Seasonal variation of total mercury and condition indices of Arctic charr (Salvelinus alpinus) in Northern Québec, Canada

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Abstract

The winter ecology of anadromous Arctic charr has remained poorly detailed in the literature beyond descriptions of seasonal fasting and resulting declines in condition. However, prolonged periods of reduced feeding can have significant consequences for other variables, such as tissue contaminant levels. To more thoroughly detail seasonal changes, biological information (fork length, total weight, age, sex, somatic condition), stable isotopes ($\delta^{13}$C, % carbon, $\delta^{15}$N, % nitrogen), dorsal muscle % lipid, caloric densities, and total mercury (THg) concentrations were assessed in anadromous Arctic charr collected from Deception Bay, Canada, during the summer and over-wintering periods. Significant reductions in somatic condition, total weight, and % nitrogen, consistent with prolonged periods of fasting, were found for post-winter captured Arctic charr, but % lipid and caloric densities were significantly higher in these fish. THg also varied seasonally and was significantly higher in summer collected tissue. When tested individually via linear regression, significant relationships were seasonally dependent, but limited in number. All previously mentioned parameters were then incorporated into multi-variable models which better explained variations in the data. While there was no clear best model for explaining the % lipid values, caloric densities, and THg, season, condition, and stable isotope values (% carbon and % nitrogen) were the best indicators of % lipid content and caloric densities. THg concentrations were best explained by total weight, somatic condition, and $\delta^{13}$C. Seasonal variation in fish condition measures and THg may be indicative of condition selective mortality that yields apparent improvement through the disproportionate removal of poorer conditioned fish from the population during the over-wintering period. This hypothesis was further supported by mortality estimates and the results of the multi-predictor variable models. Collectively, this research highlights the importance of seasonal dynamics for anadromous Arctic charr populations and suggests that future studies further consider seasonality when evaluating this species.
Keywords

Arctic charr; total mercury; Nunavik; seasonal variation; condition; lipid content; caloric density
1. Introduction

Arctic charr (*Salvelinus alpinus*) are the most northerly distributed freshwater fish species on Earth with populations ranging from southern temperate locations in eastern North America and the European Alps to the northern extent of land masses in Eurasia and North America (Johnson 1980). These fish are considered habitat generalists, occupying lakes, streams, rivers, and marine environments depending on the time of year and life history form (Power et al. 2008). Arctic charr exhibit diverse life history strategies (Jonsson et al. 1988) that include anadromous, lacustrine, and partial migratory types, where anadromous and non-anadromous fish co-exist (Jonsson and Jonsson 1993), with the decision to migrate dictated by environmental conditions, sex, genetics, and ontogenetic niche shifts (Jonsson and Jonsson 1993; Jørgensen and Johnsen 2014; Klemetsen et al. 2003a). Both sexually mature and immature fish perform migrations and first time migrants can be anywhere from 3 to 8 years of age (Johnson 1989; Nordeng 1983).

Seaward migrations of anadromous Arctic charr begin after ice break up in the spring (Johnson 1983; Klemetsen et al. 2003a), with fish generally remaining in coastal areas for 1-2 months before returning to freshwater environments (Klemetsen et al. 2003a; Mathisen and Berg 1968). The relative abundance of anadromous individuals is variable within and among populations (Johnson 1980; Svenning et al. 1992) and dependent on conditions, such as freshwater growth opportunities and differences in productivity between freshwater and marine environments (Gross et al. 1988) and/or the physical characteristics of the migratory route (Kristoffersen 1994; Moore et al. 2016). However, at the northern extremes of their distribution lacustrine residency is favoured regardless of access to the marine environment (Power et al. 2008).

For Arctic charr, the period of marine residency is characterized by rapid growth, with anadromous fish growing faster than resident freshwater fish (Berg and Berg 1989; Johnson 1980; Moore and Moore 1974). Fish may double their body weight and experience a five-fold increase in lipid content while in the marine environment (Finstad and Heggberget 1993; Jørgensen et al. 1997; Mathisen and Berg...
1968), with the carcass (head, skeleton, and skin) and muscle tissue accounting for 50% and 35-40%, respectively, of the total body lipid content when fish re-enter freshwater from the sea (Jobling et al. 1998; Jørgensen et al. 1997; Jørgensen and Johnsen 2014). During the marine period fish feed opportunistically on zooplankton, amphipods, pelagic and benthic fishes (Dempson et al. 2002; Grønvik and Klemetsen 1987; Power et al. 2008).

Historically, studies of anadromous Arctic charr have focused on the period of marine residency and its consequences for growth and maturation (Berg and Berg 1989; Jørgensen et al. 1997; Murdoch et al. 2015), but have traditionally lacked winter data (Mulder et al. 2018b). The available over-wintering data that does exist have focused on resident lacustrine Arctic charr condition, diet, and habitat during the ice-covered months (Amundsen and Knudsen 2009; Eloranta et al. 2013; Klemetsen et al. 2003b). Thus the ecology of over-wintering anadromous Arctic charr has remained poorly described in the literature beyond the noted cessation of feeding and the resulting reduction of condition (e. g. (Boivin and Power 1990; Jørgensen et al. 1997; Rikardsen et al. 2003). More recently, telemetry studies of over-wintering Arctic charr in southern Labrador have indicated reduced activity patterns (Mulder et al. 2018b) and temperature-dependent diurnal movement patterns (Mulder et al. 2019) occurring within a narrow range of temperatures (0.5–2 °C) that suggest a use of strategies to lower metabolic costs and minimize over-winter energy expenditure (Mulder et al. 2018a).

During winter, whole body lipid body reserves have been reported to decline by up to 30% for non-reproductive anadromous Arctic charr and between 35 – 46% for post-spawning individuals (Dutil 1986; Jørgensen et al. 1997). Lipids are depleted from all tissues, but mobilization of muscle and carcass deposits has been determined to be the most significant during this period (Jobling et al. 1998; Jørgensen et al. 1997). Additionally, emaciation is greatest in females with individuals on average losing approximately 80% of their lipid stores during spawning and overwintering (Jobling et al. 1998; Jørgensen et al. 1997). The documented declines in energy reserves, the minimization of movement, use of colder water temperatures to reduce metabolism (Mulder et al. 2018b), lack of suitable prey (Boivin and Power 1990), and the implications of light restrictions on prey capture efficiency (Mazur and
Beauchamp 2003; Turesson and Brönmark 2007) all suggest significantly reduced feeding during the winter months. While in lacustrine populations of Arctic charr, winter feeding has been documented to some extent in the literature (e.g., (Eloranta et al. 2013; Klemetsen et al. 2003b; Power et al. 2009)), it has only been inferred for anadromous individuals based on activity and habitat use (Boivin 1987; Mulder et al. 2018b).

In addition to losses in lipids and overall reductions in condition, prolonged periods of reduced feeding can have other significant consequences for over-wintering fish. Critical among those effects are the possible associated changes in tissue contaminant levels. Jørgensen et al. (2006) demonstrated that winter fasting and subsequent emaciation of anadromous Arctic charr resulted in the redistribution of lipophilic PCBs to sensitive organs (e.g., liver and brain). This was accompanied by significant increases in hepatic biomarker activity and disruptions to endocrine mechanisms, immune function, and smoltification processes. The results suggests that seasonal lipid dynamics may result in increased sensitivity of Arctic animals to certain contaminants undergoing seasonal fasting (Jørgensen et al. 2002; Jørgensen et al. 2006). Methylmercury (MeHg) is one contaminant of specific concern due to its neurologically toxic health effects (Mergler et al. 2007) and its ability to bioaccumulate in aquatic food webs to reach high concentrations in large-bodied predatory fish (Gantner et al. 2010; Kidd et al. 1995; van der Velden et al. 2013a). In the muscle tissue of these fish, MeHg concentrations are typically greater than 90% of total mercury, THg, (Eagles-Smith et al. 2016; Hall et al. 1997; Lockhart et al. 2005), and generally increase with size and age (Lescord et al. 2018), making THg concentrations a suitable proxy for MeHg levels and of particular concern to the Inuit people who consume large quantities of fish (Lemire et al. 2015).

