1	Working title: Behavioural tactics across thermal gradients align with partial						
2	morphological divergence in brook charr						
3	Author names: Aliénor Stahl <sup>1</sup> , Marc Pépino <sup>1,2</sup> , Andrea Bertolo <sup>1</sup> , Pierre Magnan <sup>1</sup>						
4	Affiliations:						
5	(1) Centre de Recherche sur les Interactions Bassins Versants – Écosystèmes						
6	Aquatiques (RIVE), Université du Québec à Trois-Rivières, Trois-Rivières, QC,						
7	Canada						
8	(2) Current address: Direction de la Gestion de la Faune Mauricie – Centre-du-						
9	Québec, Ministère de l'Environnement, de la Lutte Contre les Changements						
10	Climatiques, de la Faune et des Parcs, Trois-Rivières, QC, Canada						
11	Corresponding author: Aliénor Stahl, alienor.stahl@uqtr.ca						
12	Journal: Journal of Animal Ecology						
13	Keywords (up to 8): thermal stratification; behavioural thermoregulation; vertical forays;						
14	diel movement patterns; morphological divergence; acoustic telemetry; Salvelinus						

15 fontinalis

#### 16 Abstract (350w max):

17 Understanding how animals balance environmental constraints is essential for predicting 18 species persistence under climate change. In thermally stratified lakes, cold-water fishes 19 such as brook charr (Salvelinus fontinalis) must navigate vertical gradients in temperature 20 and oxygen to optimize foraging while avoiding physiological stress. We hypothesized that 21 individuals would exhibit behavioral tactics that reflect a trade-off between accessing 22 warm surface waters to exploit profitable prey and avoiding thermal stress, with greater 23 constraints expected on epilimnetic use as surface temperatures rise. Using high-24 resolution acoustic telemetry, we quantified fine-scale patterns of thermal habitat use, 25 vertical foray behavior, and diel timing in a wild brook charr population across the summer 26 stratification period. We also assessed whether behavioral thermoregulation aligned with 27 morphological differences, testing whether divergent behaviours reflect partial ecotypic divergence. As surface temperatures rose, brook charr reduced their use of the warm 28 29 epilimnion, making fewer, and shorter vertical forays. Hypolimnion use increased 30 concurrently but was unrelated to limiting oxygen concentrations, indicating that deep-31 water use was not physiologically constrained and may reflect an alternative foraging 32 behavior. Epilimnetic forays peaked at dusk and varied with moon phases, consistent with 33 crepuscular visual foraging. Hypolimnetic use peaked at dawn and dusk but showed no 34 response to moonlight. Diel patterns shifted seasonally: in warmer months, epilimnetic 35 access was restricted to twilight hours, while cooler months saw broader surface use 36 throughout the day. Principal component analysis of vertical movement and temperature exposure traits revealed two behavioral tactics: a "warm" tactic, characterized by 37 38 frequent epilimnetic forays and warmer average thermal exposure, and a "cool" tactic, 39 associated with greater hypolimnetic use and cooler average temperatures. These tactics

40 corresponded with partial morphological divergence within the two sexes. Linear discriminant analysis showed that males following the warm tactic were morphologically 41 42 distinct from cool ones, differing in traits related to feeding and swimming performance, 43 while females showed weaker morphological differentiation. Together our findings reveal 44 repeatable habitat use and consistent thermal tactics that reflect trade-offs between 45 foraging and thermal stress. Vertical gradients in temperature and resource distribution may thus promote fine-scale individual specialization and phenotypic divergence in cold-46 47 water species facing lake warming.

48

49 Acknowledgements: We thank the numerous students and research assistants involved 50 in this project for their invaluable field and laboratory assistance: Pierre-André Bordeleau, 51 Olivier Chouinard, Alexandre East, Antoine Filion, Chantal Fournier, Matteo Giacomazzo, 52 Natalie Godbout, Benjamin Gosselin, Georey Marselli, Vickie Lapointe, Riwan Leroux, Vincent Rainville, and Irene T. Roca. We also thank Olivier Roy and Éric Harnois from the 53 54 Mastigouche Wildlife Reserve for their logistical support. We are grateful to the VEMCO 55 staff, especially Stephanie Smedbol, Dale Webber, Jeremy Kuehner, and Richard Vallée, 56 for their constructive comments and judicious advice on the VEMCO Positioning System. 57 This work was supported by grants from Natural Sciences and Engineering Research 58 Council of Canada (NSERC) and the Canada Research Chair Program to PM. Aliénor Stahl was supported by funds from the NSERC to PM and a Postdoctoral fellowship from the 59 60 University du Québec à Trois-Rivières.

61 Author contributions (with statement on inclusion):

Data acquisition: MP, AB, PM. Conceptualization: AS, MP, AB, PM. Data curation: AS, PM.
Formal analysis: AS. Coding: AS, MP. Methodology: AS, MP, AB, PM. Visualization: AS.
Writing – original draft: AS. Writing – review and editing: AS, MP, AB, PM. All authors read
and approved the final manuscript. Our study brings together authors from a number of
different countries, including scientists based in the country where the study was carried
out. Whenever relevant, literature published by scientists from the region was cited;
efforts were made to consider relevant work published in the local language.

**Data availability statement:** Data and code will be made available on Borealis.

**Conflict of interest statement:** The authors declare there are no competing interests.

### 73 Introduction

74 Fish, as ectotherms, are strongly constrained by the thermal conditions of their 75 environment. Ambient water temperature regulates their distribution, activity, and 76 performance, with direct consequences for survival, growth, and reproduction 77 (Magnuson et al. 1979; Kieffer et al. 1998). When temperatures deviate from optimal 78 ranges, fish adjust their behavior -primarily through active habitat selection and 79 behavioral thermoregulation- to maintain physiological performance (Haesemeyer 2020; Amat-Trigo et al. 2023). However, their capacity to adapt to thermal stress is 80 81 asymmetrical. According to Brett's heat-invariant hypothesis, lower thermal limits may 82 shift with local adaptation, whereas upper limits remain relatively fixed -potentially 83 constraining persistence in warming environments (Brett 1971; Martin and Huey 2008; 84 Buckley et al. 2022; Ern et al. 2023).

85 Thermal stratification in lakes creates steep vertical temperature gradients that divide the 86 water column into distinct layers (i.e., epilimnion, metalimnion, and hypolimnion) which 87 strongly influence fish distribution and activity (Flood et al. 2021). Many species remain 88 within preferred temperature zones to optimize metabolic efficiency (Guzzo et al. 2019), 89 yet may periodically enter in suboptimal thermal environments to access resources or 90 avoid predators (Thomas et al. 2023). This trade-off is especially pronounced in cold-91 water species such as salmonids, which often ascend into warmer epilimnetic waters to 92 forage despite the associated physiological costs (Guzzo et al. 2017). These excursions elevate metabolic demands by increasing oxygen consumption and energy expenditure 93 94 through heat transfer from the surrounding water (Clarke and Johnston 1999). To mitigate 95 these costs, salmonids often employ behavioral tactics such as brief foraging bouts

96 followed by rapid returns to cooler refugia, a pattern documented across multiple species 97 in stratified systems (Bertolo et al. 2011; Armstrong et al. 2016). Although the general 98 relationship between fish behavior and thermal gradients is well established, fine-scale 99 patterns of vertical forays across thermal layers remain poorly understood in natural 100 settings. Most studies have focused on broad habitat use or diel vertical shifts (Goyer et 101 al. 2014; Cote et al. 2020), while few have quantified behavioral metrics such as foray 102 frequency, duration, and timing across thermal strata. In particular, the role of the 103 hypolimnion remains understudied (but see e.g. Roberts et al., 2012; Rodrigues et al., 104 2022), despite its increasing importance in stratified systems experiencing surface 105 warming and deep-water hypoxia (Bartosiewicz et al. 2019).

106 The physiological constraints of different thermal layers differ markedly. In the epilimnion, 107 temperature can reach lethal levels, strictly limiting how long fish can remain in surface 108 waters before incurring thermal stress (Pépino et al. 2015). In contrast, the hypolimnion 109 is often characterized by reduced thermal stress but also low oxygen availability 110 (hypolimnetic hypoxia), which can constrain fish movement and influence habitat 111 selection (Fang and Stefan 2009; Roberts et al. 2012; Rodrigues et al. 2022). Unlike acute 112 thermal stress, short term exposure to hypoxia exerts more gradual effects, impairing 113 aerobic performance without causing immediate physiological failure (Chapman and 114 Mckenzie 2009). Climate change is intensifying these constraints by warming surface 115 water and worsening hypolimnetic hypoxia through stronger thermal stratification and 116 diminished oxygen replenishment. This combination of stressors is of particularly 117 concern for salmonid species, which already occupy a narrow thermal window for 118 optimal growth and survival (McCullough et al. 2009). As warming and hypoxia intensify, 119 understanding how salmonid species navigate thermal gradients to balance

physiological stress and foraging needs is critical for predicting their future distributionand persistence under climate change.

122 Brook charr (Salvelinus fontinalis) provide an ideal model for studying how fish navigate 123 thermal constraints due to their reliance on both warm and cold-water habitats (Pépino 124 et al. 2024). First, the species colonized many postglacial lakes of the Canadian Shield 125 which exhibit pronounced vertical temperature gradients creating distinct thermal zones 126 that brook charr must navigate to balance thermal stress, oxygen availability, and foraging 127 needs (Smith et al. 2020). Second, brook charr display resource polymorphism in many 128 of these lakes, where a littoral ecotype feeds mainly on zoobenthos and a pelagic ecotype 129 feeds mostly on zooplankton implying habitat-specific foraging tactics (Bourke et al. 130 1997; Proulx and Magnan 2004; Pépino et al. 2018; Rainville et al. 2021a). However, no 131 direct relationship between morphology and behavior has yet been demonstrated. 132 Similar patterns of habitat-related resource polymorphism are well documented in other 133 Salvelinus species, particularly Arctic charr (S. alpinus), where morphs specialized for 134 deep, pelagic, and littoral zones show distinct morphological and behavioral adaptations 135 to their thermal and foraging environments (Jonsson 2001; Hawley et al. 2016). Behavioral 136 studies similarly show that brook charr use the epilimnion, metalimnion, and 137 hypolimnion at different times, demonstrating flexible habitat use (Pépino et al. 2024). 138 This flexibility makes them an excellent model for studying behavioral thermoregulation 139 in response to temperature gradients and resource availability.

This study focuses on fish behavioral thermoregulation by addressing key gaps in our understanding of fine-scale vertical movement in stratified lakes, pinpointing on how fish foraging behavior varies across thermal zones and shifts with changing summer

143 temperatures. Specifically, we quantify the temporal patterns of vertical forays in relation 144 to temperature and oxygen availability. Beyond individual metrics, we classified fish into 145 distinct thermoregulation behavioral tactics based on their use of thermal strata, 146 potentially shaped by asymmetrical physiological constraints. We then tested whether 147 these thermoregulation behaviors were associated with morphological differences, exploring whether thermal niches align with potential foraging morphotypes. We 148 149 hypothesize that: (1) as surface temperatures rise, brook charr will reduce forays into the 150 epilimnion due to thermal stress; (2) like Arctic charr (Jonsson 2001), some individuals 151 may increasingly use deeper zones such as the hypolimnion to access cooler 152 temperatures, but remain constrained by low oxygen availability; (3) epilimnetic forays 153 will peak at dusk, coinciding with brook charr foraging activity patterns (Bourke et al. 154 1996), while forays into the hypolimnion will show less predictable patterns due to 155 competing influences of temperature, food availability, and oxygen constraints; and (4) 156 thermoregulation behaviors will correspond to distinct morphotypes, suggesting 157 integrated thermal and trophic tactics. By combining fine-scale behavioral metrics, 158 thermal profiles, and morphological data, this study provides new insight into how a cold-159 water species manage competing environmental constraints and whether alternative 160 tactics are linked to phenotypic divergence.

