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Breeding stage and tissue isotopic consistency suggests colonylevel flexibility in niche breadth of an Arctic marine bird

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1	Breeding stage and tissue isotopic consistency suggests colony-level flexibility in niche
2	breadth of an Arctic marine bird
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10	
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20 Abstract

21 Organisms must overcome environmental limitations to optimize their investment in life history 22 stages to maximize fitness. Human-induced climate change is generating increasingly variable 23 environmental conditions, impacting the demography of prey items and therefore the ability of 24 consumers to successfully access resources to fuel reproduction. While climate change effects 25 are especially pronounced in the Arctic, it is unknown whether organisms can adjust foraging 26 decisions to match such changes. We used a 9-year blood plasma δ^{13} C and δ^{15} N dataset from 27 over 700 pre-breeding Arctic common eiders (Somateria mollissima) to assess breeding-stage 28 and inter-annual variation in isotopic niche, and whether inferred trophic flexibility was related to colony-level breeding parameters and environmental variation. Eider blood isotope values 29 30 varied both across years and breeding stages, and combined with only weak relationships 31 between isotopic metrics and environmental conditions suggests that pre-breeding eiders can 32 make flexible foraging decisions to overcome constraints imposed by local abiotic conditions. 33 From an investment perspective, an inshore, smaller isotopic niche predicted a greater probability to invest in reproduction, but was not related to laying phenology. Proximately, our 34 35 results provide evidence that eiders breeding in the Arctic can alter their diet at the onset of 36 reproductive investment to overcome increases in the energetic demand of egg production. 37 Ultimately, Arctic pre-breeding common eiders may have the stage- and year-related foraging 38 flexibility to respond to abiotic variation to reproduce successfully.

39 Keywords: isotopic niche, foraging flexibility, phenology, trophic interactions, reproductive

- 40 investment, adaptive capacity, carbon-13, nitrogen-15
- 41

42 Introduction

Investment in various life history stages is significantly influenced by an individual's ability to 43 obtain energetic resources and optimally allocate them to meet the energetic demands of 44 associated life history (McNamara and Houston 1996). Consequently, resource limitation is one 45 46 of the strongest constraints influencing the ability of an individual to optimize foraging decisions 47 to maximize fitness (Stephens and Krebs 1986; Newton 1998). Since the quality and quantity of 48 available dietary resources can be influenced by variability in the surrounding environment, 49 variation in environmental conditions can play an important role in shaping the relationship 50 between resource acquisition, and fitness and population demography (Boggs 1992). However, 51 as climate change continues to generate increases in mean annual temperatures and variability in 52 climatic conditions (IPCC 2018), animals are expected to be increasingly impacted by mounting 53 instability in resource availability (Cushing 1990; Tylianakis et al. 2008). Relative to other temperate and tropical ecosystems, the Arctic is experiencing amplified 54 55 rates of climate change (Wassmann et al. 2011), which is reducing sea-ice extent (Johannessen et 56 al. 2004; Comiso et al. 2008; Hoegh-Guldberg and Bruno 2010; Ciancio et al. 2016), and leading to bottom-up trophic disruptions (Wassmann et al. 2011; Boeitus et al. 2013; Jones et al. 2014; 57 58 Meier et al. 2014). As such, many key prey sources are declining or their distributions are 59 shifting (Both et al. 2006), complicating species' ability to acquire resources and their ability to 60 optimize investment decisions in energetically demanding life history stages, such as 61 reproduction (Ward et al. 2009; Seyboth et al. 2016). Whether Arctic species have the required adaptive capacity to be flexible enough to optimize breeding decisions to proximately keep pace 62 63 with the current rate of environmental change has therefore become an important topic of

64 investigation (Kovacs et al. 2010; Moore and Huntington 2011; Descamps et al. 2017).