Measured concentrations of THg are often related to biological variables such as: fish size, age, trophic position (δ¹⁵N), feeding strategies and habitat use (δ¹³C), growth, and somatic condition (Dittman and Driscoll 2009; Power et al. 2002; Wiener et al. 2003), but also depend on lipid and protein contents (Eisler 1987; Kahilainen et al. 2016; Swanson and Kidd 2010). THg in fish tissues is derived almost exclusively from prey consumption (Hall et al. 1997) and after digestion THg is translocated to the liver.
via blood and subsequently stored in muscle tissues (Oliveira Ribeiro et al. 1999; Wang and Wang 2015) where it is almost exclusively bound within proteins (Amlund et al. 2007; Bury et al. 2003; Mason et al. 1995). Growth during the summer can lower THg concentrations during the growing season as individuals accumulate tissue faster than THg (Karimi et al. 2007; Lepak et al. 2012; Olk et al. 2016), while lipid losses prompted by a cessation of feeding during the winter months can result in a phenomenon termed starvation–concentration (Cizdziel et al. 2003; Cizdziel et al. 2002) that increases mercury in remaining protein-containing tissues (Kahilainen et al. 2016). The result is higher THg during the ice-covered period (Keva et al. 2017; Olk et al. 2016). While seasonal variations in THg have been reported in lacustrine resident Arctic charr and other fish species (Kahilainen et al. 2016; Keva et al. 2017; Olk et al. 2016), such variations are relatively understudied and have yet to be documented for anadromous Arctic charr. Additionally, quantifying seasonal variations in THg concentrations in anadromous Arctic charr may help determine differences in seasonal risks for capture and consumption associated with the winter fishery for this species, which provides an important component of the year-round diet of Inuit people, including those of the Nunavik region of northern Québec (e.g., (Boivin and Power 1990)). Similar to other areas of Arctic Canada, anadromous Arctic charr in Deception Bay, Nunavik, Québec, are of significant cultural and economic importance to the local Indigenous communities (e.g., Salluit and Kangiqsujuaq). However, literature on Arctic charr populations in this region remains scarce (e.g., (Boivin and Power 1990; Murdoch et al. 2013; Murdoch and Power 2013)), with only one study having directly examined issues related to seasonality (Boivin and Power 1990).

Here samples collected from Deception Bay, Nunavik, where Arctic charr migrate and feed in summer, and lakes Duquet and Françoys-Malherbe, where Arctic charr spawn and over-winter, are used to investigate patterns of seasonal change in fish condition measures and THg concentrations. Specifically, it was hypothesized that (i) over-wintering anorexia and fasting (e.g., as represented by reduced total weight and somatic condition (Fulton’s K) (Jobling 1981; Jørgensen et al. 1997) would result in significant declines in percent lipid (% lipid) and caloric densities of anadromous Arctic charr collected during post-winter sampling as stored energy reserves are mobilized to meet metabolic demands
during this period (Jobling et al. 1998; Jørgensen et al. 1997). Additionally, seasonal relationships with co-measured biological information (fork length, total weight, age, sex, and somatic condition) and stable isotopes used as proxies for feeding behaviour, habitat use, and trophic position ($\delta^{13}$C, % carbon (C), $\delta^{15}$N, % nitrogen (N)) were evaluated.

Samples were also used to test a contamination hypothesis: (ii) that THg values would be higher during the post-winter sampling period than in the late summer sampling period as a result of fasting-induced emaciation and the subsequent increase of mercury in the remaining tissues (Kahilainen et al. 2016; Keva et al. 2017; Olk et al. 2016). Again, seasonal correlations with biological information and stable isotope ratios were assessed, with specific relationships between measured THg concentrations % lipid and caloric density values being evaluated. Finally, data were used to model dorsal muscle % lipid content, caloric density, and THg concentrations as a function of multi-variable statistical models inclusive of combinations of the above tested biological information, stable isotope values, and their interactions. Specifically, we hypothesized that (iii) season and/or fish condition would best describe % lipid values and caloric content (Dutil 1986; Thompson et al. 1991; Todd et al. 2008) and variables associated with bioaccumulation (fork length, total weight and/or age and trophic position represented by $\delta^{15}$N) (Cizdziel et al. 2002; Power et al. 2002; van der Velden et al. 2013a) and season (Kahilainen et al. 2016; Keva et al. 2017; Köck et al. 1996), would best describe THg concentrations as these variables are known to significantly influence measured THg concentrations in fish tissue (Kahilainen et al. 2016; Wiener et al. 2003).

2. Methods

2.1 Study Area

The Deception River and its tributaries (Fig. 1) are located east of Salluit, Québec (62°10’46 N, 75°40’13 W) with a watershed spanning an area of 3870 km² between latitudes 61°31’26” N and 62°11’01” N. The river flows into Deception Bay, which is associated with the Hudson Strait marine ecosystem (Goldsmit et al. 2014), a deep and wide channel that connects Hudson Bay and the Foxe Basin.
with the Labrador Sea and the Davis Strait. Average daily temperatures range from -25.6°C in February, to 10.5°C in August (Environment Canada 2018a; Environment Canada 2018b), with a growing season of less than 120 days per year. In addition to traditional hunting and fishing, the area is impacted by two nickel and copper mining projects: Glencore - the Raglan Mine Project and Canadian Royalties Inc. – the Nunavik Nickel Project, and is proximate to the now closed Asbestos Hill Mine (Purtiniq). A 95 kilometre road connects the main Raglan mining site with an additional camp and deep-water port in Deception Bay. Mine personnel are present year-round and the road closely follows the Deception River and its tributaries for most of its length. Some contaminant input is believed to result from the mining facilities proximate to Deception Bay in addition to atmospheric deposition.

Arctic charr spawn and overwinter in headwater lakes Duquet (62°03’18 N, 74°31’51 W) and Françoys-Malherbe (62°00’06 N, 74°15’25 W) from October to June. Lake Duquet is less than half the size of Lake Françoys-Malherbe and the lakes are located 2.5 km and 15 km, respectively, upstream of the river mouth. There is usually a commercial fishing permit active for Deception Bay and both lakes, and a Raglan Mine sport fishing permit is active for Deception Bay. Lake Watts, a third lake in the Deception River system upstream of Lake Françoys-Malherbe, is assumed to have received some direct input of mining waste prior to the 1984 closure of the Asbestos Hill Mine located 10 km from the lake. Traditional knowledge suggests that Arctic charr are predominately non-migratory in Lake Watts and access to this lake was not possible during the course of this study.

