1	Fall composition of storage lipids is associated to the overwintering strategy of Daphnia
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29 Abstract

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31 Diapause, using dormant eggs, is a strategy some zooplankton use to avoid winter 32 months of persistent low temperatures and low food availability. However, reports of 33 active zooplankton under the ice indicate other strategies also exist. This study aimed at 34 evaluating whether fall composition of storage lipids is different between diapausing and 35 actively overwintering Daphnia. We assessed the quantity of storage lipids and fatty acid 36 composition of Daphnia species along with fatty acid content of seston in six boreal, 37 alpine and subarctic lakes at the onset of winter, and evaluated the association of storage 38 lipids to Daphnia overwintering strategy. We found that active overwintering Daphnia had 39 >55% of body fat and the highest fatty acid concentrations. Polyunsaturated fatty acids, 40 especially stearidonic acid (18:4n3; SDA) and high ratios of n3:n6 were preferentially 41 retained in active overwintering Daphnia compared to those that entered diapause. The 42 Daphnia fatty acid compositions were independent of that of the seston diet, indicating 43 Daphnia adjusted their storage lipids according to the physiological requirements of a 44 given overwintering strategy. The occurrence of the active overwintering strategy has 45 consequences to zooplankton community structure, and can have important implications 46 for the transfer of high quality energy in higher trophic levels. 47 48 **Keywords:** freshwater, winter, fatty acids, ice, life history strategies, zooplankton, SDA 49 50 51 52

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55 Abbreviations

56	FA	Fatty acid(s)
57	FAME	Fatty acid methyl esters
58	EPA	Eicosapentaenoic, 20:5n3
59	MUFA	Monounsaturated fatty acid(s)
60	PUFA	Polyunsaturated fatty acid(s)
61	SDA	Stearidonic acid, 18:4n3
62	SFA	Saturated fatty acid(s)

63 Introduction

64

65 Life history strategies of organisms are often determined by environmental cues, such as 66 temperature and food availability [1–3]. This means that organisms can adjust their 67 physiology and reproductive cycles to changes in their environment. One of the best 68 documented life-history adaptations to adverse changes in the environment is the 69 production of resting eggs, ephippia, in Daphnia (Cladocera) [4, 5]. Increases in 70 population density and decreases in temperature, light and food supply are some of the 71 most common triggers that induce resting egg production, and characterize lakes in late 72 summer – fall. It is therefore commonly accepted that cladocerans prepare for winter by 73 producing ephippia and are absent from the water column during the winter months with 74 persistent low temperatures and limited food. However, there is increasing awareness of 75 the presence of active zooplankton throughout the ice cover period [6-8], indicating that 76 at least for certain species there exist also other adaptation mechanisms to overwinter. 77 While most of the historic research has focused on environmental conditions initiating 78 overwintering diapause, especially ephippia production in Daphnia [9], much less is 79 known about physiological adaptations of cladocerans that overwinter actively. 80 81 An effective physiological overwintering mechanism known for fish and marine

82 zooplankton is to accumulate storage fat [5, 10, 11]. Storage fats and their fatty acids 83 (FA) provide high-density energy reserves used to maintain metabolic function and 84 reproduction during low food supply [12–14]. Studies on freshwater copepods indicate 85 that both the quantity of lipid reserves [15] and the high overall percentage of 86 polyunsaturated fatty acids (PUFA) [16] are common characteristics in the fall for 87 copepods that overwinter actively suggesting that there could be differences in lipid 88 accumulation depending on whether cladocerans enter diapause or overwinter actively. 89 The link between storage fats and FA, and zooplankton life history strategies has 90 however not been explored in detail, especially in freshwater systems. 91 92 Typically the function of particular fatty acids is determined by the degree of saturation, 93 and fatty acids can be grouped accordingly. Saturated and monounsaturated fatty acids 94 (SFA and MUFA) tend to be used for storage fats [17]. PUFA are particularly important 95 for reproduction and early development [17], and are used to compensate for the loss of

96 cell membrane flexibility in cold temperatures [18]. Further, consumers differ in their

97 composition and thus in their requirement for specific FA [19]. The growth and 98 reproduction of cladocerans has e.g. been strongly affected by the EPA 99 (eicosapentaenoic, 20:5n3) content [20] but whether EPA or any other FA is linked or 100 required to the active overwintering Daphnia is not known. Little is also known about how 101 composition or relative differences between certain FAs account for any species specific 102 or individual metabolic requirement, such as an organism's reproduction, thermal 103 compensation, or life-history stage. Generally a higher quantity of PUFA and a higher 104 ratio between omega-3 (n3) and omega-6 (n6) PUFA are favorable for all organisms, 105 however the optimum dietary ratio of n3:n6 seems to be highly species specific [21] and 106 very little is known about changes in seasonal requirements for these FA ratios.