65 Determining the downstream influence of environmental variation on reproductive investment first requires consideration of which prey items organisms consume to fuel 66 reproduction (Walther et al. 2002; Rutschmann et al. 2016). The use of tissue stable isotope 67 68 measurements to represent foraging niche is well established in several taxa (Bearhop et al. 69 2004) and is increasingly used to assess individual- and population-level responses to rapid 70 environmental change (Dawson and Siegwolf 2011; Mancinelli and Vizzini 2015; Pethybridge et 71 al. 2017). Specifically, the combination of stable isotope measurements of carbon (δ^{13} C), which provides information on basal sources reflective of habitat use (i.e., inshore vs offshore), and 72 73 nitrogen (δ^{15} N), which allows relative estimation of trophic level (Peterson and Fry 1987), 74 provides a non-lethal method of quantifying isotopic niche (Newsome et al. 2007; Matich and 75 Shipley 2021). Quantifying isotopic niche space or volume can provide insight on the degree of 76 trophic specialization (Seamon and Adler 1996), where groups with larger isotopic niches are predicted to have more diverse diets (i.e., more generalist; Moreno et al. 2010). The degree of 77 78 trophic specialization can then be used to predict how resilient organisms or populations will be 79 to further environmental change (Polito et al. 2015). For example, groups with a more generalist 80 isotopic niche might be expected to be more successful investing in reproduction, regardless of 81 environmentally induced trophic disruptions because of their greater dietary flexibility (Seamon and Adler 1996). Further, isotopic niche provides a useful tool to assess shifts in community-82 83 wide trophic dynamics in response to environmental change (Hobson 1992a, b, 1999; Newsome 84 et al. 2007; Herman et al. 2017), and insight into how foraging decisions might affect key fitness-related decisions (Hutchinson 1957; Vandermeer 1972; Alatalo 1982; Bolnick et al. 85 86 2003). Finally, quantification of isotopic niche also provides a testable framework in which to 87 investigate how environmental variability mediates trophic dynamics and ultimately influences

88	reproductive investment decisions (Chesson 1986; Leibold 1995); however, this framework
89	relies on a number of assumptions (Matich and Shipley 2021). The primary assumption being
90	that baseline isotope values remain constant or are accounted for through periods of
91	investigation. Secondly, tracing nutrients isotopically during reproduction is complex and
92	depends on the degree to which birds may be income or capital breeders (Hobson 2006, Hobson
93	et al. 2015, Whiteman et al. 2021). Even considering these caveats, establishing temporal
94	variability in the isotopic niches of populations can be a useful tool for examining the influence
95	of environmental change in polar regions which are historically relatively slow to change over
96	time compared to more southern regions (Yurkowski et al. 2020).
97	Female common eiders (Somateria mollissima) make an ideal study species to test these
98	linkages given that a key environmental factor limiting Arctic-nesting eiders is sea ice cover,
99	particularly during spring migration and the pre-laying period which can restrict access to
100	foraging grounds of this diving seabird (Jean-Gagnon et al. 2018). Additionally, female eiders
101	must meet a minimum body condition threshold to invest in reproduction (Hennin et al. 2016),
102	and their ability to do so is influenced by access to resources just prior to breeding (Love et al.
103	2010; Jean-Gagnon et al. 2018). This is an important consideration because female eiders that
104	can fatten more quickly lay earlier (Hennin et al. 2017, 2019), invest in larger clutches
105	(Descamps et al. 2011a; Hennin et al. 2018) and ultimately recruit more ducklings into the
106	breeding colony (Love et al. 2010; Descamps et al. 2011b). Since eiders rely on lipid and protein
107	sources consumed on the breeding grounds to produce their eggs (Sénéchal et al. 2011), and prey
108	items used during egg formation differ in nutritional value (Paiva et al. 2013), the ability to
109	access nutritious prey likely influences breeding decisions and outcomes (e.g., Kitaysky et al.
110	2010), especially under climatically unpredictable conditions (Barbraud et al. 2012). With

isotopic niche capturing a general snapshot of female foraging prior to and during laying, it may
act as an important predictor of variation in reproductive investment (Sénéchal et al. 2011).

113 Here we used a 9-consecutive-year (2010-2018) isotopic dataset collected from over 700 114 Arctic-nesting female common eiders (hereafter eiders) to investigate the linkages between 115 environmental variability, variation in important breeding metrics, and temporal variation in 116 isotopic niche at an Arctic-breeding colony. Specifically, our first objective was to quantify 117 variation in isotopic niche across breeding stages and years. Given the nutritional demands of 118 egg formation (Descamps et al. 2010; Love et al. 2010; Sénéchal et al. 2011; Hennin et al. 2015, 119 2016, 2018, 2019), we predicted that as birds approached laying, they would have higher $\delta^{15}N$ 120 values and higher δ^{13} C values as birds targeted higher trophic level prey (Forero et al. 2002; 121 Becker et al. 2007; Gonzálex-Medina et al. 2018) found closer to shore, respectively. Likewise, 122 given the energetic demands of initiating and fueling egg formation, we predicted that in years 123 when the colony exhibited a larger isotopic niche (more generalized foraging strategy) more 124 birds would have greater access to a diversity of resources, and therefore be able to fatten more 125 quickly, generating shorter mean delays before laying, earlier mean laying dates, and higher 126 colony-level breeding propensity (Love et al. 2010; Descamps et al. 2011; Hennin et al. 2015, 127 2016, 2018). Finally, we predicted significant inter-annual variation in isotopic niches, driven by 128 inter-annual variation in climate and trophic dynamics. Considering the highly variable sea ice 129 conditions during the pre-breeding period (Love et al. 2010; Jean-Gagnon et al. 2018), and that 130 local environmental conditions can influence seabird foraging behaviour (Hobson 1999; Paiva et 131 al. 2013), our second objective was to examine whether broad-scale environmental conditions 132 explained inter-annual variation in isotopic niche. We predicted that in years with warmer 133 winters, warmer springs, and higher overall ambient temperatures (as indicated by the North

