Bioaccumulation of mercury in invertebrate food webs of Canadian Rocky Mountain streams

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Abstract: Methylmercury (MeHg) is a contaminant of concern because of its ability to biomagnify in aquatic food webs, resulting in potentially harmful concentrations in higher consumers. Beaver impoundments in the southern Canadian Rockies release bioavailable MeHg to downstream food webs. We examined the magnitude of uptake and trophic transfer of exported Hg (total Hg [THg] and MeHg) to grazing and predatory invertebrates and controls on these transfers by site-specific (dissolved organic C [DOC], MeHg in water, MeHg in diet) and individual (body size, trophic level) variables. Bioconcentration factors (BCFs) were high (79,756 ± 68,204) relative to values from the literature, declined with increasing MeHg in water, and did not differ above and below beaver ponds. Biomagnification factors (BMFs) for uptake from periphyton to grazers (18.3 ± 11.7) were high and increased with increasing DOC, BMFs from grazers to predators (2.1 ± 1.2) were low, and neither of these BMFs differed above and below ponds. Invertebrate body size had no effect on MeHg concentration. However, the relative difference in trophic level from prey to consumer was an important driver of BMFs, and MeHg in the diet was negatively associated with BMFs. Rates of uptake and transfer were greater at low than at higher concentrations, but overall trophic magnification through the food web (average trophic magnification factor = 2.3) was on the lower end of the typical range observed worldwide. Thus, the limited risk to wildlife and humans who consume fish from these systems is a result, in part, of a dampening of bioaccumulation as it is transmitted through these invertebrate food webs that exhibit weak size structuring relative to other aquatic systems.

Key words: methylmercury, invertebrates, trophic transfer, streams, bioaccumulation, biomagnification, stable isotopes, food webs

Bioaccumulation, the process by which an organism attains a higher concentration of a chemical relative to its environment (Borgå et al. 2011), is highly variable among chemicals, organisms, and ecosystems. Bioaccumulation is a key endpoint criterion identified during hazard assessment, which traditionally attempts to define persistence, bioaccumulation, and toxicity (PBT). Bioconcentration and biomagnification are 2 components of bioaccumulation that are used to define rates of uptake and subsequent trophic transfer of a chemical. Bioconcentration in aquatic organisms is the uptake of a chemical across respiratory/dermal surfaces from the water. The bioconcentration factor (BCF) is the ratio of the concentration of a substance in an organism to that in the water. Biomagnification is a special case of bioaccumulation that occurs when the chemical concentration is greater in an organism than in its prey because dietary absorption occurs faster than elimination (Borgå et al. 2011). The biomagnification factor (BMF), sometimes called the trophic transfer factor (TTF; DeForest et al. 2007), is the ratio of the concentration of a substance in an organism to that in its diet (Gobas et al. 2009). The sum of these processes yields the bioaccumulation factor (BAF), the ratio of the concentration of a substance in an organism to that in the environment including both water and dietary sources (Gobas et al. 2009). BCF and BAF are bioconcentration and bioaccumulation endpoints that have been used in the development of environmental guidelines for risk assessment (Arnot and Gobas 2006). For example, Environment Canada identifies substances with BCF and BAF values ≥5000 as bioaccumulative under the Canadian Environmental Protection Act (CEPA 1999).

Some trace elements, such as Hg, are persistent in fresh waters, and their bioavailability is heavily influenced by geochemical factors (DeForest et al. 2007). Therefore, rates of Hg bioaccumulation are highly variable among locations (Lavoie et al. 2013) and defining critical PBT levels can be difficult (DeForest et al. 2007). For Hg and other trace elements, such as Se, Zn, Cu, and Pb, BCFs tend to be highest (indicating hazard) at low concentrations of exposure (low potential for toxicity) and lowest (indicating reduced haz-
ard) at higher concentrations of exposure (high potential for toxicity). This tendency complicates risk assessments and suggests that exposure concentration may contribute as much as other metal- and species-specific determinants of bioaccumulation to BAFs (McGeer et al. 2003, DeForest et al. 2007). Therefore, BCFs and BAFs are important for describing the link between exposure concentrations and tissue concentrations. For example, high exposure concentrations may be of little consequence if the rate of transfer is low, leading to absent or minimal toxic effects.

MeHg is a potent neurotoxin with the potential to pose a serious health risk to humans and wildlife (Mergler et al. 2007). Thus, understanding the potential for MeHg to transfer from abiotic to biotic compartments and move further up the food web to higher trophic levels is imperative. Based on a large compilation of available data from peer-reviewed literature and technical documents, DeForest et al. (2007) reported that BAFs for MeHg range from 100,000 to 48,000,000 across species and trophic levels, including invertebrates, small fish, and large fish. This wide range highlights the importance of understanding MeHg bioaccumulation and biomagnification and their contributing variables in aquatic systems.

Levels of MeHg are elevated in water downstream from impoundments (Roy et al. 2009), and this elevation leads to higher downstream concentrations in algae and invertebrates (Painter et al. 2015). Increases in MeHg concentration downstream from reservoirs, including North American beaver (Castor canadensis) impoundments, are thought to be the consequence of flooding of organic matter, which enhances microbial decomposition and, therefore, Hg methylation (St Louis et al. 2004, Roy et al. 2009). The magnitude of change from up- to downstream varies among ecological compartments and sites and ranges from a 0.5 to 2.5 × increase in algae to a 0.6 to 5.0 × increase in predatory invertebrates (Painter et al. 2015). These ranges indicate that other factors may modulate responses within the food web. For example, concentrations in water and the diet are inversely related to BCFs and BMFs, respectively, in laboratory and field studies (DeForest et al. 2007), and low pH can lead to higher BMFs in some organisms (Jardine et al. 2013).