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108 As the source of zooplankton lipids is diet, any seasonal accumulation of lipids in 109 zooplankton is linked to food sources. Seasonal phytoplankton food availability has been 110 reported to be the main driver for organisms to store FA [3, 17] and there is evidence that 111 freshwater copepods use fall phytoplankton to build up the fats that allow for reproduction 112 under the ice [8, 16]. Changes in phytoplankton community and FA composition are to 113 some extent mirrored in the zooplankton FA composition [22, 23]. However, in fall, a 114 simple reflection of the bulk phytoplankton community, where many taxa may be 115 senescing, would not be a winning strategy to survive the winter months under the ice. 116 Rather, a selective accumulation of FAs could be used as an energy source during 117 winter. Given the predominance of ice cover in northern latitudes, it is important to 118 understand the environmental and physiological mechanisms determining life history 119 strategies. Ultimately the presence or absence of an active zooplankton in a lake affects 120 the lake's community structure and ecological functioning, such as winter food web 121 dynamics [24, 25].

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123 In this study we investigate the link between winter life history strategy of *Daphnia* and 124 the quantity of their lipids and FA composition. Daphnia are well adapted to unstable 125 habitats due to short life cycle, and have a wide phenotypic plasticity [6, 26]. These 126 characteristics make Daphnia an ideal genus to compare the physiology and 127 overwintering strategies among different populations in different parts of their 128 environmental range. Our primary goal of this study was to compare the FA composition 129 of Daphnia from boreal, alpine and subarctic lakes at the onset of winter. In this wide 130 geographical range, different species compositions of Daphnia and different

131 overwintering strategies emerged, including Daphnia that entered diapause for winter, 132 Daphnia that remained active under the ice, and lakes where both overwintering 133 strategies were present. We hypothesized that in lakes where *Daphnia* overwinter 134 actively, they will store lipids during the fall, particularly PUFA, in order to compensate for 135 low temperature and limited food availability under the ice. Since the dormant strategy 136 simply avoids the harsh winter conditions, we assumed that *Daphnia*, which enter 137 dormancy will not invest in storing lipids. We include further discussions on the 138 evolutionary implications of winter strategies, and the composition and physiology of 139 storage lipids.

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141 Materials and methods

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143 Study sites and sample collection

144 We sampled Daphnia from six lakes across three geographically distinct regions in 2009-145 2011 (Table 1). The *Daphnia* were the dominant zooplankton species in three of the sites 146 (Big Rock, Malla South and Mekkojärvi) while in the other sites they contributed to a 147 zooplankton community dominated by copepods. Due to the wide geographical range the 148 dataset included five different Daphnia species and many lakes had different Daphnia 149 assemblages (Table 1). In the text they are collectively referred to as *Daphnia*. The lakes 150 were divided to three categories based on the occurrence of *Daphnia* in winter. The lakes 151 Malla South and Saanajärvi with no record of ephippia-carrying *Daphnia* in earlier 152 zooplankton samples or paleolimnological studies (e.g. [27]) were considered lakes with 153 actively overwintering Daphnia. The lakes where all Daphnia entered diapause as 154 ephippia due to freezing solid (Big Rock) or winter anoxia (Mekkojärvi) were called 155 diapause lakes. The lakes Lunzersee and Simoncouche, where both ephippia and active 156 individuals were present, were considered lakes with both overwintering strategies. 157 Daphnia were present in winter in the alpine Lunzersee (0.0005 ind L⁻¹, n=1, December) 158 with the shortest ice cover period of three months, in boreal Simoncouche (2.46±5.5 ind 159 L^{-1} , n=12, December-April) with six months of ice and the two subarctic lakes Malla South 160 and Saanajärvi (0.02±0.01 ind L⁻¹, n=8, November-May) with up to nine months of ice. 161 The winter abundance was up to two fold lower than in summer, however, 162 parthenogenetically reproducing individuals were found in both lakes in winter. 163

164 From these six sites we compared the percent body fat and FA composition of adult 165 Daphnia at the onset of winter. Considering the wide latitudinal variation in our study sites 166 this period was defined as maximum the two months before ice cover for each water 167 body (hereafter referred to as fall). Temperature in all sites was close to 10°C during the 168 sampling and decreasing from the maximum summer values. *Daphnia* were collected by 169 vertical tows of a 50 µm mesh net at the deepest point of the lake site, or by horizontal 170 pulls in the shallow pond Big Rock. The *Daphnia* were hand sorted, aiming for at least 171 0.5-1.5 mg dry weight (DW), when possible triplicates were made. Seston, representing 172 the dietary source of FA available for *Daphnia*, was taken from a depth-integrated water 173 sample that was pre-filtered through a 50 µm net to remove larger organisms, then 174 filtered onto a GF/F filter using 1-3 L per triplicate. All samples were freeze dried and 175 stored in -80 °C until lipid and FA analysis.

