Multiple species drive flexible lake food webs with warming

Timothy J. Bartley, Tyler D. Tunney, Nigel P. Lester, Brian J. Shuter, Robert H. Hanner, and Kevin S. McCann

1 Department of Biology, University of Toronto Mississauga, 3359 Mississauga Road, Mississauga, Ontario, Canada, L5L 1C6
2 Department of Integrative Biology, University of Guelph, 50 Stone Road East, Guelph, Ontario, Canada N1G 2W1
3 Gulf Fisheries Centre, Fisheries and Oceans Canada, Moncton, New Brunswick E1C 9B6, Canada
4 Center for Limnology, University of Wisconsin–Madison, Madison, WI, USA 53706
5 Science and Research Branch, Ontario Ministry of Natural Resources, Peterborough, Ontario, Canada K9J 7B8
6 Department of Ecology and Evolution, University of Toronto, Toronto, Ontario, Canada M5S 3G5
7 Biodiversity Institute of Ontario, University of Guelph, 50 Stone Road East, Guelph, Ontario, Canada N1G 2W1

*correspondence to T.J.B.

e-mail: timothy.bartley@gmail.com
phone: 1 (519) 546-4652
Abstract
Climate change is rewiring the food webs that determine the fate of diverse ecosystems. Mobile generalist consumers are responding to climate change by rapidly shifting their behaviour and foraging, driving food webs to flex. Although these responsive generalists form a key stabilizing module in food web structure, the extent to which they are present throughout whole food webs is largely unknown. Here, we show that multiple species comprising key trophic roles drive flexible lake food webs with warming. By examining lakes that span a 7°C air temperature gradient, we found significant reductions in nearshore derived carbon and nearshore habitat use with increased temperature in three of four fish species. We also found evidence that the response of lake trout to increased temperatures may reduce their biomass and cascade to release their preferred prey, the pelagic forage fish cisco. Our results suggest that climate warming will shift lake food webs toward increased reliance on offshore habitats and resources. We argue that species across trophic levels broadly couple lake macrohabitats, suggesting that potentially stabilizing responsive consumers are present throughout food webs. However, climate change appears to limit their ability to responsively forage, critically undermining a repeated stabilizing mechanism in food webs.

Keywords: behavioral response, boreal shield, climate change, fish ecology, generalist module, habitat coupling, responsive foraging, mobile consumer, stable isotopes, trophic level


**Introduction**

Ongoing rapid climate change is altering species interactions and reorganizing the food webs that determine the fate of diverse ecosystems (Barton *et al.* 2009; Tunney *et al.* 2014; Kortsch *et al.* 2015). As environmental conditions change, food web structure changes—or ‘flexes’—with it (McMeans *et al.* 2016). Identifying these changes in food web structure is central to understanding the consequences of climate change on the major energy pathways of carbon and nutrient flow in ecosystems but is challenging given that food web structure is inherently complex (Polis 1991). Importantly, climate change is expected to ‘rewire’ food webs (Bartley *et al.* n.d.) by adding or removing whole trophic links (e.g., Kortsch *et al.* 2015) and by modifying the strengths of existing interactions (e.g., Barton & Schmitz 2009). These changes in the flow of energy among species are likely to have myriad consequences in ecosystems, from species to whole ecosystem function (Křivan & Schmitz 2003; Kondoh 2010; Guzzo *et al.* 2017). If so, future warming may alter food webs in ways that lead to ecosystems that differ substantially from past and present conditions (Kortsch *et al.* 2012). Despite the potential for these drastic changes, our understanding of food webs responses to climate change and the consequences of these response remain limited.

Food webs are flexible because of the foraging responses of mobile generalist consumers that can rapidly shift their foraging behaviour in response to environmental change. Many generalist consumers move across the landscape to forage in various spatially-distinct macrohabitats (Rooney *et al.* 2008) and can feed omnivorously across trophic levels (Thompson *et al.* 2007).

As a result, these generalist consumers can respond in two key ways: by changing the degree to which they couple different macrohabitats (i.e., ‘horizontal’ shifts between energy pathways, Schindler & Scheuerell 2002; Vander Zanden & Vadeboncoeur 2002; Rooney *et al.* 2008) and
by shifting their degree of omnivory (i.e., ‘vertical’ foraging shifts along an energy pathway, Barton & Schmitz 2009; Sentis et al. 2014; Ruiz-Cooley et al. 2017). Generalists that exhibit both horizontal and vertical foraging flexibility form the generalist module (sensu McMeans et al. 2016), a widely documented patterning of food web interactions (Polis et al. 1997; Nakano & Murakami 2001; Rooney et al. 2008; Eloranta et al. 2015) that can expand and contract food webs in response to changing environmental conditions (Tunney et al. 2012). This generalist module may be a powerful stabilizing structure, allowing mobile generalists to respond to changing densities in different habitats in a manner that buffers population dynamics (McCann & Rooney 2009; McMeans et al. 2016). Despite this potential, few studies have empirically documented the presence of fundamental module throughout food webs or how human impacts like climate change may interrupt this module.