134 Atlantic Oscillation [NAO] index), there ought to be a greater degree of open water, and

135 therefore a wider variety of possible prey items available to eiders, resulting in a broader colony-

wide isotopic niche. 136

137

138 Methods

139 *Study species and breeding parameters*

140 From 2010 to 2018, sampling was undertaken at the largest and longest continually monitored 141

colony of common eiders in the eastern Canadian Arctic; at East Bay Island (EBI), which is

142 found within in the Qaqsauqtuuq Migratory Bird Sanctuary, Nunavut, Canada (64°02'N,

143 81°47′W). EBI is a small (800 m x 400 m), low lying island (<8m elevation). Females breeding

144 at this colony migrate from their wintering grounds off the western coast of Greenland and the

145 northern coast of Newfoundland and Labrador, Canada in May (Mosbech et al. 2006; Steenweg

146 et al. 2017), arrive at EBI in early to mid-June, and lay their eggs in mid-June to early-July

147 (Hennin et al. 2015; Jean-Gagnon et al. 2018). Using flight nets, we captured female common

148 eiders as they flew over the colony in mid-June, coinciding with their timing of arrival at the

149 breeding grounds (Descamps et al. 2010; see Supplementary Materials Table S1).

150 After capture, females were blood sampled within three minutes of capture (Romero and 151 Reed 2005) from the tarsal vein using a 1-mL heparinized syringe and 23G thin-wall, 0.5-inch 152 needle (see Hennin et al. 2015, 2016 for details). Plasma was separated from red blood cells, and 153 both components were frozen at -20°C. After sampling, females were assigned a metal band and 154 alpha-numeric Darvic bands, then affixed with a unique combination of coloured and shaped 155 nasal tag plastic discs using UV degradable monofilament. Females were identified within the 156 breeding colony by their nasal tags using spotting-scopes from seven permanent blinds

157 positioned around the periphery of the island. By observing the females and their associated 158 behaviours, we obtained: breeding propensity (probability of breeding) and lay date (and therefore the interval in days between arrival at the colony and laying). From these data, we 159 160 assigned reproductive stage to all birds as either laying (LAY), rapid follicle growth (RFG), pre-161 recruiting (PR), and non-breeding (NB) (see Hennin et al. 2015 for details; see Supplementary 162 Material). Laying and incubating females were determined based on careful observations from 163 blinds and monitoring of nests, along with noting the presence of an egg in the oviduct at 164 capture. All work was approved by the animal care committees of the University of Windsor 165 (AUPP 11-06 and 19-11) and Environment and Climate Change Canada (EC-PN-15-026).

166

167 Environmental indices

168 We selected climate variables predictive of the storm activity and ice conditions eiders face 169 during the pre-breeding period (see Supplementary Materials Table S2). The North Atlantic 170 Oscillation (NAO) index was used as a proxy for inter-annual variation in environmental 171 conditions, as it impacts a wide area across the Northern Atlantic Ocean. The relationship 172 between temporal variation in NAO values and energetic constraints has been validated in 173 multiple seabird species (Stenseth et al. 2003; Hallett et al. 2004; Sandvik and Erikstad 2008), 174 including eiders (Descamps et al. 2010). We calculated the average winter NAO (December-175 March), which directly impacts the arrival body condition of female eiders (Descamps et al. 176 2010) and the average spring NAO (April-July; pre-breeding conditions for eiders at MI). All 177 NAO values were obtained from the National Weather Service 178 (https://www.cpc.ncep.noaa.gov/). As a proxy for localized environmental conditions at the 179 breeding grounds, we used air temperature (Ta) measured at the Coral Harbour Airport Weather Station (70 km from the breeding colony). We first calculated the mean Ta for each individual on the dates that coincided with the isotopic half-life of plasma (i.e., air temperature roughly 3 days prior to sample collection; Hobson and Clark 1993; Hahn et al. 2012). This value was then averaged across all the individuals in the colony for each year.