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177 Fatty acid analyses

178 Fatty acids were analysed using a three-step extraction-methylation protocol [27]. Freeze 179 dried and weighed Daphnia and seston samples were extracted using a chloroform-180 methanol wash cycle and concentration of extracted lipids determined by gravimetry. The 181 body fat percentage was calculated as a function of extracted lipid weight and Daphnia 182 DW. The extracted lipids were methylated using toluene and sulphuric acid-methanol and 183 then solubilized in hexane. Fatty acids (C14–C24) were identified as fatty acid methyl 184 esters (FAME) using a gas chromatograph equipped with a Supelco™ SP-2560 column 185 (100 m, 25 mm i.d., 0.2 µm film thickness) and a flame ionization detector (TRACE GC-186 FID THERMOTM). Total fatty acid concentrations (Σ FAME) were calculated using 187 calibration curves based on known external standard concentrations and are reported as 188 μ g FAME per mg carbon weight (μ g FA mgC⁻¹). Fatty acid concentrations were calculated 189 using calibration curves of known standard concentrations. Daphnia dry weights were 190 converted to carbon (C) weight using a Daphnia summer average of 42% C and 15% for 191 seston that were available from mass spectrometry (Thermo Finnigan DELTAplus 192 Advantage) and stable isotope analyses (Mariash unpubl). Only the FAME greater than 193 0.1% were used in the analysis. All FAME results are presented as a proportion of the 194 total FAME concentration in moles. Individual FA are summarized in results into general 195 lipid classes of saturated- (SFA), monounsaturated- (MUFA), polyunsaturated- (PUFA) 196 fatty acids or as ratios of the sum of omega- 3 to omega-6 FAME (n3:n6). 197

198 Statistical analysis

199 Differences in seston and Daphnia fatty acid composition (data normalized, Euclidean 200 distances), and Daphnia fall FAME characteristics (SFAME, %PUFA, %SFA, %SDA, 201 SFA:PUFA and Σ n3:n6) and body fat levels among strategies (active, diapause, both 202 strategies) and lakes (Malla South, Saanajärvi, Simoncouche, Lunzersee, Big Rock, 203 Mekkojärvi) were tested with a 2 factor (lake nested in strategy, strategy fixed and lake 204 random) Permutational (Multivariate for fatty acid compositions) Analysis of Variance 205 (PERMANOVA with 999 permutations). While comparing FAME characteristic values can 206 be done with classical ANOVA, permutational ANOVA was preferred as it did not require 207 observations fitting the normality assumptions (see [29]). The nested design was used to 208 control for variation among lakes within strategy. When only a restricted number of permutations were possible, Monte Carlo p-values were used. Non-metric multi-209 210 dimensional scaling (nMDS) was used to visualize Daphnia FA composition among 211 factors. Linear regressions were carried out between Daphnia and seston FAME 212 variables that were significantly different in Daphnia in different overwintering strategies 213 (Σ FAME, %PUFA, %SDA, Σ n3:n6) to test for potential diet influence on *Daphnia* FAME 214 composition. Pair-wise comparison tests among strategies were carried out for Daphnia 215 FAME composition, total FAME, PUFA, SDA and n3:n6 in PERMANOVA with adjusted 216 significance level method from Holm [30]. All analyses were conducted using 217 PRIMER+PERMANOVA version 7.0.9. A significance level α = 0.05 was used for all

statistical tests.

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220 Results

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222 The Daphnia total lipid amount and FA profiles were different for Daphnia with different 223 strategies (Table 2, Table 3, Fig. 1). In the lakes with actively overwintering Daphnia, the 224 percent body fat was highest (58%) although not statistically different from the other 225 strategies (Table 3a). In the lakes with diapause or mixed overwintering strategy Daphnia 226 had lower body fat ($\sim 30\%$) (Table 2a). Total FAME concentrations (mean ±SD) were 227 statistically higher for active *Daphnia*, than for the diapause strategy (Table 2b, Table 3b, 228 Fig. 2). The *Daphnia* FA compositions among the strategies were marginally different (p =229 0.068) from each other (Table 3c; Fig 1). When the lake factor was not considered all 230 strategies were highly statistically different from each other (p < 0.003 for all multiple pair-231 wise comparisons) and sample scores from the active and diapausing populations were