One well-documented example of a responsive generalist consumer that drives food web flexibility is lake trout (Salvelinus namaycush), a common generalist top predator in lakes of boreal North America. Lake trout forage on fishes and invertebrates in both of the two thermally distinct lake macrohabitats: the nearshore (i.e., littoral) habitat and the offshore (i.e., pelagic) habitat (Vander Zanden & Vadeboncoeur 2002). Because lake trout is cold-adapted and thermally sensitive, the foraging habits of lake trout are mediated by thermal accessibility of the relatively warm nearshore habitat (Plumb & Blanchfield 2009; Guzzo et al. 2017). Lake trout show reduced nearshore habitat use with increasing nearshore temperature (Tunney et al. 2014; Guzzo et al. 2017). When lake trout avoid entering the physiologically taxing nearshore habitat, they forage less on nearshore resources and more on fish (Tunney et al. 2012, 2014). This reduced access to abundant nearshore prey may undermine lake trout’s ability to responsively forage to resource densities in different macrohabitats (Tunney et al. 2014; Guzzo et al. 2017).
Importantly, this phenomenon may not be limited to lake trout; many fishes are generally highly mobile and flexible foragers (Dill 1983), and boreal lakes in North America contain numerous cold-adapted fish species that may exhibit similar behavioural and foraging responses due to thermal limitations (Vander Zanden & Vadeboncoeur 2002; Hasnain et al. 2013). In contrast, the species that prefer warmer waters may not be thermally limited and thus may not exhibit behavioural or foraging responses to warming. However, the roles that species other than lake trout play in determining how lake food webs flex are largely unexplored. To understand how whole food webs are being altered by climate change requires that we examine how many species across trophic levels and with various thermal preferences respond to warming.

Here, we use a natural climate gradient of approximately 7°C to examine how multiple species behavioural and foraging responses drive flexes in food web structure. We investigate the habitat use and diet of four widespread and abundant species of boreal shield fishes that represent different key roles and trophic levels in the food web: the cold-adapted generalist top predator lake trout, the cold-adapted pelagic planktivorous cisco (Coregonus artedi), the cool-adapted piscivorous top predator walleye (Sander vitreus), and the cool-adapted mesopredator yellow perch (Perca flavescens) (Coker et al. 2001; Hasnain et al. 2013). We use stable-isotope-based food-web indices of nearshore feeding and trophic position along with catch-per-unit-effort based metrics of behaviour to show that several species across trophic levels and from multiple thermal guilds respond predictably to changes in temperature. We also use catch-per-unit-effort data to show that lake trout show reduced biomass index under increased temperatures, consistent with a reduction of the nearshore resource availability. Our results suggestion that multiple species comprising key trophic roles drive flexible lake food webs with warming. We argue that by studying the foraging and behavioral responses of this set of key
players, we can predict how food webs will flex with climate change. We end by discussing how these changes suggest climate change will impact the function and stability of lake ecosystems, and how many generalist consumers may be a fundamental feature of food web architecture.

Methods

Lake Selection

The Canadian boreal shield includes hundreds of thousands of lakes that span various natural environmental gradients, including climate (Gunn et al. 2004; Keller 2007). We used data for 66 lakes in the province of Ontario, all of which have been used previously to study food web structure through stable isotope analysis (Dolson et al. 2009; Tunney et al. 2014, 2018). We use lakes that have both nearshore and offshore intermediate consumers to ensure that changes in food web structure or predator behaviour are not driven by differences in the presence or absence of whole trophic groups. For 59 of our 66 lakes, we used catch-per-unit-effort (CUE) data from the Ontario Ministry of Natural Resources and Forestry (OMNRF) Broad-scale Fisheries Monitoring (BSM) Program (Sandstrom et al. 2013). In summary, for each lake, standardized fish community surveys took place one time between May and September from 2008 to 2012 using overnight sets of two types of multipaneled mesh gillnets: North American standard multipaneled gill nets (with 8 mesh sizes varying from 38 to 127 mm) and small mesh gillnets (with 5 mesh sizes from 13 to 38 mm). Netting occurred randomly throughout the lake and depending on the maximum depth of each lake, in each of up to 8 depth strata: 1-3m, 3-6m, 6-12m, 12-20m, 20-35m, 35-50m, 50-75m, and >75m. Complete details can be found in Sandstrom et al. 2013. Catch data were not available for the 7 lakes from Dolson et al. (2009).
Lake Attributes

We used two measures of temperature for our analysis: average recent air temperature in °C growing degree days > 5 °C (Table S1). Our primary temperature variable was average recent air temperature, which was calculated as the average hourly air temperature in °C for the 30 days prior to field sampling because it corresponds to the time period reflected by the isotopic signature of fish muscle tissue (Peterson & Fry 1987). This temperature variable was derived from FetchClimate (Grechka et al. 2016) using the latitude and longitude of each lake. In addition to the average recent air temperature, we corroborated our results with and growing degree days above 5°C from 1981 to 2010 (provided by the Ontario Ministry of Natural Resources and Forestry), which measures the accumulation of heat over time. We also account for several physical and chemical factors that have been previously identified as important drivers of habitat coupling or trophic position in boreal shield lakes (Table 1, all provided by the Ontario Ministry of Natural Resources, Dolson et al. 2009; Tunney et al. 2014, 2018): lake surface area in hectares, mean lake depth in metres, Secchi depth in metres, and total phosphorus in μg•L⁻¹, and lake shape (calculated the shoreline development index):

\[ SDI = \frac{Shoreline Distance}{2\sqrt{\pi \times Surface Area}} \]