2.2 Sample Collection

Summer collection of anadromous Arctic charr was completed using multi-mesh experimental gill nets (25 – 150 mm mesh panels with a length of 120 m and a hanging depth of 2 m) set coincident with the returning upstream migration period. Arctic charr were caught at eight locations equally distributed throughout Deception Bay and from the mouth of the Deception River in August of 2016. A post-winter sample was obtained from lakes Françoys-Malherbe (May 8th – 11th, 2017) and Duquet (May 12th – 13th, 2017), approximately a month prior to ice break-up (as reported for the area by the Canadian Ice Service (2018) as June 26th, 2017). Post-winter dorsal muscle samples were collected from fish.
captured throughout the lakes with jigging lines by Nunavik Research Centre (NRC) staff in collaboration with Inuit elders during the Salluit community Elder’s Spring Fishing Event hosted by Qaqqalik Landholding Corporation and supported by Raglan Mine. All fish captured were sacrificed after capture with a sharp blow to the head and all sample collection was performed in accordance with standards dictated by the Ministère des Forêts, de la Faune, et des Parcs – Direction de la gestion de la faune du Nord-du-Québec (permis de gestion de la faune #2016-02-199-152-10-G-P N/D : 9053_36).

Fish were sexed, weighed (± 1 g) and measured for fork length (± 1 mm). Measurements were used to calculate Fulton’s condition factor (\( K = 10^5 \times \frac{W}{L^3} \)), which was only determined after performing standardized weight-length regressions and ensuring that the slope of this regression did not significantly deviate from a value of three (Ricker 1975). A sample of dorsal muscle tissue (mass ≈ 10 g wet weight) was removed from above the lateral line and posterior to the dorsal fin on the left side of each Arctic charr (van der Velden et al. 2013a) and frozen at -20°C for subsequent laboratory analyses. A random sample of 50 fish from the summer sampling was selected for THg analysis. A sub-sample of 40 fish was used for lipid analysis and a sub-sample of 30 fish was used for bomb calorimetry. From the winter sampling a total of 55 fish dorsal muscle samples were provided by the Salluit community Elder’s Spring Fishing Event. To guard against inadvertent use of resident fish from the post-winter sampling, anadromy was confirmed with \( \delta^{34} \text{S} \) stable isotope analysis following methods described in Doucett et al. (1999). After removal of resident fish, the remaining anadromous fish were used for THg analyses. Similar to the summer sampling, a sub-sample was retained for lipid and bomb calorimetry analyses. Dorsal muscle samples were generally used for all (lipid, bomb calorimetry and THg) analyses except where tissue was limited.

To minimize accidental metal contamination, tissues for THg analysis were placed in Eppendorf polypropylene tubes. These tubes had been acid washed in 15% HNO₃ for at least 24 hours before being rinsed 5 times with distilled water, twice with ultrapure water, and then dried under a laminar-flow fume hood before use. The upper gastrointestinal tracts (e.g., esophagus and stomach) of all post-winter collected fish were examined for evidence of short-term winter feeding. Aging of all sampled fish was
completed by NRC staff, with fish ages determined by submersing the otolith in water and examining the
distal surface with reflected light under a dissecting microscope (Chilton and Beamish 1982). Ages were
used to estimate von Bertalanffy growth equations (Ricker 1975) for each seasonal sample from which
the von Bertalanffy growth coefficient (K) was used to estimate mortality following (Jensen 1996).

2.3 Stable Isotope Analysis

For stable isotope analyses, dorsal muscle tissue was dried at 50°C for 48 h and pulverized into a
homogenate powder with a mortar and pestle. After being weighed to 0.275 – 0.300 mg (UMX2, Mettler-
Toledo GmbH, Greifensee, Switzerland), each tissue sample was simultaneously analyzed for carbon and
nitrogen stable isotope ratios (δ¹³C, δ¹⁵N) at the Environmental Isotope Laboratory, University of
Waterloo, Ontario, Canada with a Delta Plus Continuous Flow Stable Isotope Ratio Mass Spectrometer
(Thermo Finnigan, Bremen, Germany) coupled to a Carlo Erba Elemental Analyzer (CHNS-O EA1108,
Carlo Erba, Milan, Italy) following methods described in van der Velden et al. (2013a). Elemental
compositions were expressed in percentage terms based on pre-analysis weights. All stable isotope
measurements were expressed using standard delta notation (δ) as parts per thousand differences (‰)
with respect to the international reference standards of Vienna Peedee Belemnite carbonate rock for δ¹³C
(Craig 1957) and nitrogen gas in the atmosphere for δ¹⁵N (Mariotti 1983):

\[ \delta R = \left[ \frac{R_{\text{sample}} - R_{\text{standard}}}{R_{\text{standard}}} \right] \times 1000 \]

where δR is the measured carbon (¹³C/¹²C) or nitrogen (¹⁵N/¹⁴N) isotope ratio expressed with respect to
the appropriate international standard. Machine analytical precision was determined to be ± 0.2‰ for δ¹³C
and ± 0.3‰ for δ¹⁵N and was established via repeat analysis of internal laboratory working standards
(IAEA-N₁ + N₂, IAEA-CH₃ + CH₆) cross calibrated to International Atomic Energy Agency (IAEA)
standards: CH₆ for δ¹³C and N₁ and N₂ for δ¹⁵N. Internal standards were placed at the beginning, middle
and end of every run of samples and repeatability was assessed by repeat analysis of 1 in 10 samples. As
C:N ratios were consistently below the 4.0 threshold above which lipid extraction is required (Jardine et
al. 2011; Logan et al. 2008; Sanderson et al. 2009), δ13C values were not lipid extracted or mathematically normalized for lipid content.

### 2.4 Lipid Analysis and Bomb Calorimetry

Lipid analysis and bomb calorimetry were performed at the University of Waterloo, Waterloo, Ontario, Canada. A modified version of the procedure outlined in Folch (1957) was used for lipid extraction as this method provides accurate estimates of lipid content when lipids comprise greater than 2% of tissue (Iverson et al. 2001). After freeze drying (Freezone Plus 2.5 Liter Cascade Benchtop Freeze Dry Systems, Labconco, Kansas City, USA), Arctic charr dorsal muscle tissue was ground with a mortar and pestle and weighed to approximately 30.0 mg ± 0.1 mg (XS205DU Analytical Balance, Mettler Toledo, Mississauga, Canada). 2 mL of a 2:1 chloroform-methanol solution and 1.6 mL of a 0.9% KCl solution were then added to the ground tissue. The resulting solution was homogenized with a vortex (Fisherbrand Analog Vortex Mixer, Fisher Scientific, Hampton, USA) and centrifuged (Fisherbrand™ Centrifuge Model 225A Benchtop Centrifuge, Fisher Scientific, Hampton, USA) at 2000 RPM for 5 minutes until the KCl, tissue, and chloroform-methanol layers were completely separated. The lipid containing solution was extracted via Pasteur pipette through the KCl and residual biomass layers. Three iterations of the procedure were performed until a final lipid solution of 8 mL was obtained. The lipid-containing solution was then evaporated to dryness. Once dry, an additional 2 mL of 2:1 chloroform-methanol solution were added to the dried material and two, 100 μL aliquots were then transferred to pre-weighed tin cups. The solution was evaporated at room temperature overnight until only dry lipids remained. Remains were weighed on a micro-balance (UMX2, Mettler-Toledo GmbH, Greifensee, Switzerland) to determine the percentage of lipid in the dorsal muscle tissue expressed as:

\[
\% \text{ Lipid} = \left( \frac{\text{Mass}_{\text{Dry}}(\text{g})}{\text{Mass}_{\text{Ground}}(\text{g})} \right) \cdot (1 - P_{\text{Water}}) \cdot 100\% \cdot 20;
\]

where Mass_{Dry} is the weight of the dried lipid following the extraction procedure, Mass_{Ground} is the initial ground mass of the tissue preceding extraction, P_{Water} is the proportion of water in the analyzed dorsal muscle tissue (wet tissue mass (g) – dry tissue mass (g)), and 20 represents a correction as a result of
using only a subset of the final extraction solution volume for establishing the final dried mass (Folch 1957).