232 furthest away from each other (Fig. 1). The difference among differently overwintering 233 Daphnia FA compositions was largely due to differences in their total PUFA, stearidonic 234 acid (18:4n3; SDA) and the n3:n6 ratio (Fig. 2). The Daphnia PUFA were marginally 235 different between strategies (p = 0.061; Table 3d) with higher values in actively 236 overwintering Daphnia (Table 2c). SDA was significantly different among strategies 237 (Table 3f, Fig. 2) and more than double in active winter Daphnia (26% of total FAME) 238 compared to Daphnia with mixed strategies (12%) and 9-fold higher compared to the 239 diapausing Daphnia (4%) (Table 2c). The SDA results were further reflected in the n3:n6 240 differences among strategies with significantly higher n3:n6 ratios for the active strategy 241 (Table 2d, Table 3h). In summary, the main lipid parameters that were most different 242 between strategies were total FAME concentration, SDA, PUFA, and n3:n6 FA ratio, as 243 illustrated in Figure 2. Further, the active and diapausing populations differed in species. 244 Lakes with active strategy had only *D. umbra* and lakes with diapause had only *D.* 245 longispina, which differed in their FA composition (SI Table 1). 246 247 FAME concentrations in seston were an order of magnitude lower than Daphnia FAME 248 concentrations (SI Table 2). Seston had high SFA:PUFA ratios, indicating that SFA 249 dominated the seston. Of all FAME in seston, between 41 to 89% were SFA. Of the 250 PUFA, seston generally contained very small amounts of 18:3n6 (GLA), 20:5n3 (EPA), 251 22:6n3 (DHA), and notably 18:4n3 (SDA) was less than 2%. Seston FA composition was 252 not different among strategies (Table 3i). Regression analyses revealed that variation in 253 seston FA composition (Σ FAME, %PUFA, %SDA, Σ n3:n6) across the six lakes did not 254 explain a significant amount of corresponding variation in the FA composition of any Daphnia community ($R^2 < 0.20$ and p > 0.10 for all comparisons). While average 255 256 concentrations of individual FAs in the Daphnia varied substantially among overwintering 257 strategies, FA concentrations in seston were more uniform and did not reflect these 258 differences. The patterns in FA composition and total fatty acid concentrations in the 259 zooplankton appeared more closely related to zooplankton overwintering strategy than 260 seston fatty acid composition. 261

262 **Discussion**

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We compared the quantity of storage lipids and FA composition of *Daphnia* at the onset of winter from lakes where there is an active winter community, lakes where the *Daphnia*

266 enter diapause for winter and from lakes where the Daphnia use both strategies but are 267 dominated by the diapausing strategy. Our results suggest that fatty acids may have an 268 important role for *Daphnia* that prepare to overwinter actively and are in accordance with 269 earlier observations from cladocerans and calanoid copepods that have demonstrated a 270 high accumulation of lipids and PUFA in fall [5, 15, 16]. Storing fat could be a 271 physiological mechanism zooplankton use to adapt to life under the ice, and therefore 272 can potentially be used as a physiological indicator for zooplankton's winter survival. 273 274 The active winter *Daphnia* stored more FA, both in % body fat and FAME concentration,

275 than the other two strategies. Specifically PUFA, SDA (18:4n3), and high ratios of n3:n6 276 were preferentially retained in active overwintering Daphnia than in Daphnia using the 277 other two strategies. While cladocerans should have fairly constant n3:n6 FA ratios, with 278 an average of 3 [19, 31, 32], our results show that there were contrasting n3:n6 ratios 279 between winter life history strategies. The active Daphnia, especially in the subarctic, had 280 more than double (8.25) the average, while the diapausing *Daphnia* were below (2.0) the 281 n3:n6 average value of three. High proportions of SDA mostly drive the n3:n6 ratios in our 282 results, highlighting that SDA could have an important role in physiological functions for 283 Daphnia winter life history strategy.

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285 We also considered the alternative explanation that the patterns in FA among Daphnia 286 may stem from differences in basal resources [33] rather than the overwintering strategy. 287 For example, changes in FA among *Daphnia* in boreal lakes were attributed to changes 288 in resource assimilation between summer and fall diets [34]. However, our results show 289 that the composition of the *Daphnia* FA indicated a preferential retention or accumulation 290 of specific FA by Daphnia, rather than a direct reflection of availability of specific FA in 291 seston during the fall. Further, as winter seston diet is more scarce in subarctic than in 292 boreal lakes, one would expect diapause to be the dominant strategy at high-latitude 293 lakes, which was not the case here. The *Daphnia* entered diapause only when 294 environmental constraints such as loss of habitat (drying, freezing solid, anoxia) 295 prevented the active stage. While the limited number of lakes in this study does not allow 296 for more accurate testing for the region's or ice cover's role on the strategy, our data 297 shows that Daphnia overwinter actively in subarctic lakes even when there is 9 months of 298 ice cover.