Species Selection

We used four species that are typical of their respective trophic groups. Stable isotope data for these species was available for 40 or more lakes and the trophic level, thermal classifications, and habitat preferences for these species are known (Coker et al. 2001; Hasnain et al. 2013). Lake trout (*Salvelinus namaycush*) is the most common top predator of offshore habitats in boreal shield lakes, and much research on boreal shield lake food webs has focused on
this coldwater species (Dolson et al. 2009; Plumb & Blanchfield 2009; Tunney et al. 2014; Guzzo et al. 2017). Cisco (or lake herring, Coregonus artedi) is one of the most common cold-adapted planktivores and is a common prey item for lake trout and walleye. Walleye (Sander vitreus) is a common cool-adapted piscivore and popular sport fish that is present in lakes across the boreal shield. Yellow perch (Perca flavescens) is a widespread and abundant cool-adapted mesopredator that is consumed by a wide variety of predatory fishes throughout its range, including both lake trout and walleye. These four species comprise a large portion of the average catch in the 59 lakes that we use here from the OMNRF’s BsM surveys (see Supplementary Information). The number of individual for each species sampled for stable isotope analysis in each lake varied from 2 to 21 (mean 13.5) for lake trout, 3 to 32 (mean 17.0) for walleye, 1 to 20 (mean 10.8) for cisco, and 1 to 18 (mean 9.1) for yellow perch. For both lake trout and walleye, only individuals greater than 250 mm were used for stable isotope analysis because these species are known to show ontogenic diet shifts (Mittelbach & Persson 1998; Sherwood et al. 2002; Galarowicz et al. 2006).

Food Web Metric Calculations using Stable Isotopes

As in many previous studies, we used stable isotopic signatures from our four fish species and baseline invertebrates to calculate both the nearshore carbon index (based on the proportion nearshore carbon, e.g., Tunney et al. 2014) and trophic position (Vander Zanden et al. 1999b, a; Post 2002). Collection and processing methods for stable isotope data can be found in Dolson et al. 2009; Tunney et al. 2014, 2018). We used baseline invertebrates from the nearshore and offshore zones to account for variability in isotopic signatures across lakes. We incorporated data for multiple trophic groups into both our nearshore and offshore baseline isotopic signatures to
reduce the number of estimated baseline values required for our analysis and to increase the
sample size for our baseline isotopic signatures. We corrected all $\delta^{13}C$ signatures using C:N
ratios as

$$\delta^{13}C_{corr} = \delta^{13}C_{raw} + (-3.32 + 0.99 \times CN)$$

where $\delta^{13}C_{corr}$ is the corrected $\delta^{13}C$ signature, $\delta^{13}C_{raw}$ is the raw $\delta^{13}C$ signature, and CN is the
C:N ratio of that tissue sample (Post et al. 2007). For lakes that were missing either nearshore or
offshore baseline isotopic signatures, we estimated the $\delta^{13}C$ and $\delta^{15}N$ signatures of the missing
baseline using the available baseline and simple linear regression between the baselines across
lakes (see Supplementary Information).

We used two source mixing models to estimate the nearshore carbon index and the
trophic position of each species in each lake based on their relative isotopic signatures (Post
2002). We calculated the nearshore carbon index for each fish species as

$$NCI_{fish} = \frac{\delta^{13}C_{fish} - \delta^{13}C_{osb}}{\delta^{13}C_{nsb} - \delta^{13}C_{osb}}$$

where $NCI_{fish}$ is the nearshore carbon index in the diet of a fish species, $\delta^{13}C_{fish}$ is the
average $\delta^{13}C$ signature for that fish species, $\delta^{13}C_{osb}$ is the average or estimated $\delta^{13}C$ signature
for all offshore baselines (i.e., mussels and/or zooplankton), and $\delta^{13}C_{nsb}$ is the average or
estimated $\delta^{13}C$ signature for all nearshore baselines (i.e., snails and/or aquatic insect larvae). The
nearshore carbon index is similar to the proportion nearshore carbon used by others (e.g., Tunney
et al. 2014) but is not constrained between 0 and 1. This approach minimizes data transformation
requirements of the stable isotopic signatures and increases the number of lakes in our analyses
while still allowing us to qualitatively understand changes in the relative contribution of nearshore resources to fish diets.

Based on the nearshore feeding index, we estimated the trophic position of each species as

$$TP_{fish} = 2 + \left( \delta^{15}N_{fish} - \left( \delta^{15}N_{nsb} \times NCI_{fish} + \delta^{15}N_{osb} \times (1 - NCI_{fish}) \right) \right) \div 3.4$$

where $TP_{fish}$ is the trophic position of a fish species, $NFI_{fish}$ is the nearshore carbon index for that fish species, $\delta^{15}N_{fish}$ is the average $\delta^{15}N$ signature for that fish species, $\delta^{15}N_{nsb}$ is the average or estimated $\delta^{15}N$ signature for all nearshore baselines (i.e., snails and aquatic insect larvae), $\delta^{15}N_{osb}$ is the average or estimated $\delta^{15}N$ signature for all offshore baselines (i.e., mussels and zooplankton), 3.4 is the assumed increase in $\delta^{15}N$ due to fractionation (Post 2002), for each trophic level, and 2 is the assumed trophic position of the baseline invertebrates.