For bomb calorimetry, Arctic charr dorsal muscle tissue was dried at 50°C for 48 hours and ground to obtain a homogenized sample using a mortar and pestle. Pellets were formed (Parr Pellet Press, Parr Instrument Company, Moline, USA) with weights not exceeding 50.0 mg ± 0.1 mg (XS205DU Analytical Balance, Mettler Toledo, Mississauga, Canada) before ignition in a Parr Semi-micro Calorimeter 6725 (Parr Instrument Company, Moline, USA) to measure caloric density (cal·g\(^{-1}\) dry mass).

The wet mass caloric density was determined by multiplying the dry mass caloric density by the proportion of final dry mass to original wet mass (Glover et al. 2010). Results were reported as wet weight (ww) caloric density means ± standard deviation. Benzoic acid pellets with a caloric density of 6318.4 cal·g\(^{-1}\) dry weight (dw) were used to standardize the calorimeter and assess recovery every 10\(^{th}\) sample. Percent recovery ± standard deviation of benzoic acid pellets (\(n = 11\)) was determined to be 100.35% ± 1.07%.

2.5 THg Analysis

Mercury analysis was performed at the Institut National de la Recherche Scientifique (INRS) in Québec City, Québec, Canada and at the University of Waterloo in Waterloo, Ontario, Canada. After freeze-drying (FTS Systems TMM, Kinetics Thermal Systems, Longueil, QC, Canada; Freezone Plus 2.5 Liter Cascade Benchtop Freeze Dry Systems, Labconco, Kansas City, USA), tissue was weighed to approximately 50.0 – 100.0 mg ± 0.1 mg (Series 321 LT 220A Balance, Precisa Gravimetrics AG, Dietikon, Switzerland; Mettler Toledo, Mississauga, Canada). Analysis was then performed with a direct mercury analyzer (DMA-80, Milestone Inc., Shelton, USA) which enables assessment of THg through thermal decomposition followed by atomic absorption spectroscopy as described in U.S. EPA method 7473 (United States Environmental Protection Agency 1998). Results were converted from dry weight to wet weight using percent moisture calculations determined from weights (± 0.1 mg) taken before and after lyophilisation (Eikenberry et al. 2015).
Past studies have indicated that age (van der Velden et al. 2013b) and/or length (Rigét et al. 2000; Swanson et al. 2011) are often strongly correlated with THg concentrations such that THg concentrations require age or length adjustment to permit comparisons among individuals. Accordingly, data were examined for evidence of significant length and/or age correlations using linear regression (log_{10} THg vs. log_{10} fork length, fork length, or log_{10} age, and age) (Tran et al. 2015). The effect of lipids on analytical results was examined by comparing differences in relationships obtained using raw and lipid corrected THg concentrations, with lipid corrected THg concentrations computed following methods described in Kahlilainen et al. (2016).

Method detection limits and percent recoveries are reported as mean percentage of certified values ± standard deviation. Tissues were evaluated in triplicate with certified reference materials from the National Research Council of Canada (NRCC) (TORT-3, DOLT-4, and DOLT-5). Blanks were used every fifth sample in the same analytical cycle to establish accuracy and recovery rates. The method detection limit, determined as 3× the standard deviation of blanks, was 2.91 ng Hg (approximately 0.003 (mg·kg^{-1})) and mean relative standard deviation of the triplicates was 5.53% (n = 108). Percent recoveries were determined to be 99.32% ± 7.42% (n = 52), 93.16% ± 9.04% (n = 47), and 88.95% ± 3.88% (n = 16) for TORT-3, DOLT-4, and DOLT-5 respectively.

2.6 Statistical Analysis

All statistical analyses were performed using JMP® statistical software (v. 13.0.0, SAS Institute, CA) and Type I error was set to \( \alpha = 0.05 \). Data consistency with normality and homoscedasticity assumptions were verified using residual diagnostic histograms, visual assessment of Q-Q plots, and the Shapiro-Wilk W test (Shapiro and Wilk 1965). Data that did not meet parametric assumptions were log_{10} transformed (Zar 2007). Linear regressions were used to determine the relationship significance between specific variables and outliers that may have unduly influenced regression results were assessed using Cook’s Distance statistic (Cook 1977) and subsequently removed. Un-paired, two-sample t-tests adjusted for homogeneity of variance assumptions were used to determine significant differences among seasons (Zar 2007) and the Wilcoxon approach was used when data did not conform to the required parametric
The Fisher exact test was used to test for significant differences among proportions (Zar 2007) and incidental correlations were assessed using Pearson's correlation coefficient. General linear models (GLM) inclusive of season, fork length, total weight, age, somatic condition, $\delta^{13}$C, $\%$ C, $\delta^{15}$N and $\%$ N were used to determine the best model to describe dorsal muscle $\%$ lipid values, caloric densities, and THg concentrations. Pearson correlation analysis (Appendix A Tables A.1 – A.3) and/or linear regression were used to establish the significance of variables expected to correlate with dorsal muscle $\%$ lipid values, caloric densities, and THg concentrations, with only significant variables being retained for use in GLM models. Significant two-way interactions among variables were similarly assessed, with significant interactions retained for use in GLM models. To reduce statistical issues associated with multicollinearity (Zar 2007), possible explanatory variables that were highly correlated ($r^2 > 0.70$) or known to have biological redundancy (e.g., fork length and total weight) were not included in the same model. To predict seasonal dorsal muscle $\%$ lipid values, caloric densities, or THg concentrations models that included all possible combinations of the feasible set of significant explanatory variables and two-way interactions, as determined above, were considered. Model selection was performed using the Akaike Information Criteria adjusted for small sample sizes ($AIC_c$). The model with the lowest $AIC_c$ score was considered the most accurate, except in circumstances where $AIC_c$ values differed by less than two (Burnham and Anderson 2003). Detailed description of the computation of the analytical metrics associated with the $AIC_c$ methodology used here can be found in Burnham and Anderson (2003).

3. Results

Of the 105 anadromous Arctic charr samples obtained, 100 were used in the comparison of summer and post-winter levels of $\log_{10}$ THg, $\log_{10}$ $\%$ lipids, and $\log_{10}$ caloric densities. A single sample from the summer sampling was removed from use because of sample desiccation prior to shipment. Testing with $\delta^{34}$S stable isotope analyses indicated 4 Arctic charr from the post-winter collection were
non-anadromous and these fish were excluded from further consideration. This resulted in the loss of 4347 Arctic charr samples from the lipid testing subset and 3 samples from the calorimetry testing subset.

3.1 Seasonal Variation in Biological Variables and Stable Isotope Values

Means ± standard deviations and ranges for biological variables and stable isotope values for all summer sampled and post-winter captured Arctic charr can be found in Table 1. Statistical testing of the slope of the standardized weight-length regression (Deception Bay sampling – 3.12, post-winter sampling – 3.03) confirmed isometric growth (Deception Bay t-test, \( p = 0.457 \), post-winter sampling t-test, \( p = 0.255 \)), thereby allowing the use of Fulton's K. Estimated seasonal von Bertalanffy growth models yielded von Bertalanffy growth coefficients from which differing mortality rates were estimated (summer mortality: 22.3%; winter mortality: 24.5%). Log_{10} somatic condition significantly declined in post-winter Arctic charr (Z_{1,98} = 4.637, \( p < 0.001 \)) indicating possible over-wintering anorexia, as a result of a significant decline in log_{10} total weight during the over-wintering period (Z_{1,98} = 2.444 , \( p = 0.015 \)). Log_{10} fork length \( t_{1,98} = 1.43, \ p = 0.156 \), log_{10} age \( t_{1,95} = 1.309, \ p = 0.194 \), % carbon \( Z_{1,98} = -0.831, \ p = 0.406 \), and \( \delta^{15}N \) values \( t_{1,98} = -1.258, \ p = 0.211 \) of captured fish did not vary seasonally. \( \delta^{13}C \) was significantly more depleted in post-winter sampled Arctic charr \( Z_{1,98} = -5.754, \ p < 0.001 \) as was % N \( Z_{1,98} = 5.623, \ p < 0.001 \). The sex ratio of summer and post-winter sampled fish did not vary between seasons (Fisher’s exact \( \chi^2 = 0.440, \ p < 0.657 \) and sex was not significantly correlated with % lipids, caloric density, or THg in any of the seasonal samples (all correlation \( p > 0.466 \)). Furthermore, no evidence of short-term winter feeding was determined via examination of the upper gastrointestinal tract.