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301 Other factors known to affect life history strategies are species type [35], environmental 302 stressors like temperature [2], and/or lake size and depth [36, 37]. Our study included five 303 different Daphnia species and it is known that different species differ in their fatty acid 304 composition and allocation of lipids [38, 39]. Lakes with active winter Daphnia had only D. 305 umbra, a species that is not known to occur in temperate lakes, and both lakes with 306 diapause had only *D. longispina, a species* common in the boreal region. It is possible 307 that lakes only support populations of species that are matched to the environment in 308 terms of their physiology, including fatty acid signatures. Hence, it could be that not all 309 Daphnia species are able to regulate their storage fats and lipid composition to the same 310 extent. The species that do not possess this ability could be geographically restricted to 311 lakes where open water period is warm and long enough to allow for the emergence from 312 ephippium and completing a life cycle before environmental conditions turn adverse 313 again. Subsequently, species not capable for lipid accumulation should be absent from 314 perennially cold arctic lakes, which could contribute to explaining the low species number 315 of zooplankton in arctic lakes [40]. However, there exists evidence that lipid allocation is 316 not only a species-specific trait but also determined by environmental conditions. Based 317 on the number of lipid droplets in D. umbra, Larsson and Wathne [5] were able to 318 separate the ephippia producing individuals with less lipid droplets from individuals that 319 accumulated large amounts of lipids to survive longer. Nevertheless, regardless of 320 whether overwintering actively is an intrinsic physiological adaptation of only certain 321 species or a more common response of *Daphnia* genus to seasonally changing 322 environmental conditions, the results here are in line with other studies that have shown 323 that zooplankton which overwinter actively accumulate lipids for the subsequent use 324 during the months when diet is scarce [8].

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326 As observed also in other studies, both strategies can coexist in the same lake [5, 36, 327 37]. Although it would be interesting to separate the different life history strategies within 328 the same lake, there is no way of visually separating between individuals that will 329 overwinter actively versus those that will make ephippia. Even at a population level, it is 330 difficult to discern the overwintering strategy. It is however, noteworthy that in lakes 331 where both strategies existed, the majority of Daphnia entered diapause based on the 332 low numbers of active *Daphnia* in water column under the ice and the high number of 333 ephippia in fall. The more similar fatty acid composition between diapausing and mixed

strategies in fall further suggests that most *Daphnia* in the mixed strategy lakes entered
diapause, possibly because they did not possess physiological means to accumulate
storage fats. FA composition between active, diapausing, and both strategies from
different lakes and different species, in this study suggests that lipid accumulation is an
important factor for *Daphnia* winter strategies, however, further experimental approach
and tests would need to be used to explicitly determine this.

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341 The ecology of the subarctic and arctic region may provide a framework to understand 342 how the active winter strategy evolved. The intense light regime in late spring at higher 343 latitudes promotes a thriving phytoplankton community even under the ice [7, 41, 42]. 344 creating favorable growing conditions for *Daphnia*. At this time ephippia would not yet be 345 primed for emergence as water temperatures are still too low $(3 - 5^{\circ}C)$ and would not be 346 able to benefit from the spring phytoplankton, as the already active community would. 347 Further, the high levels of the polyunsaturated SDA and n3 found in the active Daphnia 348 can help to combat the low spring temperatures [43]. Active winter Daphnia and their 349 parthenogenetic young are able to exploit the early spring phytoplankton, and are 350 typically larger in body size [44] giving them a decisive competitive advantage over the 351 resting egg community, made possible by energy reserves stored as > 55% body fat as 352 suggested by our study. In addition, the interaction between low temperature and 353 starvation during winter months could effectively help to extend winter survival by 354 lowering metabolism [45]. When the open water season is a major part of the year, 355 Daphnia have time to feed and regenerate within the longer productive period, and 356 ephippia have the flexibility to postpone emergence until spring conditions improve. 357

358 The evolutionary advantages of reproducing even in times of starvation are, however, 359 weighed against living longer but with increased probability of death from senescence 360 [46]. Complete abandonment of ephippia would be risky as a single season of low 361 survival and failed reproduction could eliminate an entire population. Further, the 362 diapause strategy maintains genetic diversity and ensures that a population may persist 363 during potentially fatal conditions [4]. The most profitable strategy would be to produce 364 many parthenogenetic offspring in spring plus add to the ephippia bank for insurance [1]. 365 We found the two winter life histories across the regions studied although a larger dataset 366 would be needed to confirm the occurrence and potential dominance of this strategy at a 367 global scale.

369 The occurrence of different overwintering strategies has consequences to the plankton 370 community structure and food web dynamics. Not only is the seasonal accumulation of 371 PUFA in cladocerans important to the accumulation of quality energy to higher trophic 372 levels [47], but the actual presence of winter *Daphnia* is important to winter-feeding fish 373 ([24] and ref therein) and can affect the composition of the spring phytoplankton 374 community [48]. Our results show that Daphnia using the diapause strategy do not need 375 to invest in storing a large reserve of lipids, having only half as much body fat in the fall 376 compared to the Daphnia remaining active. Therefore both population and community-377 level dynamics are affected by the life history strategies of Daphnia. Further, the 378 physiological adaptations and life history strategies zooplankton use on an annual scale 379 may provide insight into the degree of population plasticity in response to a changing 380 climate. As temperatures continue to increase around the globe, earlier spring ice off or 381 later fall ice-on will begin to affect the environmental cues (temperature and photoperiod) 382 for diapause stages [35, 49]. A longer ice-free period will change the phytoplankton 383 community and production, and could lead to changes of storage fat accumulation for 384 zooplankton.