Concentration of dissolved organic C (DOC), an indicator of wetland influence (Brigham et al. 2009, Chasar et al. 2009), explains additional variability in MeHg concentrations among sites (Painter et al. 2015), but relationships between DOC and MeHg transport and uptake are complex. For example, increased DOC concentrations in the water column of Adirondack lakes resulted in a negative correlation between DOC and BAF for Yellow Perch (Perca flavescens) (Dittman and Driscoll 2009). Dittman and Driscoll (2009) hypothesized that the negative correlation was caused by binding between DOC in the water column and MeHg, which reduced the bioavailable pool. Other studies illustrate the importance of DOC in transporting MeHg into aquatic systems (Adams et al. 2009, Brigham et al. 2009). Positive correlations between DOC concentrations and MeHg in the water column (Brigham et al. 2009) and in basal foodweb organisms (Adams et al. 2009) are common. Therefore, how DOC controls uptake and transfer of MeHg should be investigated.

We measured uptake and accumulation of MeHg in stream food webs along a MeHg gradient (up- to downstream of beaver impoundments; Painter et al. 2015) by examining the movement of this trace element among environmental compartments (water, periphyton, and herbivorous and predatory invertebrates). We used BCFs, BAFs, and BMFs as described above, and we examined other aspects of the invertebrate food web that can influence MeHg concentrations. First, we assessed trophic transfer of MeHg with the aid of N stable-isotope ratios (15N/14N; hereafter, δ15N) to identify feeding linkages in stream food webs. The habitat-independent, stepwise enrichment of 15N with trophic level has been used widely to characterize aquatic food webs (Minagawa and Wada 1984), and the relationship between δ15N and environmental contaminants, such as MeHg, has long been used as a predictor of contaminant biomagnification (Kidd et al. 1995). Second, we evaluated the relationship between body size, trophic level, and MeHg concentrations. Body size is directly correlated with MeHg concentrations in fishes because larger fishes tend to be older or feed at higher trophic levels (Kidd et al. 1995, Gewurtz et al. 2011). However, invertebrates are rarely classified based on their size in contaminant studies (Watanabe et al. 2008) because trophic-level estimates vary widely within functional feeding groups (Merritt and Cummins 1996). We expected positive relationships between δ15N, body size, and MeHg because predators are consistently larger than their prey (Riede et al. 2011) and because predators tend to occupy higher trophic levels than their prey (Anderson and Cabana 2009). Taken together, we sought to estimate the importance of physicochemical variables, such as DOC and MeHg concentrations in water and the diet, at the base of the food web and the relationships of these processes with bioconcentration and trophic transfer of MeHg.

METHODS
Field sampling and laboratory methods
From mid-July to early August 2012, we used methods described by Painter et al. (2015) to sample 15 streams with in-channel beaver impoundments and 6 free-flowing streams assumed to be uninfluenced by beaver activity on the eastern slope of the Canadian Rocky Mountain and Foothill Regions, Alberta. The streams had no known local point sources of Hg. Most streams were in conifer-dominated watersheds at elevations of 1300 to 2150 m. Sites had clear water, a high gradient with cobble and gravel substrates, moderately basic pH, and limited accumulation of leaf litter because of low canopy cover, typically <20% (Painter et al. 2015). At each sampling location, we collected water,
periphyton, and aquatic macroinvertebrates. At dammed sites, collections were made immediately above and below the impoundment (i.e., at an inflow and an outflow). Samples for THg and MeHg, DOC, total P (TP), and benthic chlorophyll-a (chl a) were collected, processed, and analyzed as described by Painter et al. (2015).

To assess body size, we sorted invertebrates, identified them to family level (Merritt and Cummins 1996), and counted and weighed them prior to drying them at 50°C for 48 h. When they were dry, we weighed invertebrates again and homogenized them using an acid-washed glass mortar and pestle. We calculated invertebrate body size as the average wet mass of individuals from 3 replicate samples collected at each site. We analyzed homogenized samples of taxa dominant across all sites, including grazing mayflies from the families Heptageniidae, Ephemerellidae, and Baetidae and predatory invertebrates from the families Perlodidae and Rhypacophilidae, as pooled samples for THg and MeHg and stable isotopes (see Painter et al. 2015 for detailed methods and quality assurance/quality control). We also analyzed all organisms collected from upstream and downstream locations at 2 beaver-affected sites (hereafter, comprehensive foodweb sites) to develop an understanding of MeHg behavior in more comprehensive invertebrate food webs in these systems. Additional taxa analyzed at those sites were Hydropsychidae, Limnephilidae, Phryganeidae, Tipulidae, Chironomidae, Siphlonuridae, Polycentropodidae, and Peltoperlidae.

