region (Carter and Hites 1992, Barclay and Zingo 1993, Robertson et al. 1993). Considering that zebra mussels bioaccumulate organic contaminants more than native Great Lakes’ bivalves (Brieger and Hunter 1993), and that they constitute between 90 and 99% of benthic invertebrate biomass (Dahl et al. 1995), it is unlikely that scaup encounter a food supply as contaminated and as abundant as that of Dreissena in western Lake Erie. Local exposure to contaminants also is suggested by chemical profiles for all three species of Lake Erie waterfowl. While concentrations of compounds differed among species, the proportional abundance of 33 congeners to total PCB concentration was consistent among species and similar to zebra mussels from Lake Erie (Mazak 1995).

Concentrations of contaminants in liver and wing were highly correlated for all but 6 of the 39 compounds studied. Mean total lipid-adjusted PCB levels were an average of 1.23 times higher in liver than in wing muscle. The correlation between concentrations of contaminants in liver and muscle was lowest for low-$K_{ow}$ compounds (e.g., QCB, PCB 28), reflecting the rapid depuration kinetics of these contaminants from liver (Morrison et al. 1995).

Many studies have demonstrated that availability of Dreissena prey can influence waterfowl abundance and distribution (Stańczykowska et al. 1990, Wormington and Leach 1992, Cleven and Frenzel 1993, Mitchell and Carlson 1993). In turn, migrant or resident waterfowl may have significant impacts on mussel population biomass and size structure (Stańczykowska et al. 1990, Cleven and Frenzel 1993). The size of mussels consumed may affect waterfowl exposure to contaminants because contaminant concentrations are higher in large than in small Dreissena (H. Morrison, personal communication). Waterfowl that feed on large mussels should experience greater exposure to contaminants. In this study, Greater Scaup consumed larger mussels and had higher contaminant concentrations than co-occurring Lesser Scaup and Bufflehead (Fig. 3; Table 1). Waterfowl exposure to organochlorine contaminants may, therefore, vary both as a function of the trophic level at which individuals feed, and by specialized (i.e., size-specific) feeding strategies employed by individual predators (Connolly and Pederson 1988, Koslowski et al. 1994). It is also possible that physiological differences among waterfowl species could account for some of the observed variation in contaminant concentrations. Observed differences were not related to sex or age of waterfowl studied (Table 1).

The relationship between contaminants and diet in scaup was site dependent, as mussel consumers from Fighting Island were more contaminated than individuals that ate macrophytes, but less contaminated than individuals from Lake Erie that consumed mussels (Table 1; Fig. 2). The latter pattern was surprising because Dreissena from Fighting Island were more contaminated than mussels from Lake Erie for a number of congeners (Table 1). This pattern may have resulted from subtle diet differences, as Lake Erie ducks consumed >97% mussels, while those from Fighting Island also consumed macrophytes (15–24%) (Fig. 2).

Differences in contaminant concentrations in conspecific plant and mussel consumers from Fighting Island were most pronounced for highly hydrophobic congeners (Table 1). Congeners such as PCBs 180, 194, and 206 were between 18 and 23 times higher in scaup that consumed Dreissena as a principal food than in those that ate mainly plants. Bufflehead that consumed mussels were also more contaminated with mid- and high-$K_{ow}$ congeners than those that ate mainly plants (Table 1). Bufflehead from Fighting Island had the highest total PCB concentration among waterfowl that consumed mainly macrophyte diets (Table 1). These individuals consumed 20% animal matter and had stable isotope values consistent with Dreissena diet (Table 1; Fig. 5). Highly hydrophobic compounds like PCBs 194 and 206 have very high-$K_{ow}$ values and correspondingly low elimination rate constants. These compounds are retained in biota to a higher degree than low-$K_{ow}$ congeners that are more easily excreted (Brieger and Hunter 1993, Morrison et al. 1995).

Herring Gulls nesting on Middle Sister Island (Lake Erie) have total PCB concentrations at least one order of magnitude higher than Lake Erie scaup from this study (G. Fox, personal communication). Greater and Lesser Scaup wet mass $p,p'$-DDE (dichlorodiphenyl-dichloroethene) levels are also at least two orders of magnitude lower than those reported for Double-crested Cormorant and Caspian Tern eggs from a variety of Great Lakes locations (Yamashita et al. 1993). These results are not surprising considering that fish-eating birds feed at a higher trophic level than waterfowl that consumed mussels in this study.
Peterson and Fry (1987) reported that δ¹⁵N ratios typically increase at least 3‰ with each increase in trophic level. Evidence from this study illustrates that trophic-level enrichment of δ¹⁵N occurred in Lake Erie waterfowl relative to *Dreissena* prey, averaging 2.6‰ in Lesser Scaup and Bufflehead and 2.3‰ in Greater Scaup (Fig. 6). Lesser and Greater Scaup from Fighting Island that consumed mussels were enriched 2.3 and 1.3‰ relative to *Dreissena*, respectively. These ducks also were enriched by 2.5 and 1.4‰, respectively, relative to co-occurring conspecifics that consumed mainly macrophytes (Fig. 6). These examples of trophic enrichment of nitrogen isotopes in waterfowl complement contaminant data.