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386 Acknowledgments

We are grateful to Kilpisjärvi and Simoncouche biological stations for logistical support.
We thank Martin Kainz and Jorge Watzke at the Wasserkluster Lunz for technical support
during fatty acid analysis and two anonymous reviewers for their constructive comments
that improved the paper. Core funding was provided by Academy of Finland with grants
19205 and 140775 to MR.

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393 References

- Thackeray SJ, Henrys P a., Jones ID, Feuchtmayr H (2012) Eight decades of
 phenological change for a freshwater cladoceran: what are the consequences of
 our definition of seasonal timing? Freshw Biol 57:345–359. doi: 10.1111/j.1365 2427.2011.02614.x
- Chen C, Folt C. (1996) Consequences of fall warming for zooplankton
 overwintering success. Limnol Oceanogr 41:1077–1086.
- 4003.Hagen W, Auel H (2001) Seasonal adaptations and the role of lipids in oceanic401zooplankton. Zoology (Jena) 104:313–26. doi: 10.1078/0944-2006-00037
- 4. Lampert W, Lampert KP, Larsson P (2010) Coexisting overwintering strategies in *Daphnia pulex*: A test of genetic differences and growth responses. Limnol
 Oceanogr 55:1893–1900. doi: 10.4319/lo.2010.55.5.1893
- 405 5. Larsson P, Wathne I (2006) Swim or rest during the winter what is best for an
 406 alpine daphnid? Arch für Hydrobiol 167:265–280. doi: 10.1127/0003407 9136/2006/0167-0265
- 4086.Slusarczyk M (2009) Extended lifespan traded for diapause in Daphnia. Freshw409Biol 54:2252–2262. doi: 10.1111/j.1365-2427.2009.02256.x
- 410 7. Rautio M, Mariash H, Forsström L (2011) Seasonal shifts between autochthonous
 411 and allochthonous carbon contributions to zooplankton diets in a subarctic lake.
 412 Limnol Oceanogr 56:1513–1524. doi: 10.4319/lo.2011.56.4.1513
- 8. Schneider T, Grosbois G, Vincent WF, Rautio M (2016) Carotenoid accumulation
 in copepods is related to lipid metabolism and reproduction rather than to UVprotection. Limnol Ocean. doi: 10.1002/lno.10283
- 416 9. Carvalho GR, Hughes RN, Animal S (1983) The effect of food availability , female
 417 culture-density and photoperiod on ephippia production in *Daphnia magna* Straus
 418 (Crustacea: Cladocera). Freshw Biol 13:37–46. doi: 10.1111/j.1365419 2427.1983.tb00655.x
- 420 10. Griffiths D, Kirkwood RC (1995) Seasonal variation in growth, mortality and fat
 421 stores of roach and perch in Lough Neagh, Northern Ireland. J Fish Biol 47:537–
 422 554.
- Hagen W, Van Vleet E, Kattner G (1996) Seasonal lipid storage as overwintering
 strategy of Antarctic krill. Mar Ecol Prog Ser 134:85–89. doi: 10.3354/meps134085
- 425 12. Tessier AJ, Goulden CE (1982) Estimating Food Limitation in Cladoceran
 426 Populations. Limnol Oceanogr 27:707–717.
- Reznick DN, Braun B (1987) Fat cycling in the mosquitofish (*Gambusia affinis*): fat
 storage as a reproductive adaptation. Oecologia 73:401–413. doi:
 10.1007/BF00385257
- 430 14. Lee RF, Hagen W, Kattner G (2006) Lipid storage in marine zooplankton. Mar Ecol
 431 Prog Ser 307:273–306.
- 432 15. van der Meeren T, Olsen RE, Hamre K, Fyhn HJ (2008) Biochemical composition
 433 of copepods for evaluation of feed quality in production of juvenile marine fish.
 434 Aquaculture 274:375–397. doi: 10.1016/j.aquaculture.2007.11.041