161

# 162 Material and methods

# 163 Lake characteristics

The study was conducted in Lake Ledoux (46°38'N, 73°15'W), located in the Mastigouche
Wildlife Reserve, Québec, Canada, from 2017 to 2018. Lake Ledoux is a small,

166 oligotrophic lake typical of temperate regions, with a surface area of 11.9 ha, an average 167 depth of 5.5 m, and a maximum depth of 18.0 m. Its physicochemical properties are 168 consistent with those reported for similar lakes in the region (Magnan 1988; Brassard et 169 al. 2023). Aquatic vegetation is primarily concentrated in shallow waters (0-2 m depth; 170 Magnan unpubl.), corresponding to the littoral zone, which serves as a key foraging habitat 171 for brook charr due to the abundance of preferred prey species. Brook charr is the sole 172 fish species present in Lake Ledoux, where sport fishing is strictly regulated by the 173 Québec Government (Bourke et al. 1996; Pépino et al. 2024).

174 To monitor thermal conditions, temperature profiles were recorded from late June 2017 175 to early September 2018 using thermographs (iButtons DS1922L in 2017 and iBCod 176 DS1921G-F5 in 2018; Alpha Mac Inc.). Sensors were deployed at 0.5 m intervals from the 177 surface to depths exceeding 10 m, with thermograph lines anchored to a floating raft 178 positioned at the lake's deepest point. While the iBCod sensors are fully submersible, 179 iButtons were encapsulated in aquarium-grade silicone (SCS1200 clear sealant) to 180 protect them from prolonged water exposure. Temperature readings were collected at 4-181 hour intervals throughout the study. Lake Ledoux exhibited thermal stratification from 182 mid-May to end of October. Stratification characteristics were analyzed using the 183 'rLakeAnalyzer' package (Winslow et al. 2013) in R (R Core Team 2017) and linearly 184 interpolated per hour. To test sensitivity to thermocline position, we also applied a 185 conservative classification: the hypolimnion was defined as starting 1 m below its 186 thermocline, and the epilimnion as ending 1 m above it. This allowed us to assess whether 187 behavioral patterns were robust to small shifts in thermal structure estimates.

Dissolved oxygen concentrations were monitored in epilimnion, metalimnion and hypolimnion with four probes (miniDOT<sup>®</sup> Loggers, PME, USA) placed at 1,5, 3, 5 and 8 m depth. Oxygen was monitored every minute between July 4 and October 23 in 2018. The probes were retrieved weekly for cleaning and to check their status. Since a dense cover of water mites was observed for the two shallower probes after the first week of deployment, the probes were subsequently equipped with automatic whippers to avoid clogging.

# 195 Acoustic telemetry system

196 We employed an acoustic telemetry positioning system to continuously monitor brook 197 charr movements and habitat use with high spatiotemporal resolution (<1 minute, <5 m) 198 using a VEMCO Positioning System (VPS). Specifically, we deployed the HR2 VPS 199 technology, designed to minimize signal 'collisions'—a common issue when multiple 200 tagged individuals are in close proximity, leading to interference during signal 201 transmission. The system comprised 23 hydroacoustic receivers (HR2-180k-100; VEMCO 202 Inc.) and four reference tags (V9TP-2x-180k-34m; VEMCO Inc.), ensuring full lake 203 coverage. The HR2 VPS was operational from June 21, 2017, to November 6, 2018. 204 Additional details on the HR2 VPS systems, as well as performance assessments, are 205 provided in the supplementary information of Pépino et al. (2024).

206 Fish capture and tagging

Brook charr were captured using Alaska traps (1.0×1.8 m mouth opening, two 1×15 m wings, 1.27 cm mesh size; Fipec Industries) over two consecutive years (12–20 June 2017 and 5–7 June 2018). Following capture, fish were temporarily held in an enclosure (3×4×6 m depth) and surgically tagged on the same day. Tagging followed standard

211 protocols (Adams and Maitland 1998; Bridger and Booth 2003), previously validated for 212 this species and study system (Goyer et al. 2014; Pépino et al. 2015, 2024). In 2017 213 (n = 30) and 2018 (n = 15), brook charr (mass: 226–600 g; total length TL: 291–391 mm) 214 were equipped with 4.0 g acoustic transmitters (V9TP-2x-180k-34m; VEMCO Inc.), which 215 alternately recorded internal temperature and fish depth. Transmitter weight never 216 exceeded 2% of fish body mass, complying with best-practice guidelines to minimize 217 behavioral and physiological impacts (Mellas and Haynes 1985; Bridger and Booth 2003). 218 All individuals were released in apparent good condition and exhibited normal post-219 release behavior. Full details are provided in the Supporting Information of Pépino et al. 220 (2024).

# 221 Data preparation and filtering

222 Detection data were filtered to retain only periods of thermal stratification: June 24 to 223 October 31 in 2017 and May 19 to September 8 in 2018 (Figure S1). This ensured that 224 analyses targeted behavior relevant to stratified lake conditions, when distinct thermal 225 habitats (epi-, meta- and hypolimnion) were clearly defined. Detections outside the lake 226 boundaries and those from individuals inferred to be dead (i.e., stationary for >24h) were 227 excluded. To reduce artifacts from post-tagging recovery, we excluded the first 10 days 228 following tagging, as fish activity was observed to stabilize thereafter.

229 Missing fish depth or temperature values were linearly interpolated within a one-minute 230 window, independently for each variable. This preserved the temporal resolution 231 necessary for accurate derivation of vertical movements and foray events for testing our 232 hypothesis that individuals engage in short, thermally constrained excursions across 233 depth layers.

#### 234 Thermal habitat use and vertical forays

235 To characterize brook charr thermal behavior and habitat use across depth strata, we 236 derived several variables from the telemetry data. These included: (1) fish depth, 237 combining sensor and interpolated values; (2) distance from the lake bottom, calculated 238 as the difference between lake bathymetric depth and fish depth at each detection point; 239 (3) distance to the nearest shoreline, derived from a shapefile based on bathymetric 240 mapping; and (4) swimming speed in body lengths per second (TL.s<sup>-1</sup>), using consecutive 241 positional data and each fish's total length (TL) recorded at tagging. These metrics 242 provided a foundation for describing individual movement and habitat use across thermal 243 and spatial gradients.

244 To assess how fish used distinct thermal habitats, we focused on vertical movements into 245 and out of the epilimnion and hypolimnion (thermal layers with different temperature and 246 oxygen characteristics, and thus likely to impose different ecological trade-offs). We 247 defined a "foray" as a movement into one of these layers followed by a return to the 248 metalimnion. A foray began at the first detection within the target layer (e.g., epilimnion) 249 and ended at the first subsequent detection in a different layer. Thermal boundaries were 250 determined using hourly thermocline depths (see Lake characteristics section), ensuring 251 that foray detection accounted for shifting thermal structure.

For each individual, we quantified multiple aspects of foraying behavior: (1) total number of forays, (2) duration of each foray (from entry to exit), and daily summary statistics including: (3) mean foray duration, (4) total daily time spent in the target layer, and (5) maximum duration of a single foray per day. These metrics allowed us to investigate fine-

scale behavioral tactics and potential ecological roles of vertical forays, particularlyunder thermal constraints.

### 258 Classifying thermal behavior and habitat use tactics

259 To identify individual-level behavioral thermoregulation and assess whether brook charr 260 segregated into consistent thermal ecotypes (as previously reported in Lake Ledoux by 261 Bertolo et al., 2011 and Goyer et al., 2014), we calculated summary behavioral metrics 262 across the entire study period. These variables included: (1) mean body temperature; (2) 263 mean foray duration into the epilimnion and hypolimnion; (3) proportion of time spent in 264 the epilimnion and hypolimnion; (4) mean swimming speed, both overall and within each 265 layer; (5) mean distance to shore; and (6) mean distance to the lake bottom. These 266 metrics were selected to reflect habitat affinity, foraging behaviors, and thermal exposure.

267 To assess whether brook charr exhibited distinct thermal tactics based on behavior alone, 268 we performed a principal component analysis (PCA, prcomp function in R) on metrics of 269 vertical movement, temperature exposure, and space use to identify dominant axes of 270 behavioral variation and reduce dimensionality. All variables were standardized prior to 271 PCA. The first principal component (PC1) was used to classify individuals into two groups 272 based on thermal habitat use: individuals with negative PC1 scores were classified as 273 adopting a "warm" tactic, while those with positive scores followed a "cool" tactic (see 274 details in Results section). To assess the consistency of this classification under peak 275 thermal stress, we repeated the PCA using data from July and August—the warmest 276 period of the summer. This seasonal comparison allowed us to evaluate whether 277 individuals consistently expressed divergent thermal tactics, as would be expected if 278 behavioral divergence is shaped by trade-offs between foraging opportunities and

thermal or oxygen stress. These classifications served as the basis for subsequentanalyses of diel activity patterns and morphology.

281 To further contextualize habitat use, we assigned each detection to a vertical habitat 282 category: benthic (within 0-2 meters from the lake bottom) or pelagic (beyond this 283 threshold). These were combined with vertical layers (epilimnion, metalimnion, and 284 hypolimnion) to define composite spatial-thermal habitat types (e.g., epilimnetic-285 pelagic). For each fish, we calculated the proportion of detections in each composite 286 category. This analysis was designed to test whether brook charr occupied distinct three-287 dimensional niches, potentially consistent with ecotypic specialization in foraging tactic and habitat preference. It also allowed us to assess whether habitat use patterns aligned 288 289 with the thermal tactics identified earlier. Having classified these tactics, we next 290 evaluated whether these behavioral types differed in the timing and structure of vertical 291 movements.

### 292 Diel patterns in foray timing

To test whether vertical forays were temporally structured, particularly with respect to diel cycles, we modeled the likelihood of foray initiation throughout the 24-hours period. This analysis was motivated by previous studies suggesting crepuscular peaks in brook charr foraging activity (Bourke et al. 1996). We hypothesized that forays into the epilimnion, potentially exposing fish to warmer temperatures, would be concentrated around dusk to optimize feeding benefits due to access to emerging insects.

We used hierarchical generalized additive models (HGAMs) with cyclic cubic regression splines to model diel variation in foray probability. Following Jannarilli et al. (2024), we split each day into 1-hour intervals and defined binary indicators (StartForay<sub>i,t,i</sub>) for

302 whether fish *i* initiated at least one foray into the target thermal layer during time interval 303 *t* on day *j*. This was done separately for forays into the epilimnion and hypolimnion. To 304 reduce model complexity, we aggregated these indicators across all days per fish and 305 time interval, modeling the counts as binomial outcomes (successes vs. failures).

306 Covariates included sunrise and sunset times, calculated using the bioRad R package 307 (Dokter et al. 2019), and lunar illumination, derived from the *lunar* R package (Lazaridis 308 2022). Each model featured: (1) a fish-specific cyclic smoother for time of day, allowing 309 for individualized diel activity curves, (2) a global cyclic smoother for time of day, which 310 acted as a prior to constrain individual-specific smoothers and capture the overall diel 311 trend across fish, and (3) a random effect for individual fish to account for repeated 312 measures. Models were fitted using the bam function from the mgcv package (Wood 313 2017), and predictions with 95% confidence intervals were obtained using *predict.bam*.