3.2 Lipid Content and Caloric Density

Means ± standard deviations and ranges for % lipid values and caloric densities are given in Table 2. Significant seasonal variation existed for log_{10} lipid content \( t_{1,74} = -6.49, \ p < 0.001 \) and log_{10} caloric density \( t_{1,55} = -11.70, \ p < 0.001 \) (Fig. 2). Log_{10} lipid content explained 28% of the variation in log_{10} caloric density of Arctic charr during the post-winter fishery \( r^2 = 0.28, \ p = 0.005 \), but values were not significantly correlated in Arctic charr captured returning from the marine environment \( r^2 = 0.09, \ p = 0.340 \).
0.118). Significant relationships between $\log_{10}$ % lipid values and $\log_{10}$ caloric densities and studied correlates are plotted in Fig. 3 and Fig. 4. A limited number of weak significant relationships were found and there was no consistent pattern of associations when comparing summer and post-winter samples.

Log$_{10}$ % lipids were not significantly correlated with any of the biological parameters (e.g., length, weight). However, log$_{10}$ % lipid values of late summer migrants significantly declined with increased offshore feeding ($r^2 = 0.12, p = 0.030$) and were generally related to % elemental composition, increasing with % C in post-winter sampled fish ($r^2 = 0.11, p = 0.048$) and decreasing with % N in both summer and post-winter sampled fish (summer ($r^2 = 0.14, p = 0.017$); post-winter ($r^2 = 0.12, p = 0.035$)) (Fig. 3). Log$_{10}$ caloric density increased with fork length in summer sampled Arctic charr ($r^2 = 0.16, p = 0.031$) (Fig. 4) and in post-winter sampled fish was positively related to % C ($r^2 = 0.30, p = 0.003$) and negatively related to % N ($r^2 = 0.43, p < 0.001$) (Fig. 4).

### 3.3 THg

Mean THg concentrations ± standard deviations are reported in Table 2 and all measured concentrations were determined to be below the regulatory 0.5 mg·kg$^{-1}$ of mercury detailed in Health Canada’s commercial guideline of maximum levels for chemical contaminants in foods (Government of Canada 2019; Health Canada 2018). Contrary to what was hypothesized, log$_{10}$ THg concentrations of summer captured Arctic charr were significantly higher than those obtained from post-winter fish ($t(1.98) = 2.59, p = 0.011$) (Fig. 2). Significant relationships between log$_{10}$ THg concentrations and tested variables in both summer and post-winter sampled Arctic charr are plotted in Fig. 5. Regressions of fork length, weight and age against log$_{10}$ THg concentrations in summer or post-winter sampled fish were either non-significant or yielded regressions that poorly explained variation in log$_{10}$ THg concentrations (summer – log$_{10}$ THg vs. age ($r^2 = 0.20, p = 0.001$); post-winter – log$_{10}$ THg vs. fork length ($r^2 = 0.09, p = 0.034$); post-winter – log$_{10}$ THg vs. weight ($r^2 = 0.14, p = 0.006$)). Additionally, significant heteroscedasticity indicative of their statistical inadequacy was displayed, as such there was no compelling statistical evidence for length or age standardization of the log$_{10}$ THg concentrations data. Examination of differences in the relationships between tested variables and raw and altered log$_{10}$ THg values also
provided no compelling evidence for the correction of log_{10} THg concentrations for lipids following (Kahilainen et al. 2016), as relationships between the parameters did not significantly change after transformation. Therefore, raw log_{10} transformed results were used in all subsequent statistical analyses.

Weak and seasonally differentiated patterns of association between log_{10}THg and stable isotope data (δ^{13}C, % C, δ^{15}N) were found (Fig. 5). Log_{10} THg increased with δ^{13}C (r^2 = 0.23, p < 0.0001) in summer and δ^{15}N in post-winter sampled fish (r^2 = 0.09, p = 0.031), and decreased with % C only in summer sampled Arctic charr (r^2 = 0.19, p = 0.002). Significant relationships between log_{10}THg concentrations and log_{10} % lipid values or log_{10} caloric densities showed significant association (negative) only with log_{10} % lipids (Fig. 5) in late summer captured Arctic charr (r^2 = 0.20, p = 0.018).

3.4 General Linear Models

Models best explaining variation in log_{10} % lipid values, log_{10} caloric densities and log_{10}THg concentrations are reported in Table 3. AICc model selection yielded multiple models with a Δ_i within 2 of the best model that were considered as essentially as good as the best model. For log_{10} % lipid values these included two models including subsets of the variables season, % C, and % N, although Akaike weight (w_i) and evidence ratio (ER_i) measures were better for the model including all three variables. For caloric density there were three equivalent models that included combinations of the variables: season, age, condition, fork length, total weight, % C and % N. Akaike weight (w_i) and evidence ratio (ER_i) measures suggested better support for the models including either age or fork length as variables, rather than weight. Similarly for log_{10}THg, three equivalent models inclusive of subsets of the variables season, total weight, condition δ^{13}C and δ^{15}N were estimated, with the model including season having the lowest likelihood of being the best approximating model for describing the log_{10} THg concentration data.

4. Discussion

Somatic condition and weight declined in post-winter captured fish as hypothesized. Significant seasonal differences in %N (decline) and δ^{13}C (increase) were also observed, whereas no significant seasonal variation was evident for the other biological parameters. Significant seasonal variation existed
for lipid content and caloric density, but contrary to what was hypothesized, values determined from Arctic charr collected during the post-winter sampling were significantly higher than those determined from fish returning from the marine environment in late summer. Percent lipid values and caloric densities were correlated in the tissue of post-winter sampled Arctic charr, but were not significantly related in late summer migrants. Additionally, significant relationships between % lipid values and caloric densities existed between some, but not all of the studied variables of interest. Seasonal variation existed for THg concentrations and was contrary to what was expected, declining in post-winter sampled Arctic charr. THg concentrations were significantly related to some, but not all, of the tested variables and were unrelated to % lipid values and caloric densities, except in late summer sampled Arctic charr where concentrations significantly declined with increasing lipid content. Models best supporting the data from the previously stated analyses better explained variation in data when inclusive of multiple parameters. While no single model was exclusively best, models generally included variables considered of traditional importance (i.e., fish size, condition and feeding patterns) as well as season.

4.1 Seasonal Variation in Biological Variables and Stable Isotope Values

Condition, driven by decreases in mean weight rather changes in mean length, declined between seasonal sampling periods, with mean K values of post-winter collected Arctic charr being significantly less than those of returning marine migrants. Similar significant over-winter reductions in somatic condition have been reported for Norwegian resident lacustrine Arctic charr despite winter feeding (Amundsen and Knudsen 2009; Klemetsen et al. 2003b). However, end of winter condition for non-reproductive individuals captured near Kangiqsualujjuaq, Nunavik, Québec, was reported to consistently exceed 1.0 (Boivin and Power 1990). Thus, variation in the extent of post-winter declines in condition is apparent among populations and likely among years.