**Behaviour Metrics and Biomass Index Using Catch-per-unit-effort**

We used catch-per-unit-effort data for each depth stratum (Sandstrom et al. 2013) to calculate a weighted average mean depth of capture for each species as

$$SMD = \frac{\sum_{i=1}^{4} d_i \times CUE_i \times p_i}{\sum_{i=1}^{4} CUE_i}$$

where SMD is the mean depth of capture of a fish species, $CUE_i$ is the CUE of that species for depth stratum $i$, $p_i$ is the proportion of the lake in depth stratum $i$, and $d_i$ is the depth of the center of the depth range for stratum $i$ (2m for stratum 1, 4.5m for stratum 2, 9m for stratum 3,
and 16m for stratum 4). We only used depth strata 1 through 4 to calculate mean depth of capture because these are the only depth strata that are sampled with both small-mesh and large-mesh gear (Sandstrom et al. 2013). The CUE data that we use here is the number of fish per 100 metre nights. For CUE biomass index, we used the kg of fish per gang per 100 metre nights, calculated by summing the area-weighted CUE (number of fish per 100 metre nights) for each depth stratum and then multiplying by the mean weight of that fish species.

Statistical Analyses

To test for the effect of temperature on each response variable (nearshore carbon index, trophic position, mean depth of capture, nearshore/offshore presence, and the log-transformed CUE biomass index of each species), we used a two-step approach. Firstly, to account for morphometric or chemical factors known or suspected to influence the foraging and behaviour of these fish species, we first constructed multiple regression models for each response variable and lake size in hectares, lake shape (SDI), Secchi depth in m, mean lake depth in m, and total phosphorous in μg·L⁻¹. We then compared all subsets of each full multivariate regression model using AIC. We selected the best model as the model with the smallest number of explanatory variables from the set of models within 2 AIC points of the lowest AIC value (Aho et al. 2014, see Supplementary Information for best models). Secondly, to test the effect of each temperature variable, we took the best model for each response variable and ran a multiple regression that included the temperature variable (Figures 2, 3, and 4, Tables 3 and 4). To meet statistical requirements, we used log₁₀ transformed lake surface area and natural log transformed mean lake depth and total phosphorous. We removed all data points with a Cook’s distance greater than 1 from their respective regression models, and we used variance inflation factor to check for
inflated coefficient estimates in multiple linear regressions. All regression analyses were performed in the R statistical language (v3.2.3), and model selection was conducted using the R package “glmulti” (Calcagno 2013).

Results

Food Web Metric Calculations using Stable Isotopes

We found strong evidence that lake food web structure varied across an approximately 7° C climate gradient. Three of the four species showed evidence of changes in nearshore feeding with increased temperature. Both top predators (lake trout and walleye) showed a significant decrease in the nearshore carbon index with increasing average recent air temperature, and the cold-adapted cisco showed a similar marginally significant decrease (Figure 2a, c, and e, Table 3). The cool-adapted yellow perch showed no significant relationship between average recent air temperature and the nearshore carbon index (Figure 2g, Table 3). Lake trout showed a significant increase in trophic position with increased average recent air temperature (Figure 2b, Table 3), but cisco, walleye and yellow perch showed no significant relationship between trophic position with increasing average recent air temperature (Figure 2d, Table 3) (see Supplementary Information and Table S3).

Behaviour Metrics and Biomass Index Using Catch-per-unit-effort

In agreement with changes in nearshore carbon index, the same three species (lake trout, walleye, and cisco) showed strong evidence of reduced nearshore habitat use. All three species showed a significant increase in mean depth of capture with increasing average recent air
temperature (Figure 3a, c, and e, Table 3). Consistent with these results, lake trout and cisco showed a significant decrease in probability of nearshore presence and walleye showed a marginally significant increase in probability of offshore presence with increasing average recent air temperature (see Supplementary Information). In contrast, yellow perch showed no relationship between either mean depth of capture or probability of offshore presence and average recent air temperature (Figure 3g and inset, Table 3).

We found evidence of a reduction in biomass with increasing temperature for one species, lake trout, which showed a significant decrease in CUE biomass index with increasing average recent air temperature (Figure 3b, Table 3). In contrast, cisco, walleye, and yellow perch showed no significant relationship between average recent air temperature CUE biomass index (Figure 3d, f and h, Table 3).

**Discussion**

Understanding how climate change is altering whole food webs is a challenging but critical endeavor for understanding the responses of whole ecosystems. Previous research showed that lake trout (*Salvelinus namaycush*), a generalist top predator in lake ecosystems, responds to increased temperature by reducing its nearshore habitat and resource use, and its degree of omnivory (Tunney et al. 2014; Guzzo et al. 2017), thus forming a flexible ‘generalist module’ (McMeans et al. 2016). Here, we extend that research to examine the impacts of climate warming on four key species that represent different key roles and trophic levels in boreal lake food webs: lake trout, walleye (*Sander vitreus*), cisco (*Coregonus artedi*), and yellow perch (*Perca flavescens*). Our stable-isotope-based food-web metrics and our catch-per-unit-effort-based behavioural metrics show a consistent response of species across multiple trophic levels to
warming to nearshore-derived carbon and offshore habitat use. In fact, temperature was the most significant variable for predicting food web structure in this study (see Supplementary Information). By harnessing the variation in food web structure across natural temperature gradients, we have detected consistent food web responses to warming in boreal shield lake ecosystems.

We found strong evidence of foraging and behavioural responses to increased temperature in three of the four key food web members that we examined. As expected, our cold-adapted species, the predatory lake trout and the planktivorous cisco, showed decreased nearshore feeding and decreased near-shore habitat use with increasing temperatures. Our results are consistent with the thermally accessibility hypothesis that has been previously shown for lake trout (Tunney et al. 2014; Guzzo et al. 2017). Because offshore fish communities are largely comprised of species that prefer cold water (Coker et al. 2001; Hasnain et al. 2013), these thermal constraints likely operate on whole offshore fish communities, causing them to respond uniformly to warming. We found that the cool-adapted piscivorous walleye showed behavioural and foraging responses through decreased coupling and decreased nearshore habitat use with higher temperature, but in contrast the cool-adapted mesopredator yellow perch showed no such response. Since walleye and yellow perch have similar thermal tolerances (Hasnain et al. 2013), the reason for the different responses of these species is unclear. Despite the lack of response in yellow perch, the foraging responses of these fish species strongly indicate that boreal shield lake food webs will collectively flex towards offshore resource and habitat use in response to warming.