Calculations

Uptake and trophic transfer are calculated as 3 factors: BCF, BMF, and BAF. BCF for animals can be measured only under controlled laboratory conditions (i.e., steady state) in which dietary uptake is deliberately excluded (Arnot and Gobas 2006, Gobas et al. 2009), but for autotrophs, such as periphyton with a single route of uptake, BCF can be calculated from field data as the ratio of the chemical concentration in the organism (C_B) to the chemical concentration in the water (C_W) as:

$$BCF = \frac{C_B}{C_W}. \quad \text{(Eq. 1)}$$

Biomagnification is the process by which the thermodynamic activity of the chemical in an organism exceeds that in its diet (Gobas and Morrison 2000, Arnot and Gobas 2006). BMF for organometals, such as MeHg, that are almost exclusively derived from the diet (Hall et al. 1997) is expressed as the ratio of C_B to the concentration in the diet (C_D) as:

$$BMF = \frac{C_B}{C_D}. \quad \text{(Eq. 2)}$$

BMF > 1 indicates that the chemical is a probable bioaccumulative substance (Gobas et al. 2009). For these analyses, we classified Heptageniidae, Ephemerellidae, and Baetidae as herbivorous grazers, and Perlodidae and Rhypacophilidae as predatory grazers and assumed that each trophic level fed entirely on the trophic level below it. We pooled the 3 mayfly taxa, and we pooled the 2 predatory taxa because MeHg concentrations were not statistically different among taxa within these groups (Painter et al. 2015). We used stable N isotope data to confirm feeding linkages (described below) and to account for potential omnivory.

Bioaccumulation, the sum of all exposure routes for animals (Arnot and Gobas 2006), is expressed as:

$$BAF = \frac{C_B}{C_W}. \quad \text{(Eq. 3)}$$

For chemicals with an octanol–water partition coefficient (logKOW) ≤ 5, BCFs and BAFs ≤ 5000 L/kg wet mass indicate absence of biomagnification potential in water-respiring organisms (Gobas et al. 2009).

Stable isotopes

δ15N increases with trophic level, so it can be used to compare Hg biomagnification across systems as the slope of the regression of log-transformed MeHg vs δ15N or trophic level (Borgå et al. 2011, Lavoie et al. 2013). Calculation of a trophic-level value takes into consideration a trophic enrichment factor (TEF; increase in δ15N from diet to consumer) and assigns a discrete trophic level (TL) to a baseline organism (e.g., periphyton TL = 1) because δ15N can vary widely among sites (Caban and Rasmussen 1996). δ15N was converted to trophic level as:

$$\text{TL}_{\text{consumer}} = \frac{\delta^{15}N_{\text{consumer}} - \delta^{15}N_{\text{baseline}}}{\text{TEF}} + \text{TL}_{\text{baseline}} \quad \text{(Eq. 4)}$$

where TL_consumer = trophic level of the consumer and TL_base = trophic level of the baseline organism. We used periphyton as the baseline organism with TL = 1. We chose a TEF of 2.0‰ (McCutchan et al. 2003), a value suitable for stream food webs (Bunn et al. 2011). In an analysis of 144 records across arid, tropical, subtropical, and temperate sites, Bunn et al. (2013) found that many TEF estimates for invertebrate food webs were lower than the commonly reported 3.4‰. We chose a 2.0‰ value based on similar TEFs from the literature for stream invertebrate food webs (McCutchan et al. 2003, Bunn et al. 2013).

Trophic magnification factors (TMFs) represent the average biomagnification per TL through the entire food web and are calculated as the antilog of the slope (b) of a regression of logMeHg vs TL as follows (Borgå et al. 2011):

$$\text{TMF} = 10^b. \quad \text{(Eq. 5)}$$

When TMF = 1 (b = 0), MeHg does not biomagnify on average through the food web; when TMF > 1 (b > 0),
MeHg biomagnifies through the food web by an average of TMF/TL; and when TMF < 1 \((b < 0)\), MeHg decreases by an average of TMF/TL (Borgå et al. 2011).

Trophic magnification slopes (TMSs) also can be derived directly from the logMeHg vs \(\delta^{15}N\) regression and can be compared across systems without standardizing to a baseline (Lavoie et al. 2013). The TMS is calculated as the slope \((b)\) in the following model:

\[
\log\text{MeHg} = b\delta^{15}N + a. \tag{6}
\]

No change in logMeHg concentration with increasing \(\delta^{15}N\) produces TMS = 0, which indicates no biomagnification. An increase in logMeHg concentration with increasing \(\delta^{15}N\) produces TMS > 0, which indicates biomagnification. A decrease of logMeHg concentration with increasing \(\delta^{15}N\) produces TMS < 0, which indicates trophic dilution, the opposite of biomagnification (Gobas et al. 2009).

We were able to calculate BCFs, BArs, and BMFs for all 36 sites (hereafter, expressed as log-transformed values unless otherwise specified). However, a regression model for each site is needed to calculate TMSs and TMFs. Our sample size within sites was too small to calculate regressions, except at up- and downstream locations at the 2 comprehensive foodweb sites. Instead, we calculated overall TMF values for all sites combined into a single regression and then back-calculated TMS to compare to other studies (Lavoie et al. 2013) by dividing logMeHg vs TL slopes by the TEF.