We first constructed global models over the full study period for forays into the epilimnion and hypolimnion separately. To assess how diel patterns shifted with seasonal thermal regimes, we then fit monthly models from June through October. These models excluded covariates to focus solely on diel variation, minimizing overfitting and enabling cleaner comparisons of temporal activity shifts across months.

We also analysed hourly variation in time allocation across thermal layers per month. For each hour, we calculated the proportion of time each fish spent in a given layer (TimeForay<sub>i,t,j</sub>), defined as minutes in the layer divided by 60. These values were modeled using Gaussian HGAMs with a similar smoothing structure. Together, these analyses allowed us to evaluate whether diel behavior aligned with predictions of thermally constrained foraging.

# 325 Morphological differentiation of thermal ecotypes

326 To investigate whether thermal ecotypes (classified as "warm" or "cool" tactics based on 327 PCA-derived depth-use patterns; see Classifying thermal behavior and habitat use tactics 328 for more details) exhibited corresponding morphological divergence, we conducted a 329 morphometric analysis using both geometric landmarks and measured traits. 330 Standardized photographs were taken of each fish, and 25 landmarks were placed using 331 tpsDig2 (Rohlf, 2012; Figure S2). Following established procedures (Arbour et al. 2011; 332 Dermond et al. 2019), curvature correction was applied using four lateral-line landmarks. 333 Curvature was minimal and showed no systematic differences between ecotypes. 334 Landmark files were generated with tpsUtil (Rohlf 2012b).

335 To evaluate morphological differentiation, we performed a linear discriminant analysis 336 (LDA, using the MASS R package (Venables and Ripley 2002)) and used a forward 337 selection process to identify the best model (i.e., highest overall classification accuracy). 338 We retained 11 traits: body height, dorsal fin base length, eye width, head height, head 339 length, mouth width, pectoral fin length, pelvic fin height, peduncle height, snout-to-eye 340 distance, and lower jaw length. The LDA grouping factor was a composite of thermal tactic 341 and sex, allowing us to assess interactions between thermal habitat use and sexual 342 dimorphism. Assumptions of homogeneity of within-group covariances were verified with 343 permutation tests. Classification accuracy was assessed via leave-one-out cross-344 validation. Finally, we computed canonical correlation coefficients to assess the 345 contribution of each trait to group separation, following Zuur et al. (2009). This helped 346 identify which morphological traits best explained ecotype divergence and informed 347 potential links between form and ecological function.

348 Research permits

The research study was approved by the Animal Care Committee of the Université du Québec à Trois-Rivières (Comité de Bons Soins aux Animaux – CBSA; permit numbers: 2016-P.M.42) and by the Ministère des Forêt, Faunes et Parcs (MFFP; SEG permit numbers: 2016-06-21-080-04-S-P; 2017-04-27-051-04-S-P; 2018-05-21-050-04-S-P).

353

# 354 Results

We assessed fine-scale patterns of brook charr thermal habitat use, vertical foraying behavior, diel movement timing, and morphological divergence across behavioral types. Our results are structured to correspond to the major research questions and analytical approaches detailed in the Methods, beginning with an examination of how fish used depth and temperature strata during stratified periods.

360 Thermal habitat use and vertical forays

361 As surface temperatures increased over the summer months, brook charr showed shifts 362 in vertical foraying behavior across thermal layers (Figure 1). Consistent with 363 expectations, individuals reduced their use of the epilimnion under warmer surface 364 conditions. Specifically, the number of epilimnetic daily forays (Z = -26.35, p < 0.001), the 365 total daily duration spent in the epilimnion ( $F_{1,3037}$  = 1286,  $R^2$  = 0.29, p < 0.001), and the 366 maximum daily foray duration ( $F_{1,3037}$  = 378.7,  $R^2$  = 0.11, p < 0.001) into this layer all 367 declined significantly with increasing epilimnion temperature (Figure 1-A). In 2017, the 368 average number of daily forays dropped from over 50 in June to 21 in August; the decline 369 was even steeper in 2018, with a drop from 67 in June to just 10 in August. Similarly, total time spent in the epilimnion decreased from more than 2 hours and 30 minutes in June 2017 to under 1 hour and 30 minutes in August. In 2018, this decline was even more pronounced, from over 3 hours and 30 minutes in June to less than 1 hour in August. In contrast, the mean duration of individual forays showed a weaker relationship with temperature ( $F_{1,3037} = 112.9$ ,  $R^2 = 0.03$ , p < 0.001), varying by less than 2 minutes across months within a year and by approximately 3 minutes across years.



**Figure 1:** Metrics describing brook charr vertical habitat use as a function of day of year and average daily epilimnion temperature. Panels show summary statistics for forays into the (A) epilimnion and (B) hypolimnion, including: number of forays per day (defined as a transition into the target layer following a detection in a different layer), total daily duration of forays (in hours), maximum foray duration (in minutes), and mean foray duration (in minutes). Data are shown separately for the summers of 2017 and 2018. Colour scale reflects average daily epilimnion temperature (yellow = warm, dark purple = cold). Solid

lines represent the 50th percentile (median), and dotted lines the 95th percentile for each metric. Line colour corresponds to the thermal layer: red for epilimnion, blue for hypolimnion. See Appendix for (i) daily depth estimates of thermal layers (Figure S1); (ii) an alternative version using a more conservative threshold of  $\geq$ 1 m from the metalimnion boundary (Figure S3); (iii) a version of panel A for hypolimnion forays in 2018 coloured by average daily dissolved oxygen (Figure S4); and (iv) per-fish histograms of epilimnion foray durations (Figure S5).

391

392 In the hypolimnion, the number of forays (Z = 24.44, p < 0.001) and total time spent 393  $(F_{1,2143} = 28.67, R^2 = 0.01, p < 0.001)$  increased as epilimnion temperature rose (Figure 1B). 394 In 2017, the number of hypolimnetic forays increased from 28 in June to 46 in August, and 395 total time increased from just under 2 hours in June to over 3 hours and 15 minutes in July. 396 In 2018, forays rose from 11 in June to 46 in August, with total time increasing from just 397 over 1 hour to around 2 hours. However, both maximum daily foray duration ( $F_{1,2143}$  = 5.621, 398  $R^2 = 0.002$ , p = 0.017) and mean foray duration ( $F_{1,2143} = 0.6635$ , p = 0.41) showed weak or 399 no clear trends with temperature, varying by about 6 minutes across months and by less 400 than 1 minute on average. These patterns were consistent when using both the default 401 and conservative definitions of thermal layer boundaries (Figure S3). Oxygen availability 402 in the hypolimnion (mean = 4.9 mg/L, SD = 1.4) was not significantly related to the number 403  $(F_{1,1026} = 0.4172, p = 0.5185)$  or duration  $(F_{1,644} = 4.895, p = 0.02)$  of forays. For example, the 404 number of hypolimnetic forays remained around 35 in both July and September, despite 405 oxygen concentrations being highest in July and lowest in September. Similarly, total daily 406 foray time declined from 23 minutes in July to 15 minutes in September (Figure S4).

407 Use of habitat categories (benthic versus pelagic) showed some seasonal structure, 408 though not a consistent relationship with epilimnion temperature across years (Figure 2). 409 In early summer 2018, for instance, individuals transitioned from using epilimnetic 410 benthic habitats to greater use of benthic areas in the metalimnion, which remained the 411 most frequently used habitat across the season. In contrast, increased use of epilimnetic 412 benthic zones in fall 2017 may reflect reproductive activity rather than foraging, given the 413 known timing of brook charr spawning. This interpretation is supported by patterns in 414 Figure 1, where forays into the epilimnion become longer during the same period— 415 suggesting that reproductive forays may leave a distinct behavioral signature compared 416 to shorter, more frequent foraging events. Across all foray metrics, we observed 417 substantial inter-individual variability, with wide spreads in both the median and upper 418 quantiles (Figures 1, S3, S4, and S5). This suggests that, although there were consistent 419 population-level responses to temperature, individuals differed markedly in the extent 420 and timing of their foray behavior, as well as in their use of habitat categories.



422 Figure 2: Proportion of brook charr detections in (A) benthic and (B) pelagic zones within 423 each thermal layer (epilimnion, metalimnion, hypolimnion), across day of year and 424 average daily epilimnion temperature. The benthic zone is defined as detections occurring 425 within 2 meters of the lake bottom. Summers 2017 and 2018 are shown in separate facets. 426 Colour scale reflects average daily epilimnion temperature (yellow = warm, dark purple = 427 cold). Solid lines show the 50th percentile (median), and dotted lines the 95th percentile 428 for each metric. Line colours indicate thermal layer: red (epilimnion), green 429 (metalimnion), and blue (hypolimnion). See Appendix for daily depth estimates of thermal 430 layers, Figure S1.

431

# 432 Classifying thermal behavior and habitat use tactics

433 The first two PCA axes captured the major variations (i.e., 51.6 %) of individual behavior 434 (Figure 3). PC1, which explained the most variance (30.6%), represented a gradient in 435 vertical thermal habitat use: individuals with negative scores spent more time in warmer 436 surface waters, made prolonged forays into the epilimnion, and exhibited higher 437 swimming speed while those with positive scores used cooler, deeper waters and the 438 hypolimnion more extensively. PC2 captured variation in habitat use at intermediate 439 depths, with individuals differing in benthic versus pelagic use within the metalimnion. 440 No clear separation in benthic/pelagic use emerged within the epilimnion or hypolimnion 441 (Figure S6).



442

443 Figure 3: Variable contributions to the first two axes of a Principal Component Analysis 444 (PCA) describing brook charr vertical behaviour and habitat use. The PCA was based on 445 standardized, per-fish values for: mean body temperature (°C; Temp), proportion of time 446 spent in the epilimnion and hypolimnion (0–1; Time.Epi and Time.Hypo), average 447 swimming speed across all layers (TL/s; Speed), average speed in the epilimnion and 448 hypolimnion (Speed.Epi, Speed.Hypo), mean distance from the lake bottom (m; 449 Distance.Bottom), mean distance to shore (m; Distance.Shore), and mean foray durations 450 into the epilimnion and hypolimnion (minutes; Duration.Foray.Epi, Duration.Foray.Hypo).

Supplementary variables (not used in the PCA) represent the proportion of detections in combinations of habitat type and thermal layer (dark red). Habitat is classified as benthic (within 2 m of the lake bottom; B) or pelagic (P), combined with depth zone: Epi = epilimnion, Meta = metalimnion, Hypo = hypolimnion (e.g., B-Epi = benthic epilimnion, P-Hypo = pelagic hypolimnion). These values sum to 1 for each fish. See Appendix for a version of this figure restricted to July–August to assess seasonal variation in PCA structure (Figure S7).

458

These patterns remained consistent when the analysis was restricted to the warmer summer months (July–August; Figure S7). We used PC1 to assign individuals to two behavioral groups: a "warm" tactic (negative PC1 scores), characterized by warmer thermal exposure (mean fish temperature:  $12.8 \pm 1.5$  °C), more epilimnetic forays and higher swimming speeds; and a "cool" tactic (positive PC1 scores), characterized by cooler ambient temperatures (mean:  $10.2 \pm 1.7$  °C), greater hypolimnetic use, and a greater average distance from shore.

# 466 Diel patterns in foray timing

In the general model spanning the full study period (Figure 4), epilimnetic use peaked at dusk, as predicted. "Warm" individuals were nearly twice as likely to access the epilimnion compared to "cool" individuals, and the probability of epilimnetic forays increased with lunar illumination, suggesting a role for visual foraging. In contrast, hypolimnetic use showed no response to lunar illumination intensity. Unexpectedly, activity in the hypolimnion clearly peaked at both dawn and dusk, with "cool" individuals

exhibiting higher overall probabilities of use than "warm" individuals. Notably, "warm" fish
were more likely to access the hypolimnion at dawn than at dusk.