We found evidence that a reduction in nearshore coupling with warming may impact the biomass of lake trout. The biomass of lake trout decreased significantly with increased
temperature, but none of the other species showed this pattern (Figure 3). Such reductions in biomass with increasing temperature may be a consequence of a reduction in overall energy availability due to decreased nearshore coupling. Mobile top predators that garner energy from multiple spatially distinct macrohabitats are expected to have elevated biomass as a result of a higher overall resource availability, producing relatively top-heavy food webs (i.e., non-Eltonian, Mccauley et al. 2018; Woodson et al. 2018). Therefore, reduced access to the nearshore energy pathway in warm lakes could therefore reduce overall energy availability for lake trout and so lower lake trout biomass and population density (Guzzo et al. 2017; Tunney et al. 2018). A reduction in lake trout biomass across could subsequently reduce the top-down control of lake trout on cisco, their preferred prey. Reduced top-down control by lake trout in warmer lakes in conjunction with a bottom-up reduction in their biomass from reduced access to nearshore resources may result in no discernable trend in standing biomass of cisco with temperature, consistent with our results here. Changes in trophic biomass structure and concomitant changes in top-down control are one possible consequence of the flexes in lake food webs towards the offshore with warming. If climate change broadly decouples predators from multiple resource pools, changes in biomass structure may be a widespread signatures of climate change, possibly altering top-down effects within ecosystems (McCann et al. 2005; McCauley et al. 2018).

Our results demonstrate that many mobile generalist consumers at trophic levels and throughout compartments drive flexibility throughout food webs in boreal shield lakes. Our work builds on previous studies that have started to expose the role that single generalist top predators species play in determining food web responses to climate change in many ecosystems (Barton & Schmitz 2009; Woodward et al. 2010; Eloranta et al. 2016). The presence of flexible foragers at various trophic levels is consistent with both recent ecological theory that predicts large,
Mobile generalists ought to rapidly respond to environmental change (Rooney & McCann 2012), and is consistent with empirical evidence of flexible foraging from various ecosystems (Schmitz et al. 1997; Held et al. 2002; Vander Zanden & Vadeboncoeur 2002; Shochat et al. 2004; Rooney et al. 2008; Taipale et al. 2008; Gonçalves et al. 2012; Edmunds et al. 2016). The presence of responsive foraging throughout diverse food webs rather than just at the top of food chains would make the generalist module the type of remarkably scale-invariant feature of food web architecture that has long interested ecologists (Sugihara et al. 1989).

The presence of responsive consumers across trophic levels and throughout food web compartments has strong implications for food web stability. Responsive foraging by generalists can be potently stabilizing in food webs (Post et al. 2000; Rooney et al. 2006; Rip et al. 2010; McMeans et al. 2016). Generalist consumers can average variability in their resources (akin to the portfolio effect of Tilman et al. (1998)) by rapidly responding to resource variability in space and time by shifting away from low-density resources and towards high density resources, impeding that variability from emanating throughout food webs (Valdovinos et al. 2010). This switching ability also increases trophic redundancy, which can prevent secondary extinctions (Borrvall et al. 2000; Sanders et al. 2018). If a broad suite of organisms can make rapid, smart foraging responses to environmental change, they may accentuate the stabilizing effect of a single generalist module. The presence of many responsive consumers increases stability in complex food web models (Kondoh 2010; Valdovinos et al. 2010). This reveals a potential powerful, repeated stabilizing mechanism throughout food webs that Levin (1998) argues is fundamental to persistence of ecosystems. However, when generalist consumers decouple macrohabitats to avoid entering physiologically taxing habitats, their ability to respond to consumer densities is limited, undermining their stabilizing potential (Schmitz & Barton 2014;
Thus, the decoupling of habitats with warming may be widespread across ecosystems, forcing the food webs of many ecosystems to flex with climate change in ways that alter ecosystem stability worldwide.

Our work here adds a key piece to the mounting evidence that generalist consumers likely to play a vital role in determining how climate change rewires food webs (Bartley et al. n.d.). Generalist consumers positioned throughout food webs may ultimately reorganize whole ecosystems in response to climate change (Blanchard 2015; McMeans et al. 2016). Determining both the prevalence and positions of these responsive generalist consumers and the factors that constrain these species responses are critical tasks for predicting the myriad ways that food webs will adapt with a changing climate. Importantly, our results suggest that these species throughout food webs contribute to adaptive capacity—the ability of an ecosystem to change in response to changing conditions—a key part of the resilience of ecosystems in the face of environmental variation (McMeans et al. 2016). Although we have showed that lake food webs flex at many axes, it is unclear when this adaptive capacity will be exceeded and entire ecosystems collapse. This uncertainty highlights the clear need to forecast major changes in food web structure. Monitoring the behaviour and foraging of responsive consumers is a promising way to track changes in food web structure and give prescient warnings of future drastic changes in ecosystem dynamics (Bartley et al. n.d.; Velarde et al. 2013). This makes responsive generalist consumers critical to understanding of how ecosystems will fare through climate change.
Acknowledgements

This work was supported by the National Science and Engineering Research Council of Canada through a Discovery Grant to K.S.M., a Strategic Grant to K.S.M., B.J.S. and N.P.L., and a Canada Graduate Scholarship to T.J.B. We thank K. Armstrong, J. Wright and many other biologists and technical staff at the Ontario Ministry of Natural Resources and Forestry and the Broad-scale Fisheries Monitoring Program. We also thank M. Berger, I. Farahbakhsh, B. Graham, B. Heavenrich for assistance with gathering and organizing climate data. D. Schindler, M. Granados, M. Guzzo, and B. Robinson provided helpful comments on an early draft of this manuscript.