184

185 *Stable isotope analysis and niche metrics*

186 Female eiders caught at arrival to EBI ranged in body mass from 1.5-2 kg (Hennin et al. 2015), making the use of plasma (with a half-life of plasma approximately three days; Hobson and 187 188 Clark 1992; Hahn et al. 2012) appropriate to assess the most recent isotopic niche space occupied 189 by these individuals (Supplementary Materials Table S3). We freeze-dried 100 uL of plasma 190 from each individual until achieving a constant mass (minimum of 78 hours). All samples were 191 then ground into a homogenized, fine powder using a metal spatula. Since plasma is often high in 192 lipids we then lipid extracted all plasma samples using a 2:1 choloroform:methanol solution (Bligh and Dyer 1959). We weighed between 0.3-0.5 mg of each sample, using a four-digit 193 194 balance (Sartorius AG, Model CP2P, Gottingen, Germany), into individual 3.5x5 mm tin capsules for δ^{13} C and δ^{15} N analysis. Analyses for plasma isotopes were conducted using 195 196 continuous-flow isotope-ratio mass spectrometry (CFIRMS) at the Environment Canada Stable 197 Isotope Hydrology and Ecology Research Laboratory in Saskatoon, Saskatchewan. Encapsulated 198 plasma was combusted at 1030°C in a Carlo Erba NA1500 or Eurovector 3000 elemental 199 analyser. The resulting N₂ and CO₂ were separated chromatographcally and introduced to an 200 Elementar Isoprime or a Nu Instruments Horizon isotope ratio mass spectrometer. We used two reference materials to normalize the results to VPDB and AIR: BWBIII keratin ($\delta^{13}C = -20.18$, 201 $\delta^{15}N = +14.31$ ‰, respectively) and PRCgel ($\delta^{13}C = -13.64$, $\delta^{15}N = +5.07$ ‰, respectively). 202

Within-run (n = 5) precisions as determined from both reference and sample duplicate analyses 203 were \pm 0.1 ‰ for both $\delta^{13}C$ and $\delta^{15}N.$ 204

205	We used a bivariate approach to calculate isotopic niche size for each year and each
206	breeding stage within the SIBER package (Jackson et al. 2011) in R (R Core Team 2014,
207	Version 4.0.3). Calculated niche metrics included: mean next-neighbor distance (MNND),
208	standard ellipse area (SEA _C), X range (i.e., maximum range of δ^{13} C) and Y range (i.e., maximum
209	range of $\delta^{15}N$) (see Supplementary Material Table S3). We then ran a principal component
210	analysis (PCA) to collapse down our eight isotopic metrics and four environmental metrics. This
211	generated three principal components for isotopic metrics, and three principal components for
212	environmental metrics used for further analysis (Table 1, see Supplementary Materials).
213	
214	Statistical analyses
215	We had four goals in our statistical analyses, namely examining whether: (1) isotopic niche at the
216	colony-level varied across years, (2) breeding stages and (3) inter-annual variation in isotopic
217	metrics predicted variation in key breeding parameters (Table S1), and (4) inter-annual variation
218	in environmental indices (winter NAO, spring NAO; pre-breeding Ta PC groups; Table S2)
219	predicted inter-annual variation in isotopic metrics.
220	First, to assess annual variation in $\delta^{15}N$ and $\delta^{13}C$ values at the colony-level, we ran a
221	MANCOVA (with $\delta^{15}N$ and $\delta^{13}C$ as dependent variables) using our 9-year dataset for common
222	eider females including year, breeding stage, the interaction between year and breeding stage,
223	body mass, and relative arrival date as independent variables. Since each isotope represents
224	different trophic dynamics (i.e., δ^{15} N: trophic position; δ^{13} C: spatial foraging), we followed the
225	MANCOVA with two separate one-way ANCOVAs for $\delta^{15}N$ and $\delta^{13}C$ to disentangle the effects