Figure 4: Predicted diel vertical movement patterns of brook charr grouped into "warm" and "cool" behavioural categories based on their position along the first principal component axis (PC1) of the PCA on thermal behaviour. Panels show the predicted probability (range: 0–1) of transitioning into the epilimnion (red) or hypolimnion (blue) over a 24-hour cycle, estimated from separate hierarchical generalized additive models

(HGAMs) fitted to each behavioural group. Black vertical lines indicate sunrise and sunset times, which were included as covariates in the models; lunar illumination (percentage of moonlight) was also included. Predictions are shown for an "average" day, defined by median sunrise and sunset times across the study period. See Appendix for: (i) predictions from an HGAM fitted to a representative individual (Figure S8); and (ii) monthly models fitted separately for each behavioural category (Figure S9).

487

Individual variation in diel foray timing and intensity was substantial (Figure S8), with patterns varying among behavioral tactics. "Cool" individuals tended to concentrate their forays into deeper layers and exhibited broad diel distributions, while "warm" individuals showed more consistent crepuscular activity. This suggests that diel behavior is not uniform across the population but reflects underlying differences in thermal tactics or ecotypes.

494 Although monthly models revealed some seasonal plasticity, individual-level patterns 495 were more striking. During the warmest months (July-August), epilimnetic forays 496 occurred primarily around dusk, likely reflecting convergence of favorable conditions for 497 visual foraging -sufficient light to locate prey but cooler temperatures that reduce thermal 498 stress- rather than strictly a thermoregulatory response (Figures S9). In cooler months, 499 when surface temperatures declined seasonally and oxygen remained sufficient, these 500 temporal constraints relaxed. Fish accessed surface waters more continuously 501 throughout the day, possibly reflecting reduced thermal stress or engagement in non-502 foraging behavior such as reproductive activity. However, it is also plausible that dusk

peaks in warmer months primarily reflect increased prey availability, such as insect
emergence, rather than temperature-driven constraints alone.

#### 505 Morphological differentiation of thermal ecotypes

The assumption of multivariate homogeneity of within-group covariance was met (permutation test; n = 999,  $F_{3,37}$  = 1.838, p = 0.16), supporting the validity of the LDA. Wilks'  $\lambda$  statistic indicated a significant overall group effect ( $\lambda$  = 0.07, p < 0.001), and the first two linear discriminant (LD) axes accounted for over 90% of the total morphological variation (Figure 5-A).



**Figure 5:** Morphological differentiation in brook charr on linear discriminant analysis (LDA). (A) Individual discriminant scores (small symbols) and group centroids (large symbols) on the first two LDA axes. Fish are grouped by Classification × Sex, where classification refers to thermal type based on PCA results. orange = warm" fish; teal = "cool" fish; circles = females; squares = males. Ellipses encompass 90% of individuals per group. (B) Canonical correlation coefficients showing the contribution of each morphological trait to the LDA axes. Traits include: BH = body height, DB = dorsal fin base

length, EW = eye width, HH = head height, HL = head length, MW = mouth width, PC =
pectoral fin length, PH = pelvic fin height, PS = peduncle height, SN = snout-to-eye
distance, and SJ = lower jaw length (see Supplementary Figure S2 for a schematic of trait
locations).

523

524 LD1 explained 58.8% of the variation and primarily discriminated individuals by sex 525 (Figure 5-A). This axis was strongly and positively associated with traits related to overall 526 body and head morphology, including body height (BH), dorsal fin base (DB), pectoral fin 527 length (PC), eye width (EW), head length (HL), head height (HH), and snout-to-eye 528 distance (SE; Figure 5-B). LD2 accounted for 33.6% of the variation and partially 529 discriminated individuals by thermal tactics. It was most strongly associated with eye 530 width (EW), pelvic fin length (PH), jaw length (SJ), peduncle height (PS), body height (BH), 531 and mouth width (MW).

532 Males following the "warm" tactic were morphologically distinct from "cool" ones, with 533 no overlap along LD2 (Figure 5-A and Table 1). In contrast, females showed substantial 534 overlap between tactics and overlap also occurred between males and females of the 535 same tactic. Overall, "warm" tactic individuals tended to have larger mouths, longer 536 pelvic fins, and larger eyes, while "cool" tactic individuals had smaller mouths and pelvic 537 fins, and lower body height (Figure 6).



538

Figure 6: Mean body shapes of brook charr grouped by thermal tactic and sex. Average body shapes are shown for each classification group defined by thermal tactic (warm = orange, cool = teal) and sex. Solid coloured lines represent the mean shape of each group, while the dashed grey line indicates the grand mean shape across all individuals. Shape differences have been exaggerated by a factor of 5 to enhance visual interpretability. Representative photographs of individual fish are included alongside the outlines to illustrate morphological differences more intuitively.

546 **Table 1:** Jackknife-based classification accuracy (%) of brook charr assigned to four 547 groups based on thermal tactic ("cold" or "warm") and sex (female or male), using linear 548 discriminant analysis of morphological traits. Classification accuracy is reported for each 549 group separately (Cold-F, Cold-M, Warm-F, Warm-M) as well as the overall classification 550 rate. Expected accuracy from random classification is 25%.

	Predicte	_			
Observed	Cool-F	Cool-M	Warm-F	Warm-M	Proportion (%)
Cool-F	8	0	2	1	72.7

Cool-M	3	5	0	0	62.5
Warm-F	3	0	9	1	69.2
Warm-M	0	0	2	7	77.7

551

552 Cross-validation using a jackknife (leave-one-out) procedure yielded an overall 553 classification accuracy of 70%, compared to 25% expected by chance (Table 1). 554 Classification accuracy varied across groups, ranging from 62.5% in cold-tactic males to 555 77.7% in warm-tactic males. Misclassifications were more frequent in females, 556 particularly in distinguishing between warm and cold tactics, while males were more 557 likely to be misclassified as females of the same tactic.

558

# 559 Discussion

560 We investigated how brook charr navigate fine-scale trade-offs among temperature and 561 foraging opportunities in a thermally stratified lake. We hypothesized that individuals 562 would exhibit distinct vertical movement patterns shaped by asymmetrical thermal 563 constraints (e.g. Pépino et al., 2015) - with surface warming more strongly limiting access 564 to the epilimnion than low oxygen constrains use of the hypolimnion, consistent with 565 Brett's heat-invariant hypothesis- as well as by spatial or temporal variation in prey 566 availability (Leroux et al. 2023, 2025). We also predicted that these behaviors would be 567 associated with morphological traits. Using high-resolution telemetry, we identified two 568 consistent thermal behavioral tactics (i.e., "warm" and "cool") characterized by 569 contrasting habitat use, movement timing, and morphology. These findings are consistent 570 with our initial hypothesis that thermal constraints are asymmetrical along the thermal 571 gradient. Notably, they also reveal persistent and substantial use of the hypolimnion 572 throughout the stratified season, a pattern rarely documented in brook charr. However, 573 across both tactics, benthic zones—particularly in the metalimnion—remained the most 574 frequently used habitat type (Figure S6), highlighting the central role of benthic prey 575 resources in structuring behavior. This novel result suggests that cold, deep habitats may 576 support overlooked foraging opportunities and could play a broader ecological role in the 577 thermal behavior of cold-water species facing lake warming. Such fine-scale ecological 578 specialization within a single population underscores how intraspecific variation 579 facilitates niche partitioning and coexistence in heterogeneous freshwater environments, 580 and may enhance population resilience under environmental change.

# 581 Thermal habitat use and vertical forays

582 Brook charr adjusted their vertical movements in response to seasonal and diel changes 583 in thermal structure. During warmer months (July–August), the use of the epilimnion was 584 largely confined to crepuscular periods, particularly dusk. This pattern is likely driven not 585 by short term thermal fluctuations within the epilimnion but by prey availability, as dusk 586 corresponds to peak activity of benthic invertebrates such as chironomid pupae 587 (Grossman et al. 1980; Bourke et al. 1996) and gammarids (Grossman et al. 1980), as well 588 as zooplankton vertical migration in this system (Naud and Magnan 1988). As surface 589 waters cooled in fall, diel epilimnetic use broadened, and hypolimnion occupancy 590 increased, highlighting the importance of stable thermal refuges at depth. It is important 591 to note that these shifts may also reflect reproductive activity, particularly in October 592 (Bourke et al. 1997).

593 These patterns are consistent with studies showing that salmonids often trade off thermal
594 exposure and foraging opportunity under heterogeneous thermal environments (Brewitt

595 and Danner 2014; Armstrong et al. 2016; Rainville et al. 2022; Pépino et al. 2024). They 596 also echo findings from acoustic telemetry studies in other cold-water species such as 597 lake trout, where vertical movements reflect fine-scale behavioral thermoregulation in 598 response to both prey and habitat constraints (Guzzo et al. 2017). Hypolimnion use 599 remained stable across oxygen conditions, with values occasionally falling below 5 mg/L, 600 a commonly cited threshold for salmonid performance (Magnuson et al. 1979). This 601 suggests that either oxygen levels remained within tolerable bounds or that prey 602 availability at depth outweighed potential physiological costs—a possibility supported by 603 studies showing that fish may tolerate mild hypoxia to exploit profitable resources (Rahel 604 and Nutzman 1994; Railsback and Harvey 2002). Overall, these results highlight brook 605 charr's dynamic balancing of thermal and trophic constraints, as reflected in the fine-606 scale timing and depth of forays.

### 607 Classifying thermal behavior and habitat use tactics

608 Multivariate classification of movement and thermal exposure revealed two major behavioral tactics: "warm" individuals, which made more frequent and sustained 609 excursions into warm surface waters, and "cool" individuals, which remained 610 611 predominantly in cooler depths and used the hypolimnion more consistently. These 612 tactics were evident both across the full summer period and when analyses were 613 restricted to the warmest months (July-August), suggesting stable behavioral 614 specialization rather than transient flexibility. Importantly, while reproductive activity likely influenced behavior in the fall-particularly increased use of epilimnetic benthic 615 616 habitats-this did not alter the broader classification of individuals into warm and cool 617 thermal tactics, which remained consistent across the stratified period.

618 This behavioral classification aligns well with previous work in Lake Ledoux, which 619 identified a gradient of foraging tactics based on diel activity and space use (Bertolo et al. 620 2011; Goyer et al. 2014) and where individuals were similarly categorized into "warm" and 621 "cool" types. Our findings thus reinforce and extend this classification by linking it directly 622 to thermal habitat use and individual morphology. Although we did not assign individuals 623 to morphotypes based on capture zones (e.g., littoral vs. pelagic), the thermal tactics we 624 observed map onto similar ecological gradients described in other polymorphic 625 salmonids (Zimmerman et al. 2007; Chavarie et al. 2013). While two dominant tactics 626 emerged, we also observed considerable individual variation in metalimnion use and 627 foray behavior. This continuum of thermal responses supports the idea that behavioral 628 variation in this system is not strictly categorical but instead spans a gradient of individual 629 specialization.

630 The "cool" individuals' use of deeper, pelagic-associated zones suggests a foraging tactic 631 targeting prey in or near the hypolimnion. Stable isotope evidence supports this 632 interpretation, showing that brook charr feeding in deeper waters of similar systems have 633 distinct dietary signatures (Glaz et al. 2012) These findings point to niche partitioning 634 along thermal, spatial, and dietary axes that appear continuous rather than discrete. This 635 also aligns with observations by Rainville et al. (2021b), who found that while some 636 individuals specialize on either littoral or pelagic resources, many exploit both across 637 longer time frames -highlighting dietary flexibility and opportunism. Such plasticity may 638 be driven by temporal fluctuations in resource availability, such as seasonal mayfly 639 emergence or the late-summer decline in benthic invertebrates, and suggests that 640 individual specialization may not map cleanly onto morphotypes (Power et al. 2005; 641 Amat-Trigo et al. 2023).