Data analysis

We used paired \(t\)-tests to compare BCFs, BArs, and BMFs for all 36 sites (hereafter, expressed as log-transformed values unless otherwise specified). However, a regression model for each site is needed to calculate TMSs and TMFs. Our sample size within sites was too small to calculate regressions, except at up- and downstream locations at the 2 comprehensive foodweb sites. Instead, we calculated overall TMF values for all sites combined into a single regression and then back-calculated TMS to compare to other studies (Lavoie et al. 2013) by dividing logMeHg vs TL slopes by the TEF.
Table 1. Methylmercury (MeHg) concentrations and trophic levels (TLs) for all biological compartments in invertebrate food webs in Rocky Mountain Foothills streams, and corresponding bioconcentration factors (BCFs), bioaccumulation factors (BAFs), and biomagnification factors (BMFs) from water to periphyton (periphyton/water), water to grazers (grazers/water), water to predators (predators/water), periphyton to grazers (grazers/periphyton), and from grazers to predators (predators/grazers). BCFs, BAFs, and BMFs were log(x)-transformed. Periphyton TL is assumed to be 1. US at the end of a site code indicates sites upstream from beaver impoundments. DS indicates sites downstream from beaver impoundments. No suffix indicates sites without beaver activity.

<table>
<thead>
<tr>
<th>Site</th>
<th>Water MeHg (ng/L)</th>
<th>Periphyton MeHg (ng/g)</th>
<th>Grazer MeHg (ng/g)</th>
<th>Predator MeHg (ng/g)</th>
<th>Grazer TL</th>
<th>Predator TL</th>
<th>Log BCF (periphyton/water)</th>
<th>Log BAF (grazers/water)</th>
<th>Log BAF (predators/water)</th>
<th>Log BMF (grazers/periphyton)</th>
<th>Log BMF (predators/grazers)</th>
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<td>10.10</td>
<td>0.85</td>
<td>2.09</td>
<td>4.80</td>
<td>5.78</td>
<td>6.00</td>
<td>0.98</td>
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<td>–</td>
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described as the independent effect of variable $x_i$ given that model $k$ is the best approximating model. To incorporate uncertainty in the best approximating model, $I_{\text{weighted}}$ is calculated as:

$$I_{\text{weighted}}(x_i) = \sum_{k=1}^{K} w_k I_{\text{global}}(k, i)$$  \hspace{1cm} (Eq. 7)

where $w_k$ is the model weight as calculated from the AIC. Giam and Olden (2015) recommended reporting both metrics because $I_{\text{weighted}}$ attenuates the reduction in precision and increase in bias introduced by the incorrect inclusion/exclusion of spurious/genuine variables when the global model is used, whereas $I_{\text{global}}$ exhibits greater accuracy and precision in estimating the importance of stronger explanatory variables. $I_{\text{global}}$ also is favored in the case of small data sets with correlated explanatory variables.

We carried out statistical analyses with SYSTAT (version 13; Systat Software, San José, California) and R (version 3.2.2; R Project for Statistical Computing, Vienna, Austria) using the NADA package (Lee (2013) for the robust ROS and the relaimpo package (Grömping 2015) for RVI metrics. All data except TL were log($x$)-transformed prior to all of the above analyses.

RESULTS

General bioaccumulation trends

BCFs were in the upper ½ of the distribution of literature values for fresh waters, whereas BMFs and BAFs were generally low compared with values published in the literature (Table 1). Untransformed BCFs ranged from ~12,000 to 344,000, and untransformed BMFs ranged from ~3.8 to 48 for $BMF_{(\text{grazers}/\text{periphyton})}$ and 0.9 to 10 for $BMF_{(\text{predators}/\text{grazers})}$. BCFs, BAFs, and BMFs (log($x$)-transformed) did not differ between up- and downstream locations at beaver-affected sites ($p > 0.05$ for all comparisons; Table 2).

Mean δ$^{15}$N values were 2.5 ± 1.5% in periphyton, 3.0 ± 1.6% in grazing mayflies and 4.4 ± 1.5% in predatory invertebrates resulting in low TLs for mayflies and predators of 1.2 ± 0.5 and 2.1 ± 0.6, respectively. The slope of the logMeHg vs TL regression for all 21 sites combined was 0.34 and the slope of the logTHg vs TL regression was 0.08 (Fig. 1), resulting in $\text{TF}_{1/2}$s of 2.3 for MeHg and 1.2 for THg. These values correspond to $\text{TMS}$s of 0.18 for MeHg and 0.03 for THg. At the 2 comprehensive food web sites, MeHg TMSs were higher up- (BEV004US TMS = 0.33, BEV028US TMS = 0.27) than downstream (BEV004DS TMS = 0.01, BEV028DS TMS = 0.14) from beaver impoundments despite higher overall concentrations downstream (Fig. 2A, B). These relationships were not significant for 3 of 4 regressions (BEV004US: $r^2 = 0.61$, $p = 0.022$; BEV028US: $r^2 < 0.01$, $p = 0.957$; BEV004DS: $r^2 = 0.39$, $p = 0.099$; BEV028DS: $r^2 = 0.21$, $p = 0.182$). Confidence intervals (CIs) around slope estimates overlapped because of high variability and low sample size.

Variables contributing to Hg transfer within food webs

The logMeHg vs log-transformed mean body size (slope = –0.039, $r^2 < 0.01$, $p = 0.764$; Fig. 3A) and the TL vs body size (slope = 0.113, $r^2 = 0.02$, $p = 0.109$; Fig. 3B) relationships were not significant. The logMeHg vs TL relationship was weak but significant (slope = 0.137, $r^2 = 0.06$, $p = 0.004$; Fig. 3C).