226 of our independent variables on each of our isotopic metrics (dependent variables). In the 227 ANCOVAs, we included year, breeding stage, the interaction between year and breeding stage, 228 body mass, and relative arrival date as independent variables. Secondly to examine finer-scale, 229 breeding-stage specific changes in foraging strategies *via* variation in δ^{13} C and δ^{15} N, we 230 conducted break-point analyses. This analysis identifies sudden and significant positive or 231 negative changes in the dataset (Muggeo 2003) through estimating breakpoints by iteratively 232 fitting a model with a linear predictor. For each iteration, a standard linear model is fitted, and 233 the breakpoint value is updated until convergence occurs. We performed break point analyses for both δ^{13} C and δ^{15} N separately including the delay before laying as an independent variable using 234 235 the Segmented R package (Muggeo 2003; R Core Team 2014). Thirdly, we ran an ANCOVA to 236 test whether mean inter-annual isotopic metrics (i.e., ISOPC1 (Spatial Foraging Breadth), 237 ISOPC2 (Niche Breadth), and ISOPC3 (Trophic Position) predicted variation in colony mean 238 arrival dates and breeding parameters by including colony mean breeding parameters as 239 dependent variables (i.e., breeding propensity, the delay between arrival at the colony and laying, 240 and relative lay date). Lastly, to determine whether inter-annual variation in isotopic metrics 241 could be predicted by inter-annual variation in environmental traits, we ran separate ANCOVAs 242 with the isotopic PC scores as our dependent variables (ISOPC1, ISOPC2, and ISOPC3) and the 243 environmental PC scores (ENVPC1 and ENVPC2) as our independent variables. All our analyses met the assumptions of a parametric test. All statistical tests were run using JMP 244 245 (Version 14.1.0 SAS) unless otherwise stated.

246

247 **Results**

248 Inter-annual and breeding stage variation in isotopic values

The MANCOVA analysis detected a significant interaction between year and breeding stage in predicting variation in female eider plasma δ^{15} N and δ^{13} C (Table 2; Figs. 1-3). ANCOVAs examining the two isotopes separately, revealed that only δ^{15} N showed significant year- and breeding-stage effects (with no year by stage interaction). Variation in δ^{13} C was negatively correlated with arrival date, with earlier arriving females having higher δ^{13} C values (Table 3).

254

255 *Fine-scale changes in isotopic values across breeding stages*

256 While δ^{15} N values were relatively consistent throughout the pre-laying period, breakpoint

analyses detected a significant increase approximately two days prior to laying (breakpoint

value: 1.8 ± 5.4 days, Fig. 3b). A breakpoint was also detected for δ^{13} C where values were

relatively consistent across the pre-laying period, until 7.4 days prior to laying. This occurred at

approximately the initiation of the rapid follicle growth (RFG) stage, when δ^{13} C values began

increasing significantly (breakpoint value: 7.4 ± 2.5 days, Fig. 3a).

262

263 Using isotopic metrics to predict breeding parameters

There was a significant negative relationship between ISOPC2 (Niche Breadth) and breeding propensity (one-way ANCOVA: $F_{1,7}=15.37$, p=0.01, w=0.60 Table 4). We found that females that foraged more inshore (higher δ^{13} C values) and within a narrower niche had a higher probability of breeding. However, we found no other significant relationships between colonylevel isotopic metrics and arrival or breeding parameters (Table 4).

269

270 *Climate and isotopic metrics*

271 We found a significant negative relationship between ENVPC2 (North Atlantic Oscillation

values) and ISOPC3 (Trophic Position) (ANCOVA, F_{1,1}=8.26, p=0.03, w=0), suggesting that

273 milder spring environmental conditions were associated with more variable trophic position.

However, we did not detect any other relationships between environmental variables and isotopicmetrics (Table 3).

276

277 Discussion

278 Using a 9-consecutive-year data set we asked whether information on annual- and reproductive 279 stage-based variation in isotopic niche dynamics predict whether a breeding colony of eiders has 280 the foraging flexibility respond to current and projected environmental change. Female eiders demonstrated significant variation in their δ^{15} N values across years and breeding stages 281 282 suggesting that either access to prey or selective choice to forage at certain trophic levels differs 283 across breeding stages. However, δ^{15} N values significantly increased 2-3 days prior to laying, 284 suggesting the increased energetic demand of ovarian follicle production during the rapid follicle 285 growth period may drive a change in trophic decisions and/or that internal nutrient mobilizations associated with egg formation occurred (Whiteman et al. 2020). We also detected an increase in 286 287 δ^{13} C as birds transitioned from the pre-breeding to the rapid follicle recruitment stage. This shift 288 could similarly be associated with diet shifts and/or internal isotopic processes associated with 289 nutrient mobilization to developing eggs. Assuming a dietary effect, this, combined with our 290 result indicating isotopic niche breadth (more inshore, narrow niche) positively predicted 291 breeding propensity, suggests that eiders may shift foraging locations inshore to respond to the 292 increased energetic demand of the reproductive organ load. We also found that isotopic niche 293 breadth (more inshore, narrow niche) positively predicted breeding propensity, although not 294 laying date. Further, we found that breeding birds tended to feed more benthically when 295 producing eggs. Despite these linkages to breeding investment, we detected significant