Author Contributions

T.J.B. and K.S.M. conceived and designed this study. T.J.B. and T.D.T. collected field data and conducted lab work. T.J.B. analyzed data and wrote the initial draft of the manuscript. T.J.B., T.D.T., N.P.L., B.J.S., R.H.H., and K.S.M. all contributed substantially to writing and revisions.
References

Aho, K., Derryberry, D. & Peterson, T. (2014). Model selection for ecologists: The worldviews of AIC and BIC. *Ecology.*

Bartley, T.J., McCann, K.S., Bieg, C., Cazelles, K., Granados, M., Guzzo, M.M., et al. (n.d.). Food Web Rewiring in a Changing World. *Nat. Ecol. Evol.*

Barton, B.T., Beckerman, A.P. & Schmitz, O.J. (2009). Climate warming strengthens indirect interactions in an old-field food web. *Ecology,* 90, 2346–2351.

Barton, B.T. & Schmitz, O.J. (2009). Experimental warming transforms multiple predator effects in a grassland food web. *Ecol. Lett.,* 12, 1317–1325.

Blanchard, J.L.A. (2015). A rewired food web. *Nature,* 527, 7–8.

Borrvall, C., Ebenman, B. & Jonsson, T. (2000). Biodiversity lessens the risk of cascading extinctions in model food webs. *Ecol. Lett.,* 3, 131–136.

Calcagno, V. (2013). glmulti: Model selection and multimodel inference made easy.

Coker, G., Portt, C. & Minns, C.K. (2001). *Morphological and ecological characteristics of Canadian freshwater fishes.* Burlington, Ontario.

Dill, L.M. (1983). Adaptive Flexibility in the Foraging Behavior of Fishes. *Can. J. Fish. Aquat. Sci.,* 40, 398–408.

Dolson, R., McCann, K., Rooney, N. & Ridgway, M. (2009). Lake morphometry predicts the degree of habitat coupling by a mobile predator. *Oikos,* 118, 1230–1238.

Edmunds, N.B., Laberge, F. & McCann, K.S. (2016). A role for brain size and cognition in food webs. *Ecol. Lett.,* 19, 948–955.

Eloranta, A.P., Helland, I.P., Sandlund, O.T., Hesthagen, T., Ugedal, O. & Finstad, A.G. (2016). Community structure influences species’ abundance along environmental gradients. *J. Anim. Ecol.,* 85, 273–282.

Eloranta, A.P., Kahilainen, K.K., Amundsen, P.A., Knudsen, R., Harrod, C. & Jones, R.I. (2015). Lake size and fish diversity determine resource use and trophic position of a top predator in high-latitude lakes. *Ecol. Evol.,* 5, 1664–1675.

Galarowicz, T.L., Adams, J.A. & Wahl, D.H. (2006). The influence of prey availability on ontogenetic diet shifts of a juvenile piscivore. *Can. J. Fish. Aquat. Sci.,* 63, 1722–1733.

Gonçalves, A.M.M., Azeiteiro, U.M., Pardal, M.A. & De Troch, M. (2012). Fatty acid profiling reveals seasonal and spatial shifts in zooplankton diet in a temperate estuary. *Estuar. Coast. Shelf Sci.,* 109, 70–80.

Grechka, D.A., Berezin, S.B., Emmott, S., Lyutsarev, V., Smith, M.J. & Purves, D.W. (2016). Universal, easy access to geotemporal information: FetchClimate. *Ecography (Cop.)*
Gunn, J.M., Steedman, R.J. & Ryder, R.A. (2004). *Boreal shield watersheds: lake trout ecosystems in a changing environment*. CRC Press.

Guzzo, M.M., Blanchfield, P.J. & Rennie, M.D. (2017). Behavioral responses to annual temperature variation alter the dominant energy pathway, growth, and condition of a cold-water predator. *Proc. Natl. Acad. Sci.*, 201702584.

Hasnain, S.S., Shuter, B.J. & Minns, C.K. (2013). Phylogeny influences the relationships linking key ecological thermal metrics for North American freshwater fish species. *Can. J.* ..., 972, 964–972.

Held, S., Mendl, M., Devereux, C. & Byrne, R.W. (2002). Foraging pigs alter their behaviour in response to exploitation. *Anim. Behav.*, 64, 157–165.

Keller, W. (2007). Implications of climate warming for Boreal Shield lakes: a review and synthesis. *Environ. Rev.*, 15, 99–112.

Kondoh, M. (2010). Foraging Adaptation and the Relationship between Food-Web Complexity and Stability. *Science (80- ).*, 299, 1388–1391.

Kortsch, S., Primicerio, R., Beuchel, F., Renaud, P.E., Rodrigues, J., Lønne, O.J., et al. (2012). Climate-driven regime shifts in Arctic marine benthos. *Proc. Natl. Acad. Sci. U. S. A.*, 109, 14052–7.