435 436 437	16.	Hiltunen M, Strandberg U, Keinänen M, et al (2014) Distinctive Lipid Composition of the Copepod Limnocalanus macrurus with a High Abundance of Polyunsaturated Fatty Acids. Lipids 919–932. doi: 10.1007/s11745-014-3933-4
438 439	17.	Kattner G, Hagen W, Lee RF, et al (2007) Perspectives on marine zooplankton lipids. Can J Fish Aquat Sci 64:1628–1639. doi: 10.1139/F07-122
440 441 442	18.	Guschina IA, Harwood JL (2009) Algal lipids and effect of the environment on their biochemistry. In: Arts MT, Brett MT, Kainz MJ (eds) Lipids Aquat. Ecosyst. Springer, New York, pp 1–24
443 444 445	19.	Persson J, Vrede T (2006) Polyunsaturated fatty acids in zooplankton: variation due to taxonomy and trophic position. Freshw Biol 51:887–900. doi: 10.1111/j.1365-2427.2006.01540.x
446 447 448	20.	Müller-Navarra DC, Brett MT, Liston a M, Goldman CR (2000) A highly unsaturated fatty acid predicts carbon transfer between primary producers and consumers. Nature 403:74–7. doi: 10.1038/47469
449 450 451	21.	Yang XW, Dick TA (1994) Arctic Char (<i>Salvelinus Alpinus</i>) and Rainbow Trout (<i>Oncorhynchus Mykiss</i>) Differ in Their Growth and Lipid Metabolism in Response to Dietary Polyunsaturated Fatty Acids. Can J Fish Aquat Sci 51:1391–1400.
452 453 454	22.	Burns CW, Brett MT, Schallenberg M (2011) A comparison of the trophic transfer of fatty acids in freshwater plankton by cladocerans and calanoid copepods. Freshw Biol 56:889–903. doi: 10.1111/j.1365-2427.2010.02534.x
455 456 457	23.	Taipale SJ, Kainz MJ, Brett MT (2011) Diet-switching experiments show rapid accumulation and preferential retention of highly unsaturated fatty acids in Daphnia. Oikos 120:1674–1682. doi: 10.1111/j.1600-0706.2011.19415.x
458 459 460	24.	Eloranta AP, Mariash HL, Rautio M, Power M (2013) Lipid-rich zooplankton subsidise the winter diet of benthivorous Arctic charr (<i>Salvelinus alpinus</i>) in a subarctic lake. Freshw Biol 58:2541–2554. doi: 10.1111/fwb.12231
461 462 463	25.	Hampton SE, Moore M V, Ozersky T, et al (2015) Heating up a cold subject : prospects for under-ice plankton research in lakes. J Plankton Res 0:1–8. doi: 10.1093/plankt/fbv002
464 465	26.	Dufresne F, Hebert PDN (1995) Polyploidy and clonal diversity in an arctic cladoceran. Heredity (Edinb) 75:45–53.
466 467 468	27.	Heissernberger M, Watzke J, Kainz MJ (2010) Effect of nutrition on fatty acid profiles of riverine, lacustine, and aquaculture-raised salmonids of pre-alpine habitats. Hydrobiology 650:234–254.
469 470 471	28.	Rautio M, Sorvari S, Korhola A (2000) Diatom and crustacean zooplankton communities , their seasonal variability and representation in the sediments of subarctic Lake Saanajärvi. Methods 59:81–96.
472 473 474 475	29.	Fairclough D V., Clarke KR, Valesini FJ, Potter IC (2008) Habitat partitioning by five congeneric and abundant Choerodon species (Labridae) in a large subtropical marine embayment. Estuar Coast Shelf Sci 77:446–456. doi: 10.1016/j.ecss.2007.10.004
476 477	30.	Holm S (1979) A Simple Sequentially Rejective Multiple Test Procedure. Scand J Stat 6:65–70.

478 479 480	31.	Ahlgren G, Vrede T, Geodkoop W (2009) Fatty acid ratios in freshwaer fish, zooplanknton and zoobenthos. In: Arts MT, Brett MT, Kainz MJ (eds) Lipids Aquat. Ecosyst. Springer, New York, pp 147–178
481 482 483	32.	Hessen DO, Leu E (2006) Trophic transfer and trophic modification of fatty acids in high Arctic lakes. Freshw Biol 51:1987–1998. doi: 10.1111/j.1365-2427.2006.01619.x
484 485	33.	Brett MT, Mu C, Ballantyne AP, et al (2006) Daphnia fatty acid composition reflects that of their diet. Limnol Oceanogr 51:2428–2437.
486 487 488	34.	Galloway AWE, Taipale SJ, Hiltunen M, et al (2014) Diet-specific biomarkers show that high-quality phytoplankton fuels herbivorous zooplankton in large boreal lakes. Freshw Biol 59:1902–1915. doi: 10.1111/fwb.12394
489 490 491	35.	Sarma SSS, Nandini S, Gulati RD (2005) Life history strategies of cladocerans: Comparisons of tropical and temperate taxa. Hydrobiologia 542:315–333. doi: 10.1007/s10750-004-3247-2
492 493 494	36.	Cáceres CE, Tessier AJ (2004) To sink or swim: Variable diapause strategies among <i>Daphnia</i> species. Limnol Oceanogr 49:1333–1340. doi: 10.4319/lo.2004.49.4_part_2.1333
495 496 497 498	37.	de Senerpont Domis LN, Mooij WM, Hülsmann S, et al (2007) Can overwintering versus diapausing strategy in <i>Daphnia</i> determine match-mismatch events in zooplankton-algae interactions? Oecologia 150:682–98. doi: 10.1007/s00442-006-0549-2
499 500 501 502	38.	Demott W, Müller-Navarra D (1997) The importance of highly unsaturated fatty acids in zooplankton nutrition: evidence from experiments with <i>Daphnia</i> , a cyanobacterium and lipid emulsions. Freshw Biol 38:649–664. doi: 10.1046/j.1365-2427.1997.00222.x
503 504 505 506	39.	Elert E Von (2004) Food Quality Constraints in Daphnia: Interspecific Differences in the Response to the Absence of a Long Chain Polyunsaturated Fatty Acid in the Food Source. Hydrobiologia 526:187–196. doi: 10.1023/B:HYDR.0000041604.01529.00
507 508	40.	Patalas K (1990) Diversity of zooplankton communities in Canadian lakes as a function of climate. Verhanlungen des Int verein für Limnol 24:360–368.
509 510 511	41.	Forsström L, Sorvari S, Rautio M, et al (2007) Changes in Physical and Chemical Limnology and Plankton during the Spring Melt Period in a Subarctic Lake. Int Rev Hydrobiol 92:301–325. doi: 10.1002/iroh.200610928
512 513 514	42.	Roiha T, Laurion I, Rautio M (2015) Carbon dynamics in highly heterotrophic subarctic thaw ponds. Biogeosciences 12:7223–7237. doi: 10.5194/bg-12-7223-2015
515 516	43.	Farkas T, Herodek S (1964) The effect of environmental temperature on the fatty acid composition of crustacean plankton. J Lipid Res 5:369–373.
517 518 519	44.	Dufresne F, Hebert PDN (1998) Temperature-related differences in life-history characteristics between diploid and polyploid clones of the <i>Daphnia pulex</i> complex 1. Ecoscience 5:433–437.
520	45.	Gliwicz M, Slusarczyk A, Slusarczyk M (2001) Life history synchronization in a