Declines in condition appear linked to reduced seasonal feeding reflected in the seasonal differences of % N in the muscle tissue, with declines in % N having been reported for resident lacustrine Arctic charr after prolonged periods of fasting (Power et al. 2009). Reductions in % N have similarly been demonstrated in other species (Elliott 1976; Guerin-Ancey 1976; Van Weerd et al. 1995) and are
associated with continuous losses of nitrogenous compounds via waste products during fasting (Elliott 1976; Hobson et al. 1993; Steele and Daniel 1978). These losses are substantive when compared to continuously fed fish (Van Weerd et al. 1995). During prolonged periods of fasting, enrichment of δ¹⁵N values is anticipated via similar catabolization processes associated with reductions in % N values (Gannes et al. 1997; Hobson et al. 1993; McCutchan Jr et al. 2003). While here tests of differences in δ¹⁵N between periods showed no significant changes, the differences in mean size between periods may have masked the extent of the overall increase in δ¹⁵N occurring during the fasting period. For example, the significant relationship between fork length and δ¹⁵N often reported in the literature (e.g., (Gantner et al. 2010; van der Velden et al. 2013a) and seen here ($r^2 = 0.501, p < 0.031$) yields a δ¹⁵N prediction range (12.51-13.04l) for summer fish of a size equivalent to the observed mean size of post-winter fish that does not include the mean reported δ¹⁵N value (13.33l) found for post-winter fish. Furthermore, the overall absolute increase in δ¹⁵N between periods (0.27l) observed here fell within the 95% confidence limits (0.26-0.74l) of the mean effect size for starvation induced changes in δ¹⁵N estimated from a meta-analysis of fasting studies (Hertz et al. 2015). Overall, the evidence suggests a biologically significant increase in δ¹⁵N occurred as a result of fasting in over-wintering fish. Similarly, use of lipid reserves as an over-winter energy source (e.g., (Dutil 1986; Jørgensen et al. 1997)), typically in the range of -24.9l to -28.5l for salmonids (Aursand et al. 2000), suggests observable declines in δ¹³C would occur over the winter period as tissues equilibrated over time with catabolized lipid reserves (e.g., (Herzka and Holt 2000)). The observed absence of short term feeding by post-winter captured Arctic charr, established here through an analysis of the upper gastrointestinal tract, in conjunction with noted declines in weight, somatic condition, %N, and δ¹³C and the directional shifts in δ¹⁵N are all indicative of prolonged periods of fasting exceeding several months consistent with the over-wintering period (Hesslein et al. 1993; Power et al. 2009).

4.2 Lipid Content and Caloric Density

Somatic condition, weight loss and % N values indicative of prolonged periods of fasting are consistent with previous literature detailing the over-winter period for Arctic charr (Amundsen and...
Over-winter changes in lipid content and caloric densities ran contrary to what was hypothesized and were significantly higher in the dorsal muscle tissue of post-wintering Arctic charr. Elevated post-winter % lipid values and caloric densities might be indicative of winter feeding, as previously inferred through observations of foraging behaviour (Boivin 1987) and examination of winter movement activities (Mulder et al. 2018b). However, evidence for the absence of short term feeding provided by empty stomachs and upper gastrointestinal tracts, reductions in somatic condition, weight, and % N, is more consistent with prolonged periods of non-feeding (Hesslein et al. 1993; Power et al. 2009). While previous literature (Jobling et al. 1998; Jørgensen et al. 1997) has reported female anadromous Arctic charr exhibiting significantly greater depletion of lipid stores compared to their male counterparts during the over-wintering months, sex was not correlated with % lipid or caloric densities in this study. The lack of a significant relationship between sex and fish condition indices measures has been observed with other species (Kitts et al. 2004; Usydus et al. 2012), with differences among studies possibly related to the proportion of spawning ready adults (Henderson and Tocher 1987; Jobling et al. 1998; Medford and Mackay 1978).

Considered together, the evidence of over-winter fasting, increased mortality, reduced weight and condition, and paradoxical increases in % lipids and caloric densities suggest condition selective mortality may be operating over the winter period. Condition selective mortality acts through a range of phenotypic and genotypic variables to remove poorly conditioned individuals from a population with consequences for evolution and population demographics (Chen and Maklakov 2012; Gagliano et al. 2007; Ronget et al. 2017). Sources of condition dependent mortality often include starvation, thermal stress, predation, failure to transition between ontogenetic life stages, and the interactions between the multiple factors (Gagliano et al. 2007; Miranda and Hubbard 1994; Ronget et al. 2017). For a species such as Arctic charr, over-wintering in low productivity, oligotrophic environments, and given the evidence noted above, starvation would appear to be the mechanism of most interest.

Starvation or reduced feeding is a consistent driver of over-wintering and selective mortality and has been implicated as the cause of mortality for different life stages of several fish species, including...
Arctic charr (Biro et al. 2004; Byström et al. 2006; Post and Evans 1989). Fish size often has considerable influence, as smaller fish with higher metabolic demands and lower lipid and protein reserves relative to larger conspecifics deplete critical energy reserves at an escalated rate, resulting in more rapid starvation and higher subsequent mortality (Henderson et al. 1988; Smith and Griffith 1994; Thompson et al. 1991). For example, studies of condition selective mortality among age-0 walleye Pollock (*Theragra chalcogramma*) showed that while lipid stores and body condition for the test group as a whole were rapidly reduced by starvation, survivors had significantly higher lipid content than mortalities, with values often exceeding those of pre-starvation fish (Sogard and Olla 2000). Additionally, Searcy and Sponaugle (2001) examined mortality as a function of early life history traits (size-at-age and growth rates) at critical periods in the Bluehead wrasse (*Thalassoma bifasciatum*) and the Slippery dick (*Halichoeres bivittatus*) and similarly noted that the better conditioned fish survived. The effect of condition selective mortality, therefore, would be to shift the mean of the trait distribution in ways that would yield no apparent effect of over-wintering on lipid and body reserves, or as here, apparent improvement as a result of the systematic removal of poorer conditioned individuals. Further, condition selective mortality should increase apparent mortality within the population as was also noted here.

The life history of Arctic charr further argues for the influence of condition selective mortality. Arctic charr are a long lived fish species (Johnson 1983; Johnson 1989; Power et al. 2008) and the prevalence of condition dependent mortality has been theoretically linked with increased life spans (Chen and Maklakov 2012). Arctic charr are also fall spawning and body lipids in fish generally decrease coincident with maturation as individuals mobilize lipids into maturing gonads (Henderson and Tocher 1987). In Arctic charr, the overall change in body lipids maybe as high as 30–80% after spawning and over-wintering (Dutil 1986; Jobling 1981; Jørgensen et al. 1997) with post-spawners being much more depleted when compared to non-reproductive individuals (Dutil 1986). Spawning has also previously been linked to increased over-winter mortality in Largemouth bass, *Micropterus salmoides* (Post et al. 1998). If condition-dependent mortality was influencing the Deception River population, it would be expected to increase the mortality of spawned fish. Given the inverse relationship between body lipid
content and the gonadosomatic index (Henderson and Tocher 1987), fish preparing to spawn would have lower muscle tissue reserves in comparison to non-spawning fish. Higher mortality among spawning fish with lower muscle lipid reserves would, therefore, effectively act to remove a greater proportion of the returning migrants at the lower end of % lipid spectrum. The net effect would be an increase in measured mean % lipids among over-winter survivors, yielding the pattern of seasonal differences in % lipid and caloric density values observed here. While we did not measure the proportion of spawning ready adults, the seasonal effect can be expected to be particularly strong in years when there is a high proportion of spawning ready fish among the marine re-entry migrants.