Kortsch, S., Primicerio, R., Fossheim, M., Dolgov, A. V. & Aschan, M. (2015). Climate change alters the structure of arctic marine food webs due to poleward shifts of boreal generalists. *Proc. R. Soc. B Biol. Sci.*, 282, 20151546.

Krivan, V. & Schmitz, O.J. (2003). Adaptive foraging and flexible food web topology. *Evol. Ecol. Res.*, 5, 623–652.

Levin, S.A. (1998). Ecosystems and the biosphere as complex adaptive systems. *Ecosystems*, 1, 431–436.

McCann, K.S., Rasmussen, J.B. & Umbanhowar, J. (2005). The dynamics of spatially coupled food webs. *Ecol. Lett.*, 8, 513–523.

McCann, K.S. & Rooney, N. (2009). The more food webs change, the more they stay the same. *Philos. Trans. R. Soc. Lond. B. Biol. Sci.*, 364, 1789–1801.

McCaul, D.J., Gellner, G., Martinez, N.D., Williams, R.J., Sandin, S.A., Micheli, F., et al. (2018). On the prevalence and dynamics of inverted trophic pyramids and otherwise top-heavy communities. *Ecol. Lett.*, 21, 439–454.

McMeans, B.C., McCann, K.S., Tunney, T.D., Fisk, A.T., Muir, A.M., Lester, N.P., et al. (2016). The adaptive capacity of lake food webs: from individuals to ecosystems. *Ecol. Monogr.*, 86, 4–19.

Mittelbach, G.G. & Persson, L. (1998). The ontogeny of piscivory and its ecological consequences. *Can. J. Fish. Aquat. Sci.*, 55, 1454–1465.
Nakano, S. & Murakami, M. (2001). Reciprocal subsidies: Dynamic interdependence between terrestrial and aquatic food webs. *Proc. Natl. Acad. Sci. U. S. A.*, 98, 166–170.

Peterson, B.J. & Fry, B. (1987). Stable isotopes in ecosystem studies. *Annu. Rev. Ecol. Syst.*, 18, 293–320.

Plumb, J.M. & Blanchfield, P.J. (2009). Performance of temperature and dissolved oxygen criteria to predict habitat use by lake trout (Salvelinus namaycush). *Can. J. Fish. Aquat. Sci.*, 66, 2011–2023.

Polis, G.A. (1991). Complex trophic interactions in deserts: an empirical critique of food-web theory. *Am. Nat.*

Polis, G.A., Anderson, W.B., Anderson, R. & Holt, R.D. (1997). Toward an Integration of Landscape and Food Web Ecology: The Dynamics of Spatially Subsidized Food Webs. *Annu. Rev. Ecol. Syst.*, 28, 289–316.

Post, D.M. (2002). Using stable isotopes to estimate trophic position: models, methos, and assumptions. *Ecology*, 83, 703–718.

Post, D.M., Conners, M.E. & Goldberg, D.S. (2000). Prey preference by a top preator and the stability of linked food chains. *Ecology*, 81, 8–14.

Post, D.M., Layman, C.A., Arrington, D.A., Takimoto, G., Quattrochi, J. & Montaña, C.G. (2007). Getting to the fat of the matter: Models, methods and assumptions for dealing with lipids in stable isotope analyses. *Oecologia*, 152, 179–189.

Rip, J.M.K., McCann, K.S., Lynn, D.H. & Fawcett, S. (2010). An experimental test of a fundamental food web motif. *Proc. Biol. Sci.*, 277, 1743–1749.

Rooney, N., McCann, K., Gellner, G. & Moore, J.C. (2006). Structural asymmetry and the stability of diverse food webs. *Nature*, 442, 265–269.

Rooney, N. & McCann, K.S. (2012). Integrating food web diversity, structure and stability. *Trends Ecol. Evol.*, 27, 40–45.

Rooney, N., McCann, K.S. & Moore, J.C. (2008). A landscape theory for food web architecture. *Ecol. Lett.*, 11, 867–881.

Ruiz-Cooley, R.I., Gerrodette, T., Fiedler, P.C., Chivers, S.J., Danil, K. & Ballance, L.T. (2017). Temporal variation in pelagic food chain length in response to environmental change. *Sci. Adv.*, 3, e1701140.

Sanders, D., Thébault, E., Kehoe, R. & Frank van Veen, F.J. (2018). Trophic redundancy reduces vulnerability to extinction cascades. *Proc. Natl. Acad. Sci.*, 0, 201716825.

Sandstrom, S., Rawson, M. & Lester, N. (2013). *Manual of instructions for broad-scale fish community monitoring using North American (NA1) and Ontario small mesh (ON2) gillnets*.

Schindler, D.E. & Scheuerell, M.D. (2002). Habitat coupling in lake ecosystems. *Oikos*, 98, 177–
Schmitz, O.J. & Barton, B.T. (2014). Climate change effects on behavioral and physiological ecology of predator-prey interactions: Implications for conservation biological control. *Biol. Control*, 75, 87–96.

Schmitz, O.J., Beckerman, A.P. & O’Brien, K.M. (1997). Behaviorally mediated trophic cascades: Effects of predation risk on food web interactions. *Ecology*, 78, 1388–1399.

Sentis, A., Hemptinne, J.L. & Brodeur, J. (2014). Towards a mechanistic understanding of temperature and enrichment effects on species interaction strength, omnivory and food-web structure. *Ecol. Lett.*, 17, 785–793.