Stable isotope results for waterfowl that consumed macrophyte diets were less clear and, in some cases, in apparent contradiction to results for contaminants. For example, Redhead, Canvasback, and Bufflehead from Fighting Island had low contaminant concentrations and diets dominated (≥88.6%) by macrophytes, yet each species exhibited δ¹⁵N enrichment characteristic of *Dreissena* diet (Table 1; Fig. 5). Low contaminant concentrations in these ducks are consistent with recent consumption of macrophytes, while elevated δ¹⁵N values may reflect recent (i.e., within the past month) utilization of animal prey. Consistent with this view, Custer and Custer (1996) determined that *Dreissena* comprised between 1.3 and 50.5% of diets of Redhead and Canvasback captured on Lakes Erie and St. Clair. Hobson and Clark (1992) reported a half-life for isotopes examined of ~12.5 d for inactive ducks. Considering that all ducks surveyed here were active prior to being killed, and that active individuals have rapid turnover of proteins and of stable isotopes, 12.5 d should be treated as the maximum possible half-life (Hobson 1990; K. Hobson, personal communication). Therefore, diet and contaminant analyses indicate a vegetative diet for these waterfowl, although δ¹⁵N values and literature reports suggest that these taxa may have consumed mussels in the recent past. Because diets of Canvasback and Redhead ducks are typically composed of vegetative matter (E. Mazak, personal observation), periodic opportunistic feeding on animals (e.g., *Dreissena*) could have a large but ephemeral impact on δ¹⁵N values but not contaminant levels.

δ¹³C profiles of waterfowl that consumed *Dreissena*, as well as those suspected of eating animal matter (Redhead, Canvasback, and Bufflehead), are consistent with a *Dreissena* dietary component (Fig. 5). However, δ¹³C profiles of Mallard and Greater and Lesser Scaup from Fighting Island that consumed mainly macrophytes were not consistent with those expected of animals with a macrophyte diet (Fig. 5). Rather, δ¹³C levels in these ducks were highly variable but most similar to those of epiphytic algae and intermediate to those of terrestrial plants (~28‰) and macrophytes (approximately ~10‰) (Peterson and Fry 1987, Hamilton et al. 1992; R. Hesslein, unpublished data). Alternatively, δ¹³C signatures in these waterfowl may have been affected by earlier consumption of *Dreissena* prey.

**Reproductive impairment**

Reproductive problems associated with organochlorine contaminants have been reported for a number of waterbirds on the Great Lakes (see Fox 1993). Concern has centered on the possibility that waterfowl that consume contaminated *Dreissena* may experience reproductive impairment owing to accumulation of high contaminant burdens. Reproductive success in Tufted Ducks fed contaminated *Dreissena* was 60% lower than that of ducks fed less contaminated mussels (de Kock and Bowmer 1993). Lake Erie Greater Scaup, the most contaminated group of waterfowl in this study, had liver concentrations of PCBs 138, 153, and 180 that were between 17 and 54% of the levels found in the reproductively impaired Tufted Ducks. It is not clear whether any of the species of waterfowl included in this study have experienced contaminant-induced reproductive impairment. The number of Greater and Lesser scaup documented in all flyways of the United States and Canada during midwinter counts declined steadily between the late 1960s and the mid-1980s, though numbers appeared to increase along the Atlantic coast during the late 1980s (see Barclay and Zingo 1993). Recent surveys indicate that both Bufflehead and Common Goldeneye have declined in abundance in eastern North America (G. Fox, personal communication). Greater Scaup fly through, among other areas, the Lake Erie region during migrations to and from arctic breeding grounds. During 1993, Greater Scaup suffered an unusually high (37.5%) rate of reproductive failure in Alaska (J. Barclay, personal communication). However, it has not been established whether these scaup had previously staged on Lake Erie nor whether their diet included *Dreissena*. Lesser Scaup chicks reared on a diet of *Dreissena* from Middle Sister Island had depressed vitamin A levels and a compromised immune system (C. Tessier, personal communication). These chicks accumulated a total PCB concentration of 1600 μg/kg lipid, whereas values in Lake Erie Greater Scaup, Lesser Scaup, and Bufflehead were 4464, 2734, and 3722 μg/kg lipid, respectively. Even though waterfowl from Lake Erie that consumed *Dreissena* accumulated higher levels of PCBs than captive individuals that exhibited metabolic impairment, additional work is required to determine whether wild populations of waterfowl that frequent the Great Lakes are adversely affected by contaminated food.

In summary, three taxa of waterfowl in the lower Great Lakes that utilize *Dreissena polymorpha* as a primary food source exhibit elevated tissue concentrations for a broad suite of organochlorine congeners relative to individuals with macrophyte diet. Stable isotope analyses of conspecifics of waterfowl with *Dreissena* and macrophyte diets revealed trophic en-
richment in individuals with mussel diet. It is not yet clear whether any of the waterfowl taxa that exploited *Dreissena* experience adverse health effects in consequence.

Invasion of Lake Erie by zebra mussels has resulted in dramatic and rapid shifts in food web and contaminant dynamics (MacIsaac 1996). These changes parallel in strength, and may exceed in duration, responses by lakes subjected to intentional or unplanned bio-manipulation of fishes (e.g., Vanni et al. 1990, Scheffer et al. 1993, Carpenter et al. 1996). Dispersal and establishment of *Dreissena* in lakes and rivers throughout eastern North America may portend significant changes in many invaded systems, and provide opportunities for ecologists to address questions relating to community assembly, food web interactions, and contaminant dynamics.

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