4.3 THg

Quantification of seasonal variation in THg concentrations determined all measured values were below the regulatory 0.5 mg·kg\(^{-1}\) of mercury recommended in Health Canada’s commercial guideline of maximum levels for chemical contaminants in foods (Government of Canada 2019; Health Canada 2018). Thus the seasonal risks for capture and consumption associated with the winter fishery for anadromous Arctic charr in the Deception River system appear minimal. However, further evaluation of seasonal THg consumption risks for the system’s resident Arctic charr is recommended given previously documented higher winter THg concentrations in other lacustrine resident Arctic charr populations (Kahlilainen et al. 2016) and consistently higher THg concentrations in lake-dwelling life history forms for this species (Swanson et al. 2011; Van der Velden et al. 2015; van der Velden et al. 2013b).

Hypothesized over-wintering anorexia, prompted by a cessation of feeding by anadromous Arctic charr during the winter months, leading to seasonal energy reserve losses and increases in THg concentrations (Cizdziel et al. 2002; Kahlilainen et al. 2016; Keva et al. 2017) was not observed in the Deception fish. Rather, THg concentrations were significantly higher in tissue collected from Arctic charr sampled during the summer in the marine environment. Similar to seasonal % lipid and caloric density data, seasonal differences in THg concentrations may have resulted from the consequences of condition selective mortality, as suggested by estimated seasonal increases in mortality. The removal of poorly conditioned, lower lipid content individuals via condition selective mortality would increase measured
mean lipid content, which itself is often negatively correlated with THg concentrations (Post et al. 2007; Wiener et al. 2003). Thus, better conditioned post winter survivors with higher lipids would shift the distribution of sampled THg concentrations, yielding a decrease in mean THg.

Differences in prey THg content in the seasonally occupied habitats and elimination of tissue THg concentrations, can be reasonably excluded as potential drivers of the observed results. The absence of short term prey intake, reduced somatic condition, weight, and % N point to a significant period of fasting. Further, as elimination is a metabolically-dependent process correlated positively with temperature (Trudel and Rasmussen 1997) it is unlikely to have played a major role in determining seasonal differences because of the narrow range of cold (0.5-2°C) temperatures occupied by over-wintering Arctic charr to reduce metabolism (Mulder et al. 2018a). The continued loss of proteins during fasting via nitrogenous waste products (Elliott 1976; Hobson et al. 1993; Steele and Daniel 1978), e.g., reduced % N in post-winter fish, may have contributed to the declines in measured THg as THg in fish tissues is almost exclusively bound within proteins (Amlund et al. 2007; Bury et al. 2003; Mason et al. 1995). The increase in caloric densities, which includes consideration of protein content, in post-winter Arctic charr, however, suggests that protein loss during the over-wintering period did not substantially influence seasonal variation in THg concentrations in Deception River anadromous Arctic charr. The elimination and depuration of THg is remarkably slow (Amlund et al. 2007; Oliveira Ribeiro et al. 1999), implying that physiological processes affecting protein content were not the main driver of the significantly reduced THg concentrations observed in post-winter samples.

4.4 General Linear Models

While there were no clear best models for explaining the % lipid values, caloric densities, and THg concentrations data, all dependent variable models were better supported when containing multiple variables. The % lipids models all depended on season and % N, with the best model also including consideration of % C. Key for % lipids then are variables associated with controlling food intake (season) and/or the metabolic consequences of fasting (% N), with season and metabolism known to hold implications for body energy reserves in fish (Dutil 1986; Jørgensen et al. 1997; Thompson et al. 1991)
and changes in % N being linked to continuous, non-replaced N loses via excretion during periods of fasting (Post et al. 2007; Power et al. 2009). In contrast to % lipids, variation in caloric densities were linked to a suite of biological and feeding variables, predominantly season, and variables reflective of feeding status. Results here parallel previous research detailing seasonal feeding and its implications for somatic condition and body reserve depletion in Arctic charr (Amundsen and Knudsen 2009; Rikardsen et al. 2003; Sæther et al. 1996). Additionally, the inclusion of variables that have associations with both size and reduced feeding were consistent with the hypothesis that condition selective mortality may be exerting an influence on this population of Arctic charr, as size, starvation, and reduced feeding are commonly associated with condition selective mortality (Biro et al. 2004; Byström et al. 2006; Sogard and Olla 2000). Finally, models for describing variations in THg concentrations accorded with previous research that has detailed relationships between metal concentrations and weight and condition (Dittman and Driscoll 2009; Eastwood and Couture 2002; Swanson and Kidd 2010), season (Keva et al. 2017) and indicators of diet and feeding habits (Goutte et al. 2015; Power et al. 2002; van der Velden et al. 2013a). Collectively, the multivariate evidence for each of the studied variables suggests that single variable correlations alone are likely insufficient for understanding variations in the environmental conditions associated THg concentrations in anadromous Arctic charr.

4.5 Conclusions

In addition to addressing knowledge gaps associated with available regional and population specific data, this research has increased the limited scientific data available on patterns of metal contamination in the over-wintering anadromous Arctic charr. To our knowledge, the observed results are the first detailing higher lipid content and caloric density in Arctic charr captured during the winter months, a period that has previously been associated with significant reductions in body reserves. Condition selective mortality is argued to be the most plausible explanation for the observed results, particularly as concurrently measured variables (e.g., % N, condition, and weight loss) suggested prolonged periods of fasting coincident with the known seasonal feeding behaviour of Arctic charr. Relationships between THg and descriptive biological variables were consistent with the seasonally
dependent hypothesis, although season itself was not the single best predictor of THg concentrations. Rather, multi-predictor models better described variation in the data for THg, % lipids and caloric densities, with variables such as season, somatic condition, age, body size (length or weight) and feeding tactics ($\delta^{13}$C, $\delta^{15}$N) being implicated in varying combinations for the best description of the data sets. While there is large literature examining the associations between THg and biological or feeding variables, much less is known about the effect of season. Quantification of seasonal variation in THg concentrations has determined that winter consumption risks for anadromous Arctic charr in this region are minimal. However, further work is necessary to elucidate global patterns of seasonal THg accumulation, especially as seasonal trends may differ between life history forms for this species. Overall, our corroboration of earlier studies that highlight the importance of season suggests further work is required to understand seasonality in Arctic charr, especially in the face of climate change and the likely effects it will have on Arctic aquatic environments.
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### Tables

**Table 1** Means ± standard deviations and ranges are given for fork lengths, total weights, ages, somatic condition, δ\(^{13}\)C, % carbon (C), δ\(^{15}\)N, and % nitrogen (N) of Arctic charr used from the summer 2016 and post-winter 2017 sampling seasons. Sex ratios are also noted. Significant seasonal differences are denoted with * (p < 0.05), ** (p < 0.001), and *** (p < 0.0001).