Sherwood, G.D., Pazzia, I., Moeser, A., Hontela, A. & Rasmussen, J.B. (2002). Shifting gears: enzymatic evidence for the energetic advantage of switching diet in wild-living fish. *Can. J. Fish. Aquat. Sci.*, 59, 229–241.

Shochat, E., Lerman, S.B., Katti, M. & Lewis, D.B. (2004). Linking optimal foraging behavior to bird community structure in an urban-desert landscape: field experiments with artificial food patches. *Am. Nat.*, 164, 232–243.

Sugihara, G., Schoenly, K. & Trombla, A. (1989). Scale invariance in food web properties. *Science (80-. ).*, 245, 48–52.

Taipale, S., Kankaala, P., Tiirila, M. & Jones, R.I. (2008). Whole-Lake Dissolved Inorganic 13C Additions Reveal Seasonal Shifts in Zooplankton Diet. *Ecology*, 89, 463–474.

Thompson, R.M., Hemberg, M., Starzomski, B.M., Shurin, J.B., Hompson, R.M.T., Emberg, M.H., et al. (2007). Trophic Levels and Trophic Tangles: the Prevalence of Omnivory in Real Food Webs. *Ecology*, 88, 612–617.

Tilman, D., Lehman, C.L., Bristow, C.E., Tilman, D., Lehman, C.L. & Bristow, C.E. (1998). Notes and Comments Diversity-Stability Relationships: Statistical Inevitability or Ecological Consequence? *Am. Nat.*, 151, 277–282.

Tunney, T., McCann, K., Jarvis, L., Lester, N. & Shuter, B. (2018). Blinded by the light? Nearshore energy pathway coupling and relative predator biomass increased with reduced water transparency across lakes. *Oecologia*.

Tunney, T.D., McCann, K.S., Lester, N.P. & Shuter, B.J. (2012). Food web expansion and contraction in response to changing environmental conditions. *Nat. Commun.*, 3, 1105.

Tunney, T.D., McCann, K.S., Lester, N.P. & Shuter, B.J. (2014). Effects of differential habitat warming on complex communities. *Proc. Natl. Acad. Sci. U. S. A.*, 111, 8077–82.

Valdovinos, F.S., Ramos-Jiliberto, R., Garay-Narváez, L., Urbani, P. & Dunne, J.A. (2010). Consequences of adaptive behaviour for the structure and dynamics of food webs. *Ecol. Lett.*, 13, 1546–1559.

Velarde, E., Ezcura, E. & Anderson, D.W. (2013). Seabird diets provide early warning of
sardine fishery declines in the Gulf of California. *Sci. Rep.*, 3, 1332.

Woodson, C.B., Schramski, J.R. & Joye, S.B. (2018). A unifying theory for top-heavy ecosystem structure in the ocean. *Nat. Commun.*, 9, 23.

Woodward, G., Perkins, D.M. & Brown, L.E. (2010). Climate change and freshwater ecosystems: impacts across multiple levels of organization. *Philos. Trans. R. Soc. London B Biol. Sci.*, 365, 2093–2106.

Vander Zanden, M.J., Casselman, J.M. & Rasmussen, J.B. (1999a). Stable isotope evidence for the food web consequences of species invasions in lakes. *Nature*, 401, 464–467.

Vander Zanden, M.J., Shuter, B.J., Lester, N. & Rasmussen, J.B. (1999b). Patterns of food chain length in lakes: A stable isotope study. *Am. Nat.*, 154, 406–416.

Vander Zanden, M.J. & Vadeboncoeur, Y. (2002). Fishes As Integrators of Benthic and Pelagic Food Webs in Lakes. *Ecology*, 83, 2152–2161.
Figure 1. Nearshore foraging index and the mean log(depth) of capture of lake trout (a and b), cisco (c and d), walleye (e and f), and yellow perch (g and h) in lakes across a gradient of average recent air temperature. A solid line indicates a p-value less than 0.05, and a dashed line indicates a p-value less than 0.1. Regression model details can be found in Table S3. Fish silhouettes were taken from PhyloPic (http://www.phylopic.org).
Figure 2. Trophic position of lake trout (a), walleye (b), cisco (c), and yellow perch (d) in lakes across a gradient of average recent air temperature. A solid line indicates a p-value less than 0.05, and a dashed line indicates a p-value less than 0.1. Regression model details can be found in Table S3. Fish silhouettes were taken from PhyloPic (http://www.phylopic.org).
Figure 3. The log-transformed CUE biomass index for lake trout (a), walleye (b), cisco (c), and yellow perch (d) in lakes across a gradient of average recent air temperature. A solid line indicates a p-value less than 0.05, and a dashed line indicates a p-value less than 0.1. Regression model details can be found in Table S3. Fish silhouettes were taken from PhyloPic (http://www.phylopic.org).
a) typical cold lake

i) thermal conditions

ii) species’ responses

b) warmed lake after climate change

iii) thermal conditions

iv) species’ responses
Figure 4: Schematic of the effects of warming on four widespread and abundant fish species of boreal shield lake food web: lake trout, cisco, walleye, and yellow perch. The x axis indicates shifts in the degree of nearshore foraging based on $\delta^{13}$C signatures. The y axis indicates shifts in habitat use in terms of the mean depth of capture. Lake trout, cisco, walleye all show shifts towards offshore foraging and offshore habitat use with increased temperature. Fish silhouettes were taken from PhyloPic (http://www.phylopic.org). Figure is adapted from Tunney et al. 2014.