642 We also observed a gradient in the use of the metalimnion, with some individuals favoring 643 benthic habitats and others using pelagic areas. This pattern could explain previously 644 observed muscle carotenoid signals (Rainville et al. 2021a) in presumed "cool" 645 individuals, which suggest a non-negligible contribution of zooplankton to their diet. This 646 horizontal structuring indicates that brook charr may partition the intermediate depth 647 zone not only vertically but also horizontally, expanding the range of accessible resources 648 and potentially reducing intraspecific competition. Benthic areas in the metalimnion 649 likely support prey such as insect larvae or emerging aquatic insects, while pelagic zones 650 may provide access to zooplankton and midwater invertebrates (Knudsen et al. 2006; 651 Eloranta et al. 2011). Similar fine-scale horizontal partitioning has been documented in 652 polymorphic lake trout and Arctic charr, where individual foraging tactics are associated 653 with differential use of benthic versus pelagic microhabitats (Knudsen et al. 2010; 654 Chavarie et al. 2016). This multi-dimensional niche use-across depth, temperature, and 655 horizontal structure—highlights the potential for fine-scale individual specialization 656 throughout the water column.

657 While our primary focus was on thermal and vertical habitat use, spatial variation was 658 implicitly captured through depth-layer usage as well as fish positions relative to 659 shoreline and lake bottom. Qualitative patterns in space use suggest that spatial habitat 660 partitioning may reinforce thermal niche differentiation. For example, "cool" individuals 661 tended to avoid the shallowest areas of the lake, especially during the warmer months (June to September), whereas "warm" individuals used these areas more frequently. 662 663 Beside this seasonal pattern, horizontal distributions appeared broadly similar across 664 tactics. To illustrate these trends, we include two supplementary figures (Figs. S10–S11) 665 showing the spatial distribution of detections by month, layer, and thermal tactic. These

666 maps reveal subtle spatial segregation but also considerable overlap—patterns that may 667 be shaped by underlying individual variation, since maps were not stratified at the 668 individual level. A more detailed spatial analysis incorporating bathymetry, habitat 669 structure, and individual movement trajectories was beyond the scope of this study, but 670 represents a valuable direction for future work.

671 Diel patterns in foray timing

672 Both "warm" and "cool" individuals increased use of the hypolimnion during crepuscular 673 periods, at both dawn and dusk. These timing patterns suggest targeted foraging behavior, 674 though not necessarily linked to pelagic prey. Unlike epilimnetic forays, hypolimnetic 675 excursions did not vary with lunar phase and did not coincide with known vertical 676 movements of zooplankton in Lake Ledoux, which are primarily concentrated in the upper 677 layers (Leroux et al. 2023, 2025). Instead, the timing and depth of these forays suggest a 678 focus on benthic prey targets. Chironomid larvae, which ascend toward emergence, are 679 likely candidates due to their predictable vertical movements at dawn and dusk and their 680 accessibility near the sediment surface (Learner et al. 1990), and may confer a 681 competitive advantage to fish capable of intercepting them early in this process. Another 682 likely prey item is the large-bodied burrowing mayfly Ephemera spp., observed in 683 abundance at the surface of Lake Ledoux during summer (Magnan and Pépino, pers. 684 obs.), which emerge from the sediment during crepuscular periods and may be captured 685 prior to surfacing (Whelan 1980; Bourke et al. 1996). These preys are abundant in low-686 light, low-oxygen habitats and exhibit diel movement patterns that align with the timing of 687 charr forays, though uncertainty remains about the precise prey targets and sensory cues 688 involved. Whereas direct feeding observations would further clarify these associations,

689 the more frequent hypolimnetic forays of "cool" individuals suggest a degree of 690 specialization for exploiting these resources. This supports the interpretation that 691 hypolimnion use reflects not just thermal avoidance, but a foraging tactic that balances 692 energetic gain against the potential costs associated with low oxygen availability 693 (Railsback and Harvey 2002; Eloranta et al. 2011). However, given that observed oxygen levels in the hypolimnion occasionally fell below 5 mg/L, a commonly cited threshold for 694 695 salmonid performance (e.g., Magnuson et al., 1979), this interpretation should be made 696 cautiously. It is also important to note that oxygen was only measured at a single depth 697 (8 m) and at a single location in the lake, limiting our ability to fully characterize oxygen 698 conditions encountered by fish during these forays.

### 699 Morphological differentiation of thermal ecotypes

700 The behavioral tactics we observed were associated with subtle but consistent 701 differences in body shape. "Cool" individuals were more streamlined, with narrower 702 heads, lower body height and thinner caudal peduncles, while "warm" individuals 703 exhibited deeper bodies and broader heads. These differences align with 704 ecomorphological expectations: fusiform shapes minimize drag and support sustained 705 swimming in open water (Gatz 1979; Webb 1984), while deeper bodies and larger heads 706 enhance maneuverability and burst performance, aiding benthic foraging in complex 707 habitats (Malmquist et al. 1992; Walker 1997). This pattern is broadly consistent with 708 previous work on other salmonids (e.g., Chavarie et al., 2013).

709 Interestingly, in our system, "warm" individuals, despite having body shapes more 710 typically associated with maneuverability, exhibited higher swimming speeds than the 711 more streamlined "cool" individuals. This is unexpected, as more fusiform shapes are

generally associated with enhanced sustained swimming (e.g., Peres-Neto & Magnan,
2004). One possibility is that observed swimming speed reflects motivation or behavioral
state rather than morphological capacity. This apparent mismatch underscores the need
for integrative approaches that combine behavioral, morphological, and physiological
data to better understand performance in natural settings.

717 Our findings are consistent with those reported in other polymorphic salmonids such as 718 lake trout and Arctic charr (e.g., Chavarie et al., 2013; Hawley et al., 2016), where 719 morphological divergence corresponds to differences in depth use and foraging strategy. 720 However, this may be one of the first studies to document comparable patterns in brook 721 charr -a species not typically associated with sustained hypolimnion use. This highlights 722 the ecological relevance of deep-water habitats for cold-water fishes more broadly, and 723 supports the emerging view that such habitats may harbor overlooked axes of 724 specialization.

725 Variation in head morphology may also affect feeding efficiency. Longer lower jaws of 726 warm tactic individuals, for instance, can increase mouth gape and speed of closure, 727 improving capture of mobile prey (Kotrschal 1989; Norton 1991; Wainwright and Richard 728 1995). These traits also resemble secondary sexual characteristics common in male 729 salmonids and may involve trade-offs between reproductive investment and feeding 730 efficiency (Kazyak et al. 2013). Indeed, such trade-offs have been documented in other 731 salmonids, where sexually selected traits can reduce swimming performance or increase 732 energetic costs (Makiguchi et al. 2017). Although we did not assess reproductive status, 733 sex was included in the morphological analysis and emerged as a dominant axis of

variation, highlighting the potential for sex-linked traits to influence morphology-functionrelationships.

736 Eye size presented another intriguing pattern:, individuals following the "cool" tactic had 737 smaller eyes, contrary to our expectation that larger eyes would enhance visual sensitivity 738 in the dimly lit hypolimnion (e.g., Andersson et al., 2024). This result also contrasts with 739 findings from Rainville et al. (2021b), where pelagic ecotypes had larger eyes across 740 multiple lakes. However, the relatively larger eyes of "warm" individuals in Lake Ledoux 741 may reflect their frequent use of the epilimnion, which is thought to involve visual cues, 742 while foraging in a more complex habitat of the littoral zone (e.g., Lisney et al., 2020). 743 Smaller eyes in "cool" individuals may indicate reliance on non-visual senses (e.g., 744 chemoreception ; Keefe, 1992), or reflect trade-offs favoring hydrodynamic efficiency or 745 metabolic savings (Schmitz and Wainwright 2011). It is also possible that eye size is 746 constrained by developmental or genetic factors, reflecting local adaptation to deeper, 747 less visually demanding environments. Clarifying the functional and evolutionary drivers 748 of these differences will require further investigation into sensory ecology and 749 developmental plasticity.

Together, these results are consistent with findings in other polymorphic salmonids, where body shape is linked to habitat depth and diet (Robinson and Parsons 2002; Chavarie et al. 2013; Rainville et al. 2021b). In systems like Lake Ledoux, where morphotype distinctions may be subtle or continuous, morphological variation may not fully predict tactic but still contributes to reinforcing behavioral divergence and ecological partitioning.

756 Conclusions and implications

757 Our results highlight intraspecific behavioural specialization in brook charr facing lake 758 warming, characterized by consistent individual differences in habitat use, 759 thermoregulation behavior, and morphology. This fine-scale specialization facilitates 760 coexistence in a shared, heterogeneous environment and underscores the ecological 761 importance of intraspecific diversity (Bolnick et al. 2003; Araújo et al. 2011). By integrating 762 fine-resolution telemetry, environmental data, and morphology, our study provides novel 763 evidence for individual specialization in thermal and foraging niches within a single fish 764 population. Such diversity may buffer populations against environmental change, but 765 lake warming and deoxygenation could reduce the availability of critical habitats for the 766 sustainability of cold-water species. Understanding how individuals partition thermal and 767 trophic resources is critical for predicting species' responses to climate change and 768 conserving behavioral and morphological diversity in freshwater systems, especially as 769 behavioral plasticity may offer one of the few rapid pathways for adaptation (Sunday et al. 770 2014; Sinclair et al. 2016).

771

# 772 References

- Adams, C.E., and Maitland, P.S. 1998. The Ruffe Population of Loch Lomond, Scotland:
  Its Introduction, Population Expansion, and Interaction with Native Species. J Great
  Lakes Res 24(2): 249–262. doi:10.1016/S0380-1330(98)70817-2.
- Amat-Trigo, F., Andreou, D., Gillingham, P.K., and Britton, J.R. 2023. Behavioural
  Thermoregulation in Cold-water Freshwater Fish: Innate Resilience to Climate
  Warming? Fish and Fisheries 24(1): 187–195. John Wiley and Sons Inc.
  doi:10.1111/faf.12720.
- Andersson, M.L., Scharnweber, K., and Eklöv, P. 2024. Environmental and Ecological
  Drivers of Eye Size Variation in a Freshwater Predator: A Trade-off Between Foraging
  and Predation Risk. Funct Ecol 38(11): 2470–2477. British Ecological Society.
  doi:10.1111/1365-2435.14655.