For model set A ($BMF_{(\text{grazers}/\text{periphyton})}$), the top 4 models had $\Delta \leq 2$. The best model contained only TL$^{(\text{grazers}/\text{periphyton})}$ and the $2^{nd}$ best model contained TL$^{(\text{grazers}/\text{periphyton})}$ and DOC. Evidence ratios (ERs) of the $2^{nd}$-best model was only 1.034, suggesting almost equal weight between the 2 top models (Table 3). TL$^{(\text{grazers}/\text{periphyton})}$ and DOC had the strongest averaged parameter estimates (0.287 ± 0.121 and 0.381 ± 0.199, respectively). The averaged parameter estimates of MeHgdiet (periphyton MeHg concentration), TP, and chl $a$ were –0.317 ± 0.379, –0.296 ± 0.250, and 0.047 ± 0.116, respectively, but the large SEs made them weak predictors of $BMF_{(\text{grazers}/\text{periphyton})}$. The intercept-only (null) model had a $\Delta = 3.9$, making it a plausible model, but the top 4 models had a summed weight of 0.809 (accounted for 80.9% of the weight in the set) compared to only 3.7% for the intercept-only model. RVI metrics showed that the largest proportion of variance was accounted for by TL$^{(\text{grazers}/\text{periphyton})}$ and DOC (Table 4). The total variance explained was 33.4% for the $I_{\text{weighted}}$ model parameters and 48.7% for the $I_{\text{global}}$ model parameters.

For model set B ($BMF_{(\text{predators}/\text{grazers})}$), the top 2 models had $\text{AIC} \leq 2$ (Table 3). The top model contained MeHgdiet (here, grazer MeHg concentration) and DOC, and the $2^{nd}$-best model contained MeHgdiet, DOC, and benthic chl $a$. MeHgdiet and DOC had the strongest averaged parameter estimates (–0.631 ± 0.107 and 0.379 ± 0.119, respectively). The averaged parameter estimates for chl $a$ and TL$^{(\text{predators}/\text{grazers})}$ were 0.120 ± 0.055 and 0.084 ± 0.035, respectively, and TP had no support. The largest proportion of variance was accounted for by MeHgdiet. In the global model that accounted for 85.7% of the total variance, 79.7% was partitioned among MeHgdiet, TL$^{(\text{grazers}/\text{periphyton})}$ and DOC (Table 4), but when partitioned-$R^2$ was weighted, most of the variance (74.4% of the total 83.8%) was explained by MeHgdiet, reflecting the estimated top model. This finding could be a case of a weak correlation between explanatory variables in which the variance of the stronger variable (MeHgdiet) was overestimated by $I_{\text{weighted}}$. As such, results of both $I_{\text{global}}$ and $I_{\text{weighted}}$ should be considered.

Results of regressions between bioaccumulation measures and individual MeHg transfer variables (Fig. 4A–H) confirmed significant relationships ($p < 0.05$) between $BCF_{(\text{periphyton}/\text{water})}$ and water MeHg ($r^2 = 0.31$, $p = 0.008$; Fig. 4A), $BMF_{(\text{predators}/\text{grazers})}$ and MeHggrazers ($r^2 = 0.68$, $p =$
The far larger BCF than BMF values illustrate that processes at the base of the food web have a greater influence than trophic transfer on overall MeHg concentrations of higher consumers. At all 35 sites in our study, the magnitude of change in MeHg was greatest from water to organisms and declined from primary producers to consumers and further again from primary consumers to predators. This pattern is consistent with results of other studies, which showed that bioconcentration from water to algae is several orders of magnitude larger than biomagnification from algae to successively higher trophic levels (Weiner et al. 2007, Chasar et al. 2009) and places renewed emphasis on understanding controls on the delivery and uptake of MeHg from water to periphyton in aquatic systems.

Impoundment of stream channels by beavers increases baseline MeHg concentrations (Painter et al. 2015). However, beaver effects were partly offset by reductions in MeHg transfer among trophic levels within the food web when baseline MeHg concentration is high, and BCFs and BAFs did not change from up- to downstream of the ponds. Potential explanations include the high variability in water MeHg concentrations entering the ponds (Painter et al. 2015) and the strong effect of water MeHg concentrations on rates of uptake (Fig. 4A). The influence of the increase of water MeHg concentrations might have arisen from the absence of an accompanying increase of other basal foodweb resources (nutrients, algal and invertebrate biomass) downstream of the impoundments (Painter et al. 2015), which might have altered bioaccumulation via growth dilution (Ward et al. 2010). Nevertheless, MeHg concentrations in water and the

Table 2. Mean (±SE) upstream and downstream log(x)-transformed bioaccumulation measures (as described in Table 1) at beaver-affected sites.

<table>
<thead>
<tr>
<th>Bioaccumulation measure</th>
<th>Upstream</th>
<th>Downstream</th>
</tr>
</thead>
<tbody>
<tr>
<td>log BCF(grazers/periphyton)</td>
<td>4.79 ± 0.10</td>
<td>4.74 ± 0.09</td>
</tr>
<tr>
<td>log BAF(grazers/water)</td>
<td>6.02 ± 0.09</td>
<td>6.00 ± 0.09</td>
</tr>
<tr>
<td>log BAF(predators/water)</td>
<td>6.09 ± 0.15</td>
<td>6.21 ± 0.14</td>
</tr>
<tr>
<td>log BMF(grazers/periphyton)</td>
<td>1.18 ± 0.09</td>
<td>1.22 ± 0.07</td>
</tr>
<tr>
<td>log BMF(predators/grazers)</td>
<td>0.27 ± 0.08</td>
<td>0.25 ± 0.05</td>
</tr>
</tbody>
</table>