relationships between environmental conditions and trophic dynamics, but not spatial foraging indices (i.e., δ^{13} C). Our results suggest that common eiders may use flexible foraging strategies to overcome environmental constraints to invest in reproduction. Considering the rapid and substantial changes occurring in Arctic marine ecosystems, these findings provide a first step towards bridging key mechanistic gaps in determining how large-scale environmental processes proximately impact the foraging decisions and physiology of female eiders expected to influence colony-level resiliency.

303

304 *Links between isotopic niche variation and breeding investment decisions*

305 Given the energetic demands leading up to laying (Sénéchal et al. 2011; Hennin et al. 2015), we 306 predicted that shifts in δ^{15} N related to breeding stage would indicate adjustments in foraging strategies to the changing energetic demands of breeding. We found that $\delta^{15}N$ was indeed higher 307 308 during the pre-recruiting and RFG stages compared to the laying stage (Table 2; Fig. 2,3b). Interestingly, we did not find any changes in δ^{15} N either between pre-recruitment stage, despite 309 310 birds needing to gain significant body fat (i.e., body mass of 500-750g) to invest in breeding, and 311 initiation of rapid follicle growth (although see details below for changes in δ^{13} C values). However, we detected a significant breakpoint, and the highest $\delta^{15}N$ values, nearly two days 312 313 prior to laying (i.e., during RFG). This coincides with the most energetically demanding rapid 314 follicle growth (RFG) stage of reproduction (see Hennin et al. 2015). This suggests that females 315 may shift their prey consumption to match the energetic requirements of laying. We suggest that 316 combined, these results provide evidence that females make biologically relevant trophic 317 decisions to fuel the energetic costs of flight and diving while carrying the additional weight of 318 developing yolks and their supporting reproductive tissues (see Vézina and Williams 2002;

319 Vézina and Salvante 2010; reviewed in Williams 2012). We acknowledge however, that it is 320 plausible that isotopic effects of RFG may have led to these switches in δ^{15} N values. Previous 321 work on species relying on some degree of capital (i.e., stored) resources to fuel egg production 322 have indicated that δ^{15} N fractionation can increase in tandem with the increased energetic 323 demand associated with egg production via protein catabolism (Hobson et al. 1993; Vanderklift 324 & Ponsard 2003). It is therefore important to note that previous work at EBI has shown that pre-325 breeding eiders rely heavily on income-based resources to fuel egg production (Sénéchal et al. 326 2011) and only enter a fasting (protein catabolism) state during incubation (Bottitta et al 2003). 327 As such, we would expect minimal impact of fractionation on stable isotope values in plasma 328 due to fasting or rapid follicle growth in our focal birds. Nevertheless, we acknowledge that such 329 processes require further examination in Arctic-breeding female eiders.

330 We found that early arriving birds foraged closer to shore in agreement with our 331 predictions. This could be the result of birds needing to forage at the mouths of rivers as 332 freshwater inflow into the bay begins to open up small areas of open water prior to marine ice off 333 where birds can access resources (Jean-Gagnon et al. 2018). We also detected a significant breakpoint in δ^{13} C values in which females moved to forage at more inshore locations as they 334 335 transitioned into the RFG period. This suggests that when females are under significant energetic 336 demands at the onset of ovarian follicle recruitment, they may switch from foraging on offshore 337 benthic macroinvertebrates, to prey with lower digestive and foraging energetic costs, such as 338 amphipods (Gammarus spp.). This trophic switch may be driven by a variety of non-exclusive 339 reasons, including fueling the growth of reproductive organs and follicles. Amphipods are an 340 eider prey species found more inshore near the surface of the water (Sénéchal et al. 2011). They 341 require less metabolic energy and time to digest compared to hard-shelled benthic prey (e.g.,

blue mussels, sea urchins) (Zhang et al. 2019), while simultaneously reducing the energetic costs
of diving as females become more buoyant as they continuously accumulate needed fat stores for
reproduction (Sénéchal et al. 2011). Conserving fat stores for their upcoming incubation fast is
critical. Foraging on amphipods close to the colony may reduce the costs of commuting and
foraging, and may be an important energy-conserving strategy.