<table>
<thead>
<tr>
<th>Season of Capture</th>
<th>Sample Size</th>
<th>Fork length (mm)</th>
<th>Total weight (g)</th>
<th>Age (Years)</th>
<th>Sex</th>
<th>Condition (K)</th>
<th>δ(^{13})C (‰)</th>
<th>% C</th>
<th>δ(^{15})N (‰)</th>
<th>% N</th>
</tr>
</thead>
<tbody>
<tr>
<td>Summer 2016</td>
<td>49</td>
<td>483.5 ± 107.1</td>
<td>1380 ± 819*</td>
<td>9.6 ± 2.4</td>
<td>67% Female</td>
<td>1.06 ± 0.28***</td>
<td>-19.42 ± 0.99***</td>
<td>47.36 ± 3.06</td>
<td>13.06 ± 1.11</td>
<td>14.00 ± 0.82***</td>
</tr>
<tr>
<td></td>
<td></td>
<td>143.0; 689.0</td>
<td>20; 3300</td>
<td>5; 15</td>
<td>33% Male</td>
<td>0.57; 2.38</td>
<td>-22.97; -16.82</td>
<td>35.72; 52.19</td>
<td>8.12; 15.00</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
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<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Post-Winter 2017</td>
<td>51</td>
<td>449.4 ± 130.2</td>
<td>1031 ± 824</td>
<td>8.9 ± 2.9</td>
<td>73% Female</td>
<td>0.91 ± 0.11</td>
<td>-20.55 ± 1.00</td>
<td>48.30 ± 2.58</td>
<td>13.33 ± 1.01</td>
<td>12.92 ± 1.05</td>
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<td></td>
<td></td>
<td>221.0; 698.5</td>
<td>99.8; 3230</td>
<td>5; 20</td>
<td>27% Male</td>
<td>0.71; 1.46</td>
<td>-22.75; -16.89</td>
<td>44.03; 57.51</td>
<td>11.23; 15.22</td>
<td>8.60; 14.64</td>
</tr>
</tbody>
</table>
Table 2 Means ± standard deviations and ranges of dorsal muscle % lipid values, caloric densities, and THg concentrations of anadromous Arctic charr captured in the summer 2016 and post-winter 2017 sampling periods. *(p < 0.05), **(p < 0.001), and *** (p < 0.0001) indicate significant seasonal variation.

<table>
<thead>
<tr>
<th>Location of Capture</th>
<th>Sample Size</th>
<th>Lipids (%)</th>
<th>Sample Size</th>
<th>Caloric Density (cal·g⁻¹)</th>
<th>Sample Size</th>
<th>THg (mg·kg⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Summer 2016</td>
<td>40</td>
<td>4.08 ± 2.00</td>
<td>30</td>
<td>1327.6 ± 51.6</td>
<td>49</td>
<td>0.12 ± 0.05*</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2.15; 11.76</td>
<td></td>
<td>1180.8; 1409.8</td>
<td></td>
<td>0.06; 0.26</td>
</tr>
<tr>
<td>Post-Winter 2017</td>
<td>36</td>
<td>8.34 ± 4.85***</td>
<td>27</td>
<td>1545.8 ± 88.5***</td>
<td>51</td>
<td>0.09 ± 0.05</td>
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<tr>
<td></td>
<td></td>
<td>2.66; 27.98</td>
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<td>1371.8; 1723.8</td>
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<td>0.05; 0.35</td>
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</tbody>
</table>
Table 3 Sample size corrected Akaike information criterion (AICc) rankings of the models* that best described variation in $\log_{10}$ % lipid values (top), $\log_{10}$ caloric densities (cal·g$^{-1}$) (middle) and $\log_{10}$ THg concentrations (mg·kg$^{-1}$) (bottom) from anadromous Deception River Arctic charr. Only models within $\Delta_i = 2$ are included in the table as these were considered equivalent to the best model. Also given are the number of model fitted parameters ($k$), residual sums of squares (RSS), delta values ($\Delta_i$) defining model the difference between the estimated model AICc and the best model AICc, the Akaike weight ($w_i$) and evidence ratio (ER$_i$) for each model.

<table>
<thead>
<tr>
<th>Model</th>
<th>$k$</th>
<th>RSS</th>
<th>AICc</th>
<th>$\Delta_i$</th>
<th>$w_i$</th>
<th>ER$_i$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Variations in $\log_{10}$ % lipids</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Season, % C, % N</td>
<td>5</td>
<td>2.32</td>
<td>-38.78</td>
<td>0.00</td>
<td>0.26</td>
<td>1.00</td>
</tr>
<tr>
<td>Season, % N</td>
<td>4</td>
<td>2.42</td>
<td>-37.79</td>
<td>0.99</td>
<td>0.16</td>
<td>1.63</td>
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<tr>
<td>Variations in $\log_{10}$ caloric density</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Season, Age, Condition, % C, % N</td>
<td>7</td>
<td>0.01</td>
<td>-296.32</td>
<td>0.00</td>
<td>0.28</td>
<td>1.00</td>
</tr>
<tr>
<td>Season, Fork length, Condition, % C, % N</td>
<td>7</td>
<td>0.01</td>
<td>-296.20</td>
<td>0.12</td>
<td>0.26</td>
<td>1.06</td>
</tr>
<tr>
<td>Season, Total weight, Condition, % C, % N</td>
<td>7</td>
<td>0.01</td>
<td>-294.69</td>
<td>1.64</td>
<td>0.12</td>
<td>2.27</td>
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<tr>
<td>Variations in $\log_{10}$ THg</td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total weight, Condition, $\delta^{13}$C</td>
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<td>2.40</td>
<td>-78.34</td>
<td>0.00</td>
<td>0.27</td>
<td>1.00</td>
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<tr>
<td>Total weight, Condition, $\delta^{13}$C, $\delta^{15}$N</td>
<td>6</td>
<td>2.37</td>
<td>-77.73</td>
<td>0.61</td>
<td>0.20</td>
<td>1.36</td>
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<tr>
<td>Season, Total weight, Condition, $\delta^{13}$C</td>
<td>6</td>
<td>2.39</td>
<td>-76.49</td>
<td>1.85</td>
<td>0.11</td>
<td>2.52</td>
</tr>
</tbody>
</table>

* General linear models were estimated using the following candidate variables: season (summer 2016 and post-winter 2017), fork length (mm), total weight (g), age (years), somatic condition (K), $\delta^{13}$C (‰), % carbon (C), $\delta^{15}$N (‰), % nitrogen (N), and significant two-way interactions. Explanatory variables that were highly correlated, or known to have biological redundancy (e.g., fork length and total weight) were not included in the same model.
Figures

**Fig. 1** Map of the Deception Bay, the Deception River, and the two over-wintering lakes, Lake Duquet and LakeFrançois-Malherbe, from which Arctic charr were sampled for this study. Black circles represent mining operations present in the area, while the black square represents the sampling locations in relation to the province of Québec in eastern Canada.
Fig. 2 Seasonal variation in dorsal muscle $\log_{10}$ % lipid values, $\log_{10}$ caloric densities, and $\log_{10}$ THg concentrations from summer 2016 and post-winter 2017 sampled Deception River anadromous Arctic charr
Fig. 3 Relationships between dorsal muscle $\log_{10}$ % lipid values and biological variables (fork length, total weight, fish age, and somatic condition), as well as stable isotopes ($\delta^{13}C$, % carbon, $\delta^{15}N$, and % nitrogen)
Fig. 4 Relationships between collected dorsal muscle log$_{10}$ caloric densities and biological variables (fork length, total weight, fish age, and somatic condition), as well as stable isotopes ($\delta^{13}$C, % carbon, $\delta^{15}$N, and % nitrogen)
Fig. 5 Relationships between sampled dorsal muscle log$_{10}$ THg concentrations and biological variables (fork length, total weight, age, and somatic condition), stable isotopes ($\delta^{13}$C, % carbon, $\delta^{15}$N, and % nitrogen), and fish condition measures (log$_{10}$ % lipid values)