DISCUSSION

Our work adds to the growing body of knowledge about how Hg liberated by beaver impoundments is modified as it is transferred through food webs and offers an important contrast to work carried out at low-pH, high-nutrient streams in well vegetated watersheds (e.g., Roy et al. 2009). MeHg uptake from abiotic to biotic compartments was high, but rates of transfer within the invertebrate food webs of these Rocky Mountain streams were low. Mean untransformed BCFs were 79,756 ± 68,204, which are high compared to literature values for MeHg reported by McGeer et al. (2003) for a range of organisms including algae, invertebrates, and fish (8952 ± 24,675; n = 53). Our values probably are not indicative of hazard of toxicity but instead reflect natural conditions in which uptake is high when ambient concentrations are low (DeForest et al. 2007, Cardwell et al. 2013). Bioaccumulation factors (mean BAF(grazers/water) = 1,259,951 ± 838,812, mean BAF(predators/water) = 1,969,288 ± 1,541,130) were within the range of literature BAF values for MeHg reported by DeForest et al. (2007) (100,000–48,000,000), but rates of trophic transfer within the food web were low. Our calculated mean MeHg TMS (0.18) was near the global average for MeHg in freshwater systems (0.24 ± 0.08), but our calculated TMS for THg (0.03) was well below the global average for THg in fresh water (0.15 ± 0.11) (Lavoie et al. 2013). Accordingly, TMFs (MeHg: 2.3, THg: 1.2) also were at the lower end of the range for freshwater systems (MeHg: 8.3 ± 7.5, THg: 4.3 ± 4.8; Lavoie et al. 2013).

Figure 1. Log Hg concentration vs trophic level for methylmercury (MeHg) (slope = 0.34) and total Hg (THg) (slope = 0.08) in Rocky Mountain stream food webs.
diet strongly influenced bioaccumulation, and BCFs and BMFs were low when MeHg available for uptake was high. pH effects on Hg concentrations in streams in New Brunswick, Canada, were damped in intermediate trophic levels (primary consumers and predators) of invertebrate food webs because of inefficient Hg transfer (low BMFs) when dietary MeHg concentrations were high (Jardine et al. 2013). In our study streams, DOC appeared in our top models for predicting MeHg accumulation with a positive coefficient, but its effect generally was weak and did not differ from up- to downstream of ponds (Painter et al. 2015). In other systems, DOC increased significantly from up- to downstream of beaver ponds (Roy et al. 2009), suggesting potential for greater MeHg biomagnification at downstream sites. Thus, in locations where baseline MeHg concentrations and nutrients are high, beaver activity may have wide-reaching impacts on stream ecosystems, but prevailing chemical conditions may lead to attenuated effects on MeHg transfer further up the food chain.

Data from the comprehensive foodweb sites suggested that MeHg was taken up by periphyton and invertebrates more efficiently at the lower upstream than at the higher downstream concentrations (Painter et al. 2015). These data are consistent with the inverse relationship between concentration and uptake observed for laboratory-reared organisms (Tsui and Wang 2004, DeForest et al. 2007) and grazers and predators in New Brunswick streams (Jardine et al. 2013). Tsui and Wang (2004) suggested that the inverse relationship might be explained by saturation of binding sites for MeHg at high concentrations combined with slow turnover. In our study, MeHg was readily taken up into the food web even when water concentrations were BDL, and BCFs were highest when concentrations were low. Our model analysis further indicates that MeHg<sub>diet</sub> was negatively related to BMFs, especially from grazers to predators. Lavoie et al. (2013) found that MeHg TMSs were lowest when baseline TL Hg and atmospheric Hg deposition were highest.

DOC and baseline MeHg concentrations were positively related in our study streams (Painter et al. 2015), a pattern that has been well described in aquatic systems (Driscoll et al. 1995). Wetlands are major locations of Hg methylation, and DOC in streams is associated with the presence of wetlands in the surrounding watershed. DOC–Hg complexes wash into streams from the terrestrial environment (Brigham et al. 2009, Chasar et al. 2009). However, DOC–Hg relationships are variable across systems. For example, our study streams receive low DOC inputs, and DOC is weakly positively associated with BMFs (e.g., BMF<sub>grazers/periphyton</sub> rises from ~3 to ~50 along the low DOC gradient; Fig. 4E). BMFs for Ephemeroptera and Plecoptera at streams in a higher-DOC landscape in New Brunswick (Jardine et al. 2012) were lower (mean ± SE = 11 ± 7; Jardine et al. 2013) and not related to DOC (data not shown). This comparison suggests that DOC enhances biomagnification of MeHg through the food web at low concentrations but may inhibit trophic transfer at higher concentrations. Driscoll et al. (1995) reported that MeHg in fish was positively correlated with DOC at low concentrations (<10 mg/L) but declined when DOC was very high (>30 mg/L).