- Araújo, M.S., Bolnick, D.I., and Layman, C.A. 2011. The Ecological Causes of Individual
   Specialisation. Ecol Lett **14**(9): 948–958. doi:10.1111/j.1461-0248.2011.01662.x.
- Arbour, J.H., Hardie, D.C., and Hutchings, J.A. 2011. Morphometric and Genetic Analyses
   of Two Sympatric Morphs of Arctic Char (*Salvelinus alpinus*) in the Canadian High
   Arctic. Can J Zool 89(1): 19–30. doi:10.1139/Z10-100.
- Armstrong, J.B., Ward, E.J., Schindler, D.E., and Lisi, P.J. 2016. Adaptive Capacity at the
   Northern Front: Sockeye Salmon Behaviourally Thermoregulate During Novel
   Exposure to Warm Temperatures. Conserv Physiol 4(1). Oxford University Press.
   doi:10.1093/conphys/cow039.
- Bartosiewicz, M., Przytulska, A., Lapierre, J.F., Laurion, I., Lehmann, M.F., and Maranger,
  R. 2019. Hot Tops, Cold Bottoms: Synergistic Climate Warming and Shielding
  Effects Increase Carbon Burial in Lakes. Limnol Oceanogr Lett 4(5): 132–144. John
  Wiley and Sons Inc. doi:10.1002/lol2.10117.
- Bertolo, A., Pépino, M., Adams, J., and Magnan, P. 2011. Behavioural Thermoregulatory
   Tactics in Lacustrine Brook Charr, *Salvelinus fontinalis*. PLoS One 6(4).
   doi:10.1271/journal.page.0018602
- 799 doi:10.1371/journal.pone.0018603.
- Bolnick, D.I., Svanbäck, R., Fordyce, J.A., Yang, L.H., Davis, J.M., Hulsey, C.D., and
  Forister, M.L. 2003. The Ecology of Individuals: Incidence and Implications of
  Individual Specialization. Am Nat **161**(1): 1–28. doi:10.1086/343878.
- Bourke, P., Magnan, P., and Rodriguez, M.A. 1996. Diel Locomotor Activity of Brook
  Charr, as Determined by Radiotelemetry. J Fish Biol 49: 1174–1185.
- 805 doi:https://doi.org/10.1111/j.1095-8649.1996.tb01787.x.
- Bourke, P., Magnan, P., and Rodriguez, M.A. 1997. Individual Variations in Habitat Use
  and Morphology in Brook Charr. J Fish Biol 51: 783–794.
- 808 doi:https://doi.org/10.1111/j.1095-8649.1997.tb01999.x.
- Brassard, S.G., Rautio, M., and Bertolo, A. 2023. Vertical Distribution Patterns of
  Zooplankton Across a Gradient of Fish Predation in Boreal Lakes. Freshw Biol 68(4):
- 811 588–608. John Wiley and Sons Inc. doi:10.1111/fwb.14049.
- Brett, J.R. 1971. Energetic Responses of Salmon to Temperature. A Study of Some
  Thermal Relations in the Physiology and Freshwater Ecology of Sockeye Salmon
  (*Oncorhynchus nerkd*). Am Zool **11**(1): 99–113. doi:10.1093/icb/11.1.99.
- 815 Brewitt, K.S., and Danner, E.M. 2014. Spatio-temporal Temperature Variation Influences 816 Juvenile Steelhead (*Oncorhynchus mykiss*) Use of Thermal Refuges. Ecosphere
- 817 **5**(7). doi:10.1890/ES14-00036.1.

- Bridger, C.J., and Booth, R.K. 2003. The Effects of Biotelemetry Transmitter Presence and
  Attachment Procedures on Fish Physiology and Behavior. Reviews in Fisheries
  Science 11(1): 13–34. doi:10.1080/16226510390856510.
- Buckley, L.B., Huey, R.B., and Kingsolver, J.G. 2022. Asymmetry of Thermal Sensitivity
  and the Thermal Risk of Climate Change. Global Ecology and Biogeography **31**(11):
  2231–2244. John Wiley and Sons Inc. doi:10.1111/geb.13570.
- Chapman, L.J., and Mckenzie, D.J. 2009. Behavioral Responses and Ecological
  Consequences. *In* Fish Physiology. pp. 25–77. doi:10.1016/S1546-5098(08)00002-2.
- Chavarie, L., Harford, W.J., Howland, K.L., Fitzsimons, J., Muir, A.M., Krueger, C.C., and
  Tonn, W.M. 2016. Multiple Generalist Morphs of Lake Trout: Avoiding Constraints on
  the Evolution of Intraspecific Divergence? Ecol Evol 6(21): 7727–7741. John Wiley
  and Sons Ltd. doi:10.1002/ece3.2506.
- Chavarie, L., Howland, K.L., and Tonn, W.M. 2013. Sympatric Polymorphism in Lake
  Trout: The Coexistence of Multiple Shallow-water Morphotypes in Great Bear Lake.
  Trans Am Fish Soc 142(3): 814–823. doi:10.1080/00028487.2013.763855.
- Clarke, A., and Johnston, N.M. 1999. Scaling of Metabolic Rate with Body Mass and
  Temperature in Teleost Fish. Journal of Animal Ecology 68(5): 893–905.
  doi:10.1046/j.1365-2656.1999.00337.x.
- Cote, D., Tibble, B., Curry, R.A., Peake, S., Adams, B.K., Clarke, K.D., and Perry, R. 2020.
  Seasonal and Diel Patterns in Activity and Habitat Use by Brook Trout (*Salvelinus fontinalis*) in a Small Newfoundland Lake. Environ Biol Fishes **103**(1): 31–47.
  Springer. doi:10.1007/s10641-019-00931-1.
- osa spinigei. doi:10.1007/\$10641-019-00931-1.
- B40 Dermond, P., Sperlich, N., and Brodersen, J. 2019. Heritable Morphological
  B41 Differentiation in Salmonids from Two Distinct Stream Types. J Fish Biol 95(5):
  B42 1215–1222. Blackwell Publishing Ltd. doi:10.1111/jfb.14121.
- B43 Dokter, A.M., Desmet, P., Spaaks, J.H., van Hoey, S., Veen, L., Verlinden, L., Nilsson, C.,
  B44 Haase, G., Leijnse, H., Farnsworth, A., Bouten, W., and Shamoun-Baranes, J. 2019.
  BioRad: Biological Analysis and Visualization of Weather Radar Data. Ecography
  B46 42(5): 852–860. Blackwell Publishing Ltd. doi:10.1111/ecog.04028.
- Eloranta, A.P., Siwertsson, A., Knudsen, R., and Amundsen, P.A. 2011. Dietary Plasticity
  of Arctic Charr (*Salvelinus alpinus*) Facilitates Coexistence with Competitively
  Superior European Whitefish (*Coregonus lavaretus*). Ecol Freshw Fish **20**(4): 558–
  568. doi:10.1111/j.1600-0633.2011.00504.x.
- Ern, R., Andreassen, A.H., and Jutfelt, F. 2023. Physiological Mechanisms of Acute
   Upper Thermal Tolerance in Fish. Physiology 38(3): 141–158. American
- 853 Physiological Society. doi:10.1152/physiol.00027.2022.

854 Fang, X., and Stefan, H.G. 2009. Simulations of Climate Effects on Water Temperature, 855 Dissolved Oxygen, and Ice and Snow Covers in Lakes of the Contiguous United 856 States under Past and Future Climate Scenarios. Limnol Oceanogr **54**(6 PART 2): 857 2359–2370. American Society of Limnology and Oceanography Inc. 858 doi:10.4319/lo.2009.54.6 part 2.2359. 859 Flood, B., Wells, M., Dunlop, E., and Young, J. 2021. Vertical Oscillations of the 860 Thermocline Caused by Internal Waves Modify Coldwater Pelagic Fish Distribution: 861 Results from a Large Stratified Lake. J Great Lakes Res 47(5): 1386–1399. 862 International Association of Great Lakes Research. doi:10.1016/j.jglr.2021.06.010. 863 Gatz, A.J. 1979. Community Organization in Fishes as Indicated by Morphological 864 Features. Ecology 60(4): 711–718. doi:10.2307/1936608. 865 Glaz, P., Sirois, P., and Nozais, C. 2012. Determination of Food Sources for Benthic 866 Invertebrates and Brook Trout Salvelinus fontinalis in Canadian Boreal Shield Lakes 867 Using Stable Isotope Analysis. Aquat Biol 17(2): 107–117. doi:10.3354/ab00465. 868 Goyer, K., Bertolo, A., Pépino, M., and Magnan, P. 2014. Effects of Lake Warming on 869 Behavioural Thermoregulatory Tactics in a Cold-water Stenothermic Fish. PLoS One 870 9(3). Public Library of Science. doi:10.1371/journal.pone.0092514. 871 Grossman, G.D., Coffin, R., and Moyle, P.B. 1980. Feeding Ecology of the Bay Goby 872 (Pisces: Gobiidae). Effects of Behavioral, Ontogenetic, and Temporal Variation on 873 Diet. J Exp Mar Biol Ecol 44(1): 47–59. doi:10.1016/0022-0981(80)90100-8. Guzzo, M.M., Blanchfield, P.J., and Rennie, M.D. 2017. Behavioral Responses to Annual 874 875 Temperature Variation Alter the Dominant Energy Pathway, Growth, and Condition 876 of a Cold-water Predator. Proceedings of the National Academy of Sciences 877 114(37): 9912-9917. doi:10.1073/pnas.1702584114. 878 Guzzo, M.M., Mochnacz, N.J., Durhack, T., Kissinger, B.C., Killen, S.S., and Treberg, J.R. 879 2019. Effects of Repeated Daily Acute Heat Challenge on the Growth and 880 Metabolism of a Cold-water Stenothermal fish. Journal of Experimental Biology 881 222(12). Company of Biologists Ltd. doi:10.1242/jeb.198143. 882 Haesemeyer, M. 2020. Thermoregulation in Fish. Mol Cell Endocrinol 518: 110986. 883 Elsevier Ireland Ltd. doi:10.1016/j.mce.2020.110986. 884 Hawley, K.L., Rosten, C.M., Christensen, G., and Lucas, M.C. 2016. Fine-scale 885 Behavioural Differences Distinguish Resource Use by Ecomorphs in a Closed 886 Ecosystem. Sci Rep 6. Nature Publishing Group. doi:10.1038/srep24369. 887 Iannarilli, F., Gerber, B.D., Erb, J., and Fieberg, J.R. 2024. A 'How-to' Guide for Estimating 888 Animal Diel Activity Using Hierarchical Models. Journal of Animal Ecology. John 889 Wiley and Sons Inc. doi:10.1111/1365-2656.14213.

- Jonsson, B. 2001. Polymorphism and Speciation in Arctic Charr. J Fish Biol 58(3): 605–
   638. doi:10.1006/jfbi.2000.1515.
- Kazyak, D.C., Hilderbrand, R.H., and Holloway, A.E. 2013. Rapid Visual Assessment to
  Determine Sex in Brook Trout. N Am J Fish Manag 33(3): 665–668.
- doi:10.1080/02755947.2013.785998.
- Keefe, M. 1992. Chemically Mediated Avoidance Behavior in Wild Brook Trout, Salvelinus *fontinalis*: the Response to Familiar and Unfamiliar Predaceous Fishes and the
  Influence of Fish Diet. Can J Zool **70**(2): 288–292. doi:10.1139/z92-043.
- Kieffer, J.D., Alsop, D., and Wood, C.M. 1998. A Respirometric Analysis of Fuel Use
  During Aerobic Swimming at Different Temperatures in Rainbow Trout
  (*Oncorhynchus Mykiss*). Journal of Experimental Biology **201**(22): 3123–3133.
  doi:10.1242/jeb.201.22.3123.
- Knudsen, R., Klemetsen, A., Amundsen, P.A., and Hermansen, B. 2006. Incipient
  Speciation Through Niche Expansion: An Example from the Arctic Charr in a
  Subarctic Lake. Proceedings of the Royal Society B: Biological Sciences 273(1599):
  2291–2298. Royal Society. doi:10.1098/rspb.2006.3582.
- Knudsen, R., Primicerio, R., Amundsen, P.A., and Klemetsen, A. 2010. Temporal Stability
  of Individual Feeding Specialization May Promote Speciation. Journal of Animal
  Ecology 79(1): 161–168. doi:10.1111/j.1365-2656.2009.01625.x.
- 809 Kotrschal, K. 1989. Trophic Ecomorphology in Eastern Pacific Blennioid Fishes:
- 910 Character Transformation of Oral jaws and Associated Change of Their Biological
- 911 Roles. Environ Biol Fishes **24**(3): 199–218. Kluwer Academic Publishers.
- 912 doi:https://doi.org/10.1007/BF00001224.
- 913 Lazaridis, E. 2022. lunar: Lunar Phase & Distance, Seasons and Other Environmental
  914 Factors. doi:10.32614/CRAN.package.lunar.
- Learner, M., Wiles, R., and Pickering, J. 1990. Diel Emergence Patterns of Chironomids.
  Internationale Revue der gesamten Hydrobiologie und Hydrographie 5(75).
  doi:https://doi.org/10.1002/iroh.19900750501.
- Leroux, R., Pépino, M., Bordeleau, P.A., Bertolo, A., and Magnan, P. 2025. Animal
  Movement in Space and Time: The Role of Thermal Barriers and Individual Variation
  in Modulating Predator–prey Overlap in Stratified Lakes. Oikos. John Wiley and Sons
  Inc. doi:10.1111/oik.10528.
- 922 Leroux, R., Pépino, M., Magnan, P., and Bertolo, A. 2023. Within-lake Variatiability in
- 923 Predation Risk Shapes the Spatio-temporal Structure of the Zooplankton
- 924 Community. Freshw Biol **68**(12): 2151–2165. John Wiley and Sons Inc.
- 925 doi:10.1111/fwb.14183.