347 Interestingly, we found that an inshore, smaller isotopic niche, predicted higher breeding 348 propensity (Table 4; Fig. 1,3a). Previous studies have demonstrated that in years with later ice 349 breakup, females were less likely to breed (Jean-Gagnon et al. 2018). Additionally, previous 350 research at EBI has shown that unpredictable food shortages during the RFG stage reduce the 351 probability that an individual breeds, regardless of reproductive readiness (Legagneux et al. 352 2016), suggesting that reproductive decisions are highly responsive to environmentally driven 353 changes in resource availability typically related to ice cover and access to prey (see below). 354 Considering the short Arctic breeding season (Lepage et al. 2000) and the role that ice dynamics 355 play in impacting the decision to breed (Jean-Gagnon et al. 2018), the relationship between 356 breeding propensity and isotopic niche may only be evident in years with the greatest 357 environmental, and therefore, energetic constraints and may only impact individuals with lower 358 body condition.

359

360 *Lack of a relationship between environmental conditions and isotope values*

We predicted that in years with milder winters, milder springs, and higher overall ambient temperatures, there would be more open water and a wider variety and quantity of prey items available, resulting in a broader colony-based isotopic niche. In contrast, we only detected a significant, negative relationship between trophic position (ISOPC3) and pre-breeding NAO 365 (ENVPC2) (Table 3). In years with higher winter storm activity and milder springs at EBI (i.e., breeding grounds), females had a greater range and variance in δ^{15} N values on breeding grounds, 366 367 potentially related to accessing resources from multiple trophic levels. In years with harsh winter 368 conditions, mussel beds and other prey sources may be depleted due to the increased storm 369 activity on wintering grounds (Reusch and Chapman 1995). Consequently, eiders may employ a 370 more generalist foraging strategy upon arrival to breeding areas to compensate for diminished 371 abundance of preferred prey during winterexplaining the higher MNND values calculated during 372 these conditions. Similarly, we predicted that δ^{13} C would also be strongly related to 373 environmental indices, but this relationship was not observed (Table 3). This was surprising 374 considering that eiders are known to adjust their foraging areas based on extent of sea ice (Jean-375 Gagnon et al. 2018), and given the impacts of environmental conditions on regional sea-ice 376 dynamics near the nesting colony, although we are still learning about benthic-pelagic coupling 377 in the marine system. However, since variation in δ^{13} C is largely driven by input from primary 378 producers, it is possible that the lack of diversity of primary production around the nesting 379 colony in EBI, consistent with the circumpolar Arctic (Ramírez et al. 2017), limits our ability to detect a relationship between δ^{13} C and environmental conditions. Thus, the ability to apply bulk 380 381 stable isotopes in this context could be constrained by the biogeochemical processes of isotopic 382 incorporation (see Boecklen et al. 2011, Whiteman et al. 2021). Combining compound-specific 383 stable isotope analyses of individual amino acids and spatial tracking will provide future 384 opportunities to investigate colony-level resiliency to changes in prey availability predicted by 385 climate change.

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Table 1 - List of principal component correlation values based on plasma isotopic metrics
derived from bulk d15N and d13C values of female common eiders breeding at the East Bay
Island, Nunavut, Canada and environmental metrics (see Methods for details). Bold values
indicate statistically significant relationships at the alpha level of 0.05.

Group	Variable	ISOPC1 (Spatial Foraging)	ISOPC2 (Niche Breadth)	ISOPC3 (TP)
Isotopic	Y Range	0.20201	0.51061	-0.57323
	X Range	0.45855	0.13017	-0.12389
	MNND	-0.17996	0.40414	0.63032
	SEAc	0.47159	0.1755	0.08766
	$AVG\delta^{13}C$	0.29675	-0.58426	0.19752
	%CV δ^{13} C	-0.42799	0.08641	-0.13040
	AVG $\delta^{15}N$	-0.35635	0.21296	0.01793
	$CV \delta^{15} N$	0.30971	0.36609	0.44129
Eigenvalue		3.7445	1.4859	1.2791
Cum. Percent		46.806	18.574	15.989
Group	Variable	ENVPC1 (Spring)	ENVPC2 (NAO)	
Environment	S NAO	0.6001	-0.53296	
	W NAO	0.35475	0.84569	
	Та	-0.71696	-0.02764	
Eigenvalue		1.1305	1.0078	
Cum. Percent		37.685	33.592	

706	Table 2 - Summary of MANCOVA and ANCOVA analyses examining predictors of variance in
707	plasma δ^{13} C and δ^{15} N values in female common eiders breeding at the East Bay Island, Nunavut,
708	Canada. Bold values indicate statistically significant relationships.
709	