The difference in TL from prey to consumer can affect BMFs. For example, some concentrations were unexplained by BMF alone in a study of contaminant biomagnification in an Arctic ecosystem (Dietz et al. 2000), and the authors speculated that trophic linkages played a role. The increase
in TL from prey to predator is associated with an increase in contaminant concentration; i.e., the TL–Hg relationship is significant overall (Kidd et al. 1995). Thus, trophic transfer of Hg should be greater when the step up in TL is greater for a given predator/prey pair. When calculating trophic transfer, care should be taken when selecting the TEF because TMFs are sensitive to changes in the TEF value used (Lavoie et al. 2013). For example, the use of a commonly applied TEF of 3.4‰ (Minagawa and Wada 1984, Post 2002) would result in TMFs of 4.06 for MeHg and 1.27 for THg in our food webs, effectively doubling the rate of biomagnification of MeHg from what we calculated based on a TEF of 2.0‰.

Hg bioaccumulation models were largely developed for pelagic freshwater systems (Watras et al. 1998), where size structuring by gape limitation in fishes is strong. However, these models do not necessarily apply to stream food webs, where invertebrates are diverse and occupy many different niches, which results in weak size structuring and short food chains (Vander Zanden and Fetzer 2007, Watanabe et al. 2008, Riede et al. 2011). For example, body size increased significantly with trophic position in stream invertebrate food webs in tropical, subtropical, and temperate regions (Jardine 2014), but the relationships were weak compared to relationships in strongly size-structured food webs commonly found in the pelagic zone of temperate lakes (Hairston

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**Figure 3.** Log methylmercury (MeHg) concentration vs log body size (A), trophic level vs log body size (B), and log MeHg concentration vs trophic level (C) for invertebrates at 21 Rocky Mountain streams ($n = 131$).
The implication of this weak size structuring is that larger animals may not necessarily have the highest Hg concentrations nor occupy the highest TLs (Watanabe et al. 2008). At the comprehensive foodweb sites in our study, Tipulidae have low MeHg concentrations, low TLs, and the largest body sizes, whereas smaller invertebrates, particularly grazing mayflies, have the highest MeHg levels suggesting that other factors drive accumulation of MeHg in these organisms. In a study by Mason et al. (2000), smaller invertebrates had higher concentrations of As and Se, probably because their high surface area:volume ratios increased absorptive surface area relative to that of larger organisms, but findings for Hg were inconsistent. The relative contributions of dermal absorption and dietary uptake are less well understood for small invertebrates than fishes (Hall et al. 1997).

Small-bodied omnivores can have higher-than-expected TLs. Invertebrates can exhibit both obligate and facultative functional feeding behavior and may undergo ontogenetic shifts in feeding habits (Cummins and Klug 1979). Hepta-

Table 3. Model sets used to account for variation in biomagnification factors (BMFs) for periphyton to grazers (model set A) and grazers to predators (model set B). $\Delta_i$ is the difference between the Akaike Information Criterion for small sample size (AICc) value of the best model and a competing model. $w_i$ is the Akaike weight, with the highest ranked model in the set having the highest $w_i$. Evidence ratios are a measure of likelihood of the best model over competing models. Models shown are those used for model averaging with $\Delta_i \leq 7$. TL = trophic level, DOC = dissolved organic C, TP = total P, chl $a$ = chlorophyll $a$. See Table 1 for other abbreviations.

<table>
<thead>
<tr>
<th>Model set</th>
<th>Model</th>
<th>$\Delta_i$</th>
<th>$w_i$</th>
<th>Evidence ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>A) BMF(grazers/periphyton)</td>
<td>TL(grazers/periphyton)</td>
<td>0.000</td>
<td>0.259</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>TL(grazers/periphyton), DOC</td>
<td>0.066</td>
<td>0.251</td>
<td>1.034</td>
</tr>
<tr>
<td></td>
<td>MeHgdiet, TL(grazers/periphyton), DOC</td>
<td>0.588</td>
<td>0.193</td>
<td>1.342</td>
</tr>
<tr>
<td></td>
<td>DOC</td>
<td>1.791</td>
<td>0.106</td>
<td>2.449</td>
</tr>
<tr>
<td></td>
<td>MeHgdiet, TL(grazers/periphyton)</td>
<td>3.138</td>
<td>0.054</td>
<td>4.803</td>
</tr>
<tr>
<td></td>
<td>MeHgdiet, DOC</td>
<td>3.317</td>
<td>0.049</td>
<td>5.252</td>
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<tr>
<td></td>
<td>Intercept only</td>
<td>3.909</td>
<td>0.037</td>
<td>7.059</td>
</tr>
<tr>
<td></td>
<td>MeHgdiet, TL(grazers/periphyton), DOC, chl $a$</td>
<td>4.402</td>
<td>0.029</td>
<td>9.032</td>
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<tr>
<td></td>
<td>MeHgdiet</td>
<td>6.421</td>
<td>0.010</td>
<td>24.792</td>
</tr>
<tr>
<td></td>
<td>DOC, TP, chl $a$</td>
<td>6.765</td>
<td>0.009</td>
<td>29.442</td>
</tr>
<tr>
<td>B) BMF(predators/grazers)</td>
<td>MeHgdiet, DOC</td>
<td>0.000</td>
<td>0.456</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>MeHgdiet, DOC, chl $a$</td>
<td>0.257</td>
<td>0.401</td>
<td>1.137</td>
</tr>
<tr>
<td></td>
<td>MeHgdiet</td>
<td>3.120</td>
<td>0.096</td>
<td>4.759</td>
</tr>
<tr>
<td></td>
<td>MeHgdiet, chl $a$</td>
<td>5.796</td>
<td>0.025</td>
<td>18.136</td>
</tr>
<tr>
<td></td>
<td>TL(predators/grazers)</td>
<td>7.107</td>
<td>0.013</td>
<td>34.932</td>
</tr>
<tr>
<td></td>
<td>MeHgdiet, DOC, TP, chl $a$</td>
<td>8.988</td>
<td>0.005</td>
<td>89.483</td>
</tr>
<tr>
<td></td>
<td>MeHgdiet, TL(predators/grazers)</td>
<td>11.274</td>
<td>0.002</td>
<td>280.630</td>
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<tr>
<td></td>
<td>MeHgdiet, TL(predators/grazers), DOC</td>
<td>11.876</td>
<td>0.001</td>
<td>379.108</td>
</tr>
<tr>
<td></td>
<td>Intercept only</td>
<td>12.995</td>
<td>0.001</td>
<td>663.349</td>
</tr>
</tbody>
</table>