- Lisney, T.J., Collin, S.P., and Kelley, J.L. 2020. The Effect of Ecological Factors on Eye
   Morphology in the Western Rainbowfish, *Melanotaenia australis*. Journal of
   Experimental Biology **223**(10). Company of Biologists Ltd. doi:10.1242/jeb.223644.
- 929 Magnan, P. 1988. Interactions Between Brook Charr, Salvelinus fontinalis, and
- 930 Nonsalmonid Species: Ecological Shift, Morphological Shift, and Their Impact on
- Sooplankton Communities. Canadian Journal of Fisheries and Aquatic Sciences
  45(6): 999–1009. doi:10.1139/f88-123.
- Magnuson, J.J., Crowder, L.B., and Medvick, P.A. 1979. Temperature as an Ecological
  Resource. Am Zool 19(1): 331–343. doi:10.1093/icb/19.1.331.
- Makiguchi, Y., Nii, H., Nakao, K., and Ueda, H. 2017. Sex Differences in Metabolic Rate
  and Swimming Performance in Pink Salmon (*Oncorhynchus gorbuscha*): the Effect
  of Male Secondary Sexual Traits. Ecol Freshw Fish **26**(2): 322–332. Blackwell
  Munksgaard. doi:10.1111/eff.12278.
- 939 Malmquist, H.J., Snorrason, S.S., Skúlason, S., Jonsson, B., Sandlund, O.T., and
- 940 Jonasson, P.M. 1992. Diet Differentiation in Polymorphic Arctic Charr in
- 941 Thingvallavatn, Iceland. Source: Journal of Animal Ecology 61(1): 21–35.
  942 doi:10.2307/5505.
- Martin, T.L., and Huey, R.B. 2008. Why "Suboptimal" Is Optimal: Jensen's Inequality and
  Ectotherm Thermal Preferences. American Naturalist **171**(3). doi:10.1086/527502.
- McCullough, D.A., Bartholow, J.M., Jager, H.I., Beschta, R.L., Cheslak, E.F., Deas, M.L.,
  Ebersole, J.L., Foott, J.S., Johnson, S.L., Marine, K.R., Mesa, M.G., Petersen, J.H.,
  Souchon, Y., Tiffan, K.F., and Wurtsbaugh, W.A. 2009. Research in Thermal Biology:
  Burning Questions for Coldwater Stream Fishes. Reviews in Fisheries Science
  17(1): 90–115. doi:10.1080/10641260802590152.
- Mellas, E.J., and Haynes, J.M. 1985. Swimming Performance and Behavior of Rainbow
  Trout (*Salmo gairdneri*) and White Perch (*Morone americana*): Effects of Attaching
  Telemetry Transmitters. Canadian Journal of Fisheries and Aquatic Sciences 42(3):
  488–493. doi:10.1139/f85-066.
- Naud, M., and Magnan, P. 1988. Diel Onshore–offshore Migrations in Northern Redbelly
   Dace, *Phoxinus eos* (Cope), in Relation to Prey Distribution in a Small Oligotrophic
   Lake. Can J Zool **66**(6): 1249–1253. doi:10.1139/z88-182.
- Norton, S.F. 1991. Capture Success and Diet of Cottid Fishes: The Role of Predator
  Morphology and Attack Kinematics. Ecology 72(5): 1807–1819.
  doi:10.2307/1940980.
- 960 Pépino, M., Goyer, K., and Magnan, P. 2015. Heat Transfer in Fish: Are Short Excursions
  961 Between Habitats a Thermoregulatory Behaviour to Exploit Resources in an

962 Unfavourable Thermal Environment? Journal of Experimental Biology **218**(21): 963 3461–3467. Company of Biologists Ltd. doi:10.1242/jeb.126466. 964 Pépino, M., Magnan, P., Leroux, R., and Bertolo, A. 2024. Thermal Habitat Fragmentation 965 in Stratified Lakes Induces Resource Waves that Brook Charr Track Across Seasons. 966 Oikos. John Wiley and Sons Inc. doi:10.1111/oik.10539. 967 Pépino, M., Magnan, P., and Proulx, R. 2018. Field Evidence for a Rapid Adaptive Plastic 968 Response in Morphology and Growth of Littoral and Pelagic Brook Charr: A 969 Reciprocal Transplant Experiment. Funct Ecol 32(1): 161–170. Blackwell Publishing 970 Ltd. doi:10.1111/1365-2435.12929. 971 Peres-Neto, P.R., and Magnan, P. 2004. The Influence of Swimming Demand on 972 Phenotypic Plasticity and Morphological Integration: A Comparison of Two 973 Polymorphic Charr Species. Oecologia 140(1): 36-45. doi:10.1007/s00442-004-974 1562-y. 975 Power, M., O'Connell, M.F., and Dempson, J.B. 2005. Ecological Segregation Within and 976 Among Arctic Char Morphotypes in Gander Lake, Newfoundland. Environ Biol Fishes 73(3): 263-274. doi:10.1007/s10641-005-2137-4. 977 978 Proulx, R., and Magnan, P. 2004. Contribution of Phenotypic Plasticity and Heredity to 979 the Trophic Polymorphism of Lacustrine Brook Charr (Salvelinus fontinalis M.). Evol 980 Ecol Res 6: 503–522. Available from 981 https://www.researchgate.net/publication/228737668. 982 R Core Team. 2017. R: A Language and Environment for Statistical Computing. R 983 Foundation for Statistical Computing, Vienna, Austria. Available from 984 https://www.r-project.org/. 985 Rahel, F.J., and Nutzman, J.W. 1994. Foraging in a Lethal Environment: Fish Predation in 986 Hypoxic Waters of a Stratified Lake. Ecology 75(5): 1246–1253. 987 doi:10.2307/1937450. 988 Railsback, S.F., and Harvey, B.C. 2002. Analysis of Habitat-Selection Rules Using an 989 Individual-Based Model. Ecology 83(7): 1817. doi:10.2307/3071767. 990 Rainville, V., Dupuch, A., Pépino, M., and Magnan, P. 2022. Intraspecific Competition 991 and Temperature Drive Habitat-based Resource Polymorphism in Brook Charr, 992 Salvelinus fontinalis. Evol Ecol 36(6): 967–986. Springer Science and Business 993 Media Deutschland GmbH. doi:10.1007/s10682-022-10212-6. 994 Rainville, V., Filion, A., Lussier, I., Pépino, M., and Magnan, P. 2021a. Does Ecological 995 Release from Distantly Related Species Affect Phenotypic Divergence in Brook 996 Charr? Oecologia 195(1): 77–92. Springer Science and Business Media 997 Deutschland GmbH. doi:10.1007/s00442-020-04822-6.

- Rainville, V., Pépino, M., and Magnan, P. 2021b. Parallel Evolution of Morphological Traits
  and Body Shape in Littoral and Pelagic Brook Charr, *Salvelinus fontinalis*, Along a
  Gradient of Interspecific Competition. Oecologia **197**(2): 421–436. Springer Science
  and Business Media Deutschland GmbH. doi:10.1007/s00442-021-05028-0.
- Roberts, J.J., Grecay, P.A., Ludsin, S.A., Pothoven, S.A., Vanderploeg, H.A., and Höök,
  T.O. 2012. Evidence of Hypoxic Foraging Forays by Yellow Perch (*Perca flavescens*)
  and Potential Consequences for Prey Consumption. Freshw Biol **57**(5): 922–937.
  doi:10.1111/j.1365-2427.2012.02753.x.
- Robinson, B.W., and Parsons, K.J. 2002. Changing Times, Spaces, and Faces: Tests and
  Implications of Adaptive Morphological Plasticity in the Fishes of Northern
  Postglacial Lakes. Canadian Journal of Fisheries and Aquatic Sciences 59(11):
  1819–1833. doi:10.1139/f02-144.
- Rodrigues, T.H., Chapelsky, A.J., Hrenchuk, L.E., Mushet, G.R., Chapman, L.J., and
  Blanchfield, P.J. 2022. Behavioural Responses of a Cold-water Benthivore to Loss of
  Oxythermal Habitat. Environ Biol Fishes **105**(10): 1489–1507. Springer Science and
  Business Media B.V. doi:10.1007/s10641-022-01335-4.
- 1014 Rohlf, F. 2012a. tpsDig2. Department of Ecology and Evolution, State University of New1015 York, Stony Brook.
- 1016 Rohlf, F. 2012b. tpsUtil. Department of Ecology and Evolution, State University of New1017 York, Stony Brook.

Schmitz, L., and Wainwright, P.C. 2011. Nocturnality Constrains Morphological and
Functional Diversity in the Eyes of Reef Fishes. BMC Evol Biol 11(1).
doi:10.1186/1471-2148-11-338.

- Sinclair, B.J., Marshall, K.E., Sewell, M.A., Levesque, D.L., Willett, C.S., Slotsbo, S., Dong,
   Y., Harley, C.D.G., Marshall, D.J., Helmuth, B.S., and Huey, R.B. 2016. Can We
   Predict Ectotherm Responses to Climate Change Using Thermal Performance
   Curves and Body Temperatures? Ecol Lett 19(11): 1372–1385. Blackwell Publishing
   Ltd. doi:10.1111/ele.12686.
- Smith, D.A., Jackson, D.A., and Ridgway, M.S. 2020. Thermal Habitat of Brook Trout in
   Lakes of Different Size. Freshwater Science 39(1): 56–69. University of Chicago
   Press. doi:10.1086/707488.
- Sunday, J.M., Bates, A.E., Kearney, M.R., Colwell, R.K., Dulvy, N.K., Longino, J.T., and
   Huey, R.B. 2014. Thermal-safety Margins and the Necessity of Thermoregulatory
   Behavior Across Latitude and Elevation. Proc Natl Acad Sci U S A 111(15): 5610–
   5615. National Academy of Sciences. doi:10.1073/pnas.1316145111.