Analysis	Variable	F	df	р
MANCOVA (δ^{15} N and δ^{13} C)	Full Model	4.35	74, 1198	0.0001
	Year	5.22	16, 1198	0.0001
	Breeding stage	2.97	2,600	0.03
	Year*Breeding stage	1.41	48, 1198	0.04
	Relative arrival date	19.50	2, 599	0.0001
	Body Mass	0.50	2, 599	0.61
ANCOVA (δ^{15} N)	Full Model	5.92	37, 600	0.0001
	Year	8.67	8, 8	0.0001
	Breeding stage	2.96	3, 3	0.03
	Year*Breeding stage	1.31	24, 24	0.15
	Relative arrival date	1.41	1, 1	0.23
	Body Mass	0.13	1, 1	0.71
ANCOVA (δ^{13} C)	Full Model	2.82	37, 600	0.0001
	Year	1.15	8, 8	0.33
	Breeding stage	0.56	3, 3	0.64
	Year*Breeding stage	1.34	24, 24	0.13
	Relative arrival date	27.00	1, 1	0.0001
	Body Mass	0.51	1, 1	0.47

- 711 **Table 3** Summary of ANCOVA analyses using environmental metrics (ENVPC1 (Spring
- 712 Weather); and ENVPC2 (NAO)) to predict variation in plasma isotopic metrics (ISOPC1 (Spatial
- Foraging Location); ISOPC2 (Niche Breadth); and ISOPC3 (Trophic Position)) in female
- 714 common eiders breeding at the East Bay Island, Nunavut, Canada. Bold values indicate
- 715 statistically significant relationships.
- 716

	ISOPC1				ISOPC2			ISOPC3			
Variable	R ²	F2,6	р	R ²	F2,6	р	R ²	F _{2,6}	р		
Full Model	0.02	0.07	0.93	0.10	0.35	0.72	0.62	4.88	0.06*		
ENVPC1	-	0.12	0.74	-	0.17	0.69	-	1.50	0.27		
ENVPC2	-	0.03	0.86	-	0.53	0.49	-	8.26	0.03		

717 *Marginal significance (see Results).

718 **Table 4** - Summary of ANCOVA analyses between breeding parameters (relative arrival date

719 (RAD); percent breeding propensity (BP); delay before laying (delay); relative lay date (RLD))

and isotopic metrics (ISOPC1 (Spatial Foraging Location); ISOPC2 (Niche Breadth); and

721 ISOPC3 (Trophic Position)).

722

	RAD			BP			Delay				RLD		
PC	R ²	F _{1,1}	р	R ²	F _{1,1}	р	R ²	F _{1,1}	р	R ²	F _{1,1}	р	
Model	0.3	0.84	0.53	0.7	4.48	0.07*	0	0.02	1.00	0.2	0.51	0.70	
ISO1	-	0.16	0.70	-	0.62	0.47	-	0.03	0.87	-	0.01	0.91	
ISO2	-	1.57	0.27	-	12.67	0.02	-	0.00	0.97	-	1.25	0.31	
ISO3	-	0.79	0.42	-	0.15	0.72	-	0.02	0.91	-	0.25	0.64	

723 *Marginal significance (see Results).

Figure 1 - Inter-annual variation in plasma isotopic niche of eiders nesting at East Bay Island,
NU, Canada. Each colour indicates a different year. Ellipses represent 40% of the individuals'
isotopic values that comprise the sampled population within each year. Ellipses are used to
represent the placement of birds within each year in isotopic space and compare placement
among years.

729

Figure 2 - Breeding stage-related variation in plasma isotopic niche of eiders nesting at East Bay
Island, NU, Canada. PR (green line; pre-recruiting), shows 40% of the individuals which started
laying at least 8 days post-capture; RFG (blue line; rapid follicle growth), is characterized by
birds that are delayed between 8-1 days before breeding and Lay (red line; laying or incubating)
is the 40% ellipse of birds during laying or incubation the same day of capture.
Figure 3 - Variation in plasma δ¹³C (A), and δ¹⁵N (B) values across breeding stages of female
eiders nesting on East Bay Island. Values are represented as mean ± SEM provided for each day

738during pre-breeding (black circle), rapid follicle growth (black square), and laying (black

triangle). The size of the data points represents the sample size. Dashed, vertical lines represent

statistically significant change in δ^{13} C and δ^{15} N values detected from breakpoint analyses.