Table 4. Partitioned-$R^2$ based relative variable importance (RVI) metrics $I_{global}$ and $I_{weighted}$ for model sets A (BMF(grazers/periphyton)) and B (BMF(predators/grazers)) showing the decomposition of variance among explanatory variables dietary methylmercury (MeHgdiet), trophic level of consumers relative to their prey (TL(consumer/prey)), dissolved organic C (DOC), total P (TP), and benthic chlorophyll $a$ (chl $a$).

<table>
<thead>
<tr>
<th>RVI Metric</th>
<th>MeHgdiet</th>
<th>TL(consumer/prey)</th>
<th>DOC</th>
<th>TP</th>
<th>Chl $a$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model set A</td>
<td>$I_{global}$</td>
<td>0.041</td>
<td>0.223</td>
<td>0.180</td>
<td>0.037</td>
</tr>
<tr>
<td></td>
<td>$I_{weighted}$</td>
<td>0.012</td>
<td>0.198</td>
<td>0.122</td>
<td>0.001</td>
</tr>
<tr>
<td>Model set B</td>
<td>$I_{global}$</td>
<td>0.397</td>
<td>0.251</td>
<td>0.148</td>
<td>0.222</td>
</tr>
<tr>
<td></td>
<td>$I_{weighted}$</td>
<td>0.744</td>
<td>0.007</td>
<td>0.069</td>
<td>&lt; 0.001</td>
</tr>
</tbody>
</table>

and Hairston 1993). The implication of this weak size structuring is that larger animals may not necessarily have the highest Hg concentrations nor occupy the highest TLs (Watanabe et al. 2008). At the comprehensive foodweb sites in our study, Tipulidae have low MeHg concentrations, low TLs, and the largest body sizes, whereas smaller invertebrates, particularly grazing mayflies, have the highest MeHg levels suggesting that other factors drive accumulation of MeHg in these organisms. In a study by Mason et al. (2000), smaller invertebrates had higher concentrations of As and Se, probably because their high surface area:volume ratios increased absorptive surface area relative to that of larger organisms, but findings for Hg were inconsistent. The relative contributions of dermal absorption and dietary uptake are less well understood for small invertebrates than fishes (Hall et al. 1997). Small-bodied omnivores can have higher-than-expected TLs. Invertebrates can exhibit both obligate and facultative functional feeding behavior and may undergo ontogenetic shifts in feeding habits (Cummins and Klug 1979). Hepta-
geniid and Ephemerellid mayflies are classified as herbivorous scrapers (Merritt and Cummins 1996) but can be facultative collectors. Baetids also can be facultative feeders. They have the potential to collect particles “of animal origin” or bacteria and, therefore, may occupy a higher trophic position than expected as a consequence of omnivory (Anderson and Cabana 2007). The potential for omnivory increases the utility of stable isotopes in bioaccumulation studies, particularly when establishing the relative difference in TL from prey to consumer (TLconsumer/prey). For example,
in our study, some grazing mayflies had higher $\delta^{15}N$ than presumed predators resulting in $\text{TL}_{\text{predators/ grazers}}$ that were $<1$ (data not shown), and ephemeralids occupied the same TL as predators at 2 upstream beaver sites (data not shown). Altered size–TL–Hg relationships have implications for trophic transfer because less Hg than expected will transfer up the food chain if fish tend to eat larger invertebrates and larger invertebrates have lower Hg.

Trophic transfer is more efficient at low than high environmental MeHg concentrations, but our results suggest that MeHg in top predators is governed more by supply of MeHg at the base of the food web than by trophic transfer and trophic efficiency (Chasar et al. 2009). Hg enters ecosystems largely as Hg(II), whereas MeHg production and subsequent uptake into algae occurs within the system and is highly variable (Driscoll et al. 2013). Concentrations of MeHg in predatory fishes can remain elevated for years following impoundment and flooding of large reservoirs because of release of Hg from soils and vegetation (St Louis et al. 2004, Hall et al. 2005). In addition, atmospheric deposition of Hg is strongly linked to Hg concentrations in fish (Harris et al. 2007). Anthropogenic emissions contribute $\frac{1}{3}$ of all atmospheric deposition (Hammerschmidt and Fitzgerald 2006) and affect fish populations that are far from any direct emissions. Decreases in Hg emissions are predicted to lead to rapid declines in fish concentrations (Harris et al. 2007). These examples, taken together with our findings, further highlight the importance of processes at the base of the food web in driving uptake and trophic transfer of Hg. The relative importance of supply vs uptake and transfer in determining Hg in organisms at higher TLs is important because inputs are more readily managed than transfers through the food web.

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