- Thomas, Z.R., Beauchamp, D.A., Clark, C.P., and Quinn, T.P. 2023. Seasonal Shifts in
   Diel Vertical Migrations by Lake-dwelling Coastal Cutthroat Trout, *Oncorhynchus clarkii clarkii*, Reflect Thermal Regimes and Prey Distributions. Ecol Freshw Fish
   32(4): 842–851. doi:10.1111/eff.12725.
- 1037 Venables, W.N., and Ripley, B.D. 2002. Modern Applied Statistics with S. *In* 4th edition.
  1038 Springer, New York. doi:https://doi.org/10.1007/978-0-387-21706-2.
- Wainwright, P.C., and Richard, B.A. 1995. Predicting Patterns of Prey Use from
  Morphology of Fishes. Environ Biol Fishes 44(1–3): 97–113.
- 1041 doi:10.1007/BF00005909.
- Walker, J.A. 1997. Ecological Morphology of Lacustrine Threespine Stickleback
   *Gasterosteus aculeatus* L. (Gasterosteidae) Body Shape. Biological Journal of the
   Linnean Society 61(1): 3–50. doi:10.1111/j.1095-8312.1997.tb01777.x.
- 1045 Webb, P.W. 1984. Body Form, Locomotion and Foraging in Aquatic Vertebrates. Am Zool
  1046 24(1): 107–120. doi:10.1093/icb/24.1.107.
- Whelan, K.F. 1980. Some Aspects of the Biology of *Ephemera Danica* Müll
  (Ephemeridae: Ephemeroptera) in Irish Waters. *In* Advances in Ephemeroptera
  Biology. Springer US, Boston, MA. pp. 187–199. doi:10.1007/978-1-4613-3066-0\_16.
- Winslow, L., Read, J., Woolway, R., Brentrup, J., Leach, T., Zwart, J., Albers, S., and
   Collinge, D. 2013, September 8. rLakeAnalyzer: Lake Physics Tools.
- 1052 doi:10.32614/CRAN.package.rLakeAnalyzer.
- 1053 Wood, S.N. 2017. Generalized Additive Models: An Introduction with R. *In* 2nd edition.
  1054 Chapman and Hall/CRC. doi:https://doi.org/10.1201/9781315370279.
- Zimmerman, M.S., Krueger, C.C., and Eshenroder, R.L. 2007. Morphological and
   Ecological Differences Between Shallow- and Deep-water Lake Trout in Lake
   Mistassini, Quebec. J Great Lakes Res 33(1): 156–169. International Association of
- 1058 Great Lakes Research. doi:https://doi.org/10.3394/0380-
- 1059 1330(2007)33[156:MAEDBS]2.0.CO;2.
- Zuur, A.F., Ieno, E.N., Walker, N., Saveliev, A.A., and Smith, G.M. 2009. Mixed Effects
  Models and Extensions in Ecology with R. Springer New York, New York, NY.
  doi:10.1007/978-0-387-87458-6.

- 1064 **Supplementary material for:** Behavioural tactics across thermal gradients align with
- 1065 partial morphological divergence in brook charr
- 1066 Author names: Aliénor Stahl<sup>1</sup>, Marc Pépino<sup>1,2</sup>, Andrea Bertolo<sup>1</sup>, Pierre Magnan<sup>1</sup>
- 1067 Affiliations:
- 1068 (1) Centre de Recherche sur les Interactions Bassins Versants Écosystèmes
   1069 Aquatiques (RIVE), Université du Québec à Trois-Rivières, Trois-Rivières, QC,
   1070 Canada
- 1071 (2) Current address: Direction de la Gestion de la Faune Mauricie Centre-du-
- 1072 Québec, Ministère de l'Environnement, de la Lutte Contre les Changements
- 1073 Climatiques, de la Faune et des Parcs, Trois-Rivières, QC, Canada
- 1074 Corresponding author: Aliénor Stahl, alienor.stahl@uqtr.ca
- 1075 Journal: Journal of Animal Ecology
- 1076 Keywords (up to 8): thermal stratification; behavioral thermoregulation; vertical forays;

1077 diel movement patterns; morphological divergence; acoustic telemetry; *Salvelinus*1078 *fontinalis* 



Layer — Epilimnion — Metalimnion — Hypolimnion

1080 Figure S1: Seasonal variation in stratification metrics in Lake Ledoux: (A) Water 1081 temperature (°C) and (B) depth (m) of thermal layer boundaries as a function of day of the 1082 year. In panel A, solid lines indicate the mean daily temperature within each thermal layer: 1083 red for the epilimnion, green for the metalimnion, and blue for the hypolimnion. In panel 1084 B, solid lines represent the lower boundary of the epilimnion (red), the upper boundary of 1085 the hypolimnion (blue), and the thermocline (green). Vertical dashed lines mark October 31<sup>st</sup>, 2017 and May 19<sup>th</sup>, 2018—cut-off dates used to define the analyzed stratified period. 1086 1087 Outside this window, thermal stratification was weak or inconsistent, and layer 1088 boundaries could not be reliably identified.



**Figure S2:** Schematic representation of morphological traits measured in brook charr. Blue dashed lines indicate the 11 linear traits recorded at the beginning of the study. Measurements include: BH, body height; DB, dorsal fin base length; EW, eye width; HH, head height; HL, head length; MW, mouth width (measured dorsally); PC, pectoral fin length; PH, pelvic fin height; PS, peduncle height; SN, snout-to-eye distance; and SJ, lower jaw length. Green dots denote curve points used to "unbend" fish before conducting geometric morphometric analyses.



1098 Figure S3: Metrics describing brook charr vertical habitat use as a function of day of year 1099 and average daily epilimnion temperature. Panels show summary statistics for forays into 1100 the (A) epilimnion and (B) hypolimnion, including: number of forays per day (defined as a 1101 transition into the target layer following a detection in a different layer), total daily duration 1102 of forays (in hours), maximum foray duration (in minutes), and mean foray duration (in 1103 minutes). A fish is considered to be in the epilimnion or hypolimnion when located at least 1104 1 meter from the adjacent metalimnion boundary. Color scale reflects average daily 1105 epilimnion temperature (yellow = warm, dark purple = cold). Solid lines represent the 50th 1106 percentile (median), and dotted lines the 95th percentile for each metric. Line color 1107 corresponds to the thermal layer: red for epilimnion, blue for hypolimnion. See Figure 1108 S1 for (i) daily depth estimates of thermal layers,; (ii) a version of panel A for hypolimnion 1109 forays in 2018 colored by average daily dissolved oxygen (Figure S4); and (iii) per-fish 1110 histograms of epilimnion foray durations (Figure S5).



1112 Figure S4: Metrics describing brook charr use of the hypolimnion as a function of day of 1113 the year and average daily dissolved oxygen (mg/L) at 8 meters depth. The figure 1114 summarizes hypolimnion forays, including: (i) number of forays per day, defined as 1115 transitions into the hypolimnion following a detection in a different layer; (ii) total daily 1116 foray duration (in hours); (iii) maximum foray duration (in minutes); and (iv) mean foray 1117 duration (in minutes). Only days with available dissolved oxygen data are included. The 1118 color scale represents average daily dissolved oxygen at 8 m depth (light blue = oxygenated, dark blue = hypoxic). See Figure S1 for (i) daily depth estimates of thermal 1119 1120 layers,; (ii) an alternative version using a more conservative threshold of  $\geq 1$  m from the metalimnion boundary (Figure S3); and (iii) per-fish histograms of epilimnion foray 1121 1122 durations (Figure S5).



**Figure S5:** Individual-level histograms of epilimnion foray durations in brook charr. Histogram of foray durations into the epilimnion (in minutes) for each tagged individual. Each bar represents the number of forays of a given duration. Fish are ordered by their position along the first principal component (PC1) from the PCA of vertical habitat use. Facet title colors indicate thermal classification based on the PCA: teal for "cool" individuals and orange for "warm" individuals. Mean body temperature (°C) is shown below each fish ID for reference.



Figure S6: Proportional habitat and depth layer use per fish. Each facet represents an individual brook charr, showing the proportion of detections in different combinations of habitat (benthic = within 2 m of the lake bottom; pelagic = elsewhere) and thermal layer: epilimnion (red), metalimnion (green), and hypolimnion (blue). For each fish, the stacked bar totals 1, representing the full distribution of detections across habitat–layer categories. Fish are ordered based on their PC1 score from the PCA of vertical behavior.

- 1139 teal and "warm" fish in orange. Mean body temperature (°C) for each fish is displayed
- 1140 beneath the fish ID.



1142 Figure S7: Variable contributions to the first two axes of a Principal Component Analysis 1143 (PCA) describing brook charr vertical behavior and habitat use, based on data from July 1144 and August only. The PCA was based on standardized, per-fish values for: mean body 1145 temperature (°C; Temp), proportion of time spent in the epilimnion and hypolimnion (0–1; 1146 Time.Epi and Time.Hypo), average swimming speed across all layers (TL/s; Speed), 1147 average speed in the epilimnion and hypolimnion (Speed.Epi, Speed.Hypo), mean distance from the lake bottom (m; Distance.Bottom), mean distance to shore (m; 1148 1149 Distance.Shore), and mean foray durations into the epilimnion and hypolimnion (minutes;

- 1150 Duration. Foray. Epi, Duration. Foray. Hypo). Supplementary variables (not used in the PCA)
- 1151 represent the proportion of detections in combinations of habitat type and thermal layer.
- 1152 Habitat is classified as benthic (within 2 m of the lake bottom; B) or pelagic (P), combined
- 1153 with depth zone: Epi = epilimnion, Meta = metalimnion, Hypo = hypolimnion (e.g., B-Epi =
- 1154 *benthic epilimnion, P-Hypo = pelagic hypolimnion). These values sum to 1 for each fish.*
- 1155 This figure allows assessment of whether PCA structure is sensitive to seasonal
- 1156 environmental variation by focusing only on the peak summer months.



**Figure S8:** Individual-level diel vertical movement patterns. Predicted probability (range: 0–1) of transitioning into the epilimnion (red) or hypolimnion (blue) as a function of time of day (hour), based on HGAMs fitted separately for each fish and each layer. Models include sunrise and sunset times as fixed effects. Facet titles are colored by thermal classification based on PCA results: "cool" fish are labeled in teal, and "warm" fish in orange. Fish are ordered based on their PC1 score from the PCA of vertical behavior. Mean body temperature (°C) for each fish is displayed beneath the fish ID.



1165

Figure S9: Monthly diel vertical movement patterns of brook charr by thermal behavioral 1166 1167 category. Predicted diel probability of transitioning into the epilimnion (red) or hypolimnion (blue) for brook charr grouped into "warm" and "cool" categories based on 1168 1169 their position along the first principal component axis (PC1) from the PCA of vertical 1170 thermal behavior. Separate hierarchical generalized additive models (HGAMs) were fitted 1171 for each behavioral group within each month (June–October), allowing assessment of 1172 temporal variation in vertical movement patterns. Predictions are shown over a 24-hour 1173 cycle. Black vertical lines indicate sunrise and sunset times, which were included as 1174 covariates. An "average" day is defined by the median sunrise and sunset times for each 1175 month.



Figure S10: Monthly spatial maps of depth-layer usage by brook charr thermal tactics. Each panel shows the geographic distribution of individual fish detections within the lake, faceted by month (June–October), thermal tactic ("warm" or "cool"), and vertical layer (epilimnion, metalimnion, hypolimnion). Points represent density of individual detections, allowing visualization of how habitat use varies spatially and temporally across tactics.



- **Figure S11:** Spatial maps of depth-layer usage by brook charr thermal tactics. Each panel shows the geographic distribution of individual
- 1183 fish detections within the lake, faceted by thermal tactic ("warm" or "cool") and vertical layer (epilimnion, metalimnion, hypolimnion).
- 1184 Points represent density of individual detections, allowing visualization of how habitat use varies spatially across tactics.