521 long-lifespan single-cohort Daphnia population in a fishless alpine lake. Oecologia 522 128:368-378. doi: 10.1007/s004420100673 523 46. Tessier AJ, Henry L, Goulden CE, Durand MW (1983) Starvation in Daphnia: 524 Energy Reserves and Reproductive Allocation. Limnol Oceanogr 28:667-676. 525 47. Hartwich M, Martin-Creuzburg D, Wacker A (2013) Seasonal changes in the 526 accumulation of polyunsaturated fatty acids in zooplankton. J Plankton Res 527 35:121-134. doi: 10.1093/plankt/fbs078 528 48. Lampert W, Fleckner W, Rai H, Taylor BE (1986) A study zooplankton : on the 529 Phytoplankton control by grazing spring phase '. Limnol Oceanogr 31:478–490. 530 49. Dupuis AP, Hann BJ (2009) Climate change, diapause termination and 531 zooplankton population dynamics: an experimental and modelling approach. Freshw Biol 54:221-235. doi: 10.1111/j.1365-2427.2008.02103.x 532

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Tables and Figures

Table 1. Lake locations and physical characteristics separated by *Daphnia* winter strategy (active, both strategies, diapause).

	Country	Region	Latitude N	Longitude E	Altitude m	Area m²	Depth m	Ice cover months	Daphnia species
Active									
Malla South	Finland	subarctic	69°03	20°43	599	7850	3.0	9	D. umbra
Saanajärvi	Finland	subarctic	69°05	20°97	679	7.00E+05	24	8	D. umbra
Both strategies	6								
Simoncouche	Canada	boreal	48°25	-71°94	110	8.70E+05	6.0	6	D. galeata, D. dubia,
									D. longiremis
Lunzersee	Austria	alpine	47°51	15°04	608	6.80E+05	34	3	D. longispina
Diapause									
Big Rock	Finland	subarctic	69°10	20°43	527	47	0.5	9	D. longispina
Mekkojärvi	Finland	boreal	61°13	25°08	130	3800	4	6	D. longispina

Table 2. Average (±SD) values of *Daphnia* a) body fat (% of dry weight), b) total FAME concentration (μ gFA mgC⁻¹), c) fatty acid composition (mole % of total fatty lipids) for saturated (Σ SFA), monounsaturated (Σ MUFA), and polyunsaturated (Σ PUFA) fatty acids, d) ratios of Σ SFA:PUFA and Σ n3:n6 per overwintering strategy. n = number of replicates, nd = no data. Bold values are sums of the individual FAME.

	Active	Both Strategies	Diapause
	n=7	n=9	n=7
a) Body fat	58.6 ±11.0	30.9 ±3.3*	32.6 ±4.1
b) ΣFAME	742.6 ±132.0	189.7 ±57.3	214.7 ±10.6
c) ΣSFA	27.0 ±0.1	32.1 ±6.1	26.5 ±5.5
14:0	9.7 ±0.3	11.6 ±7.2	3.7 ±2.03
15:0	0.9 ±0.1	0.7 ±0.9	1.1 ±0.1
16:0	10.7 ±0.3	15.6 ±0.5	14.9 ±5.9
17:0	0.2 ±0	0.8 ±0.2	0.8 ±0.1
18:0	5.6 ±0.4	3.4 ±0.2	6.1 ±1.9
ΣΜUFA	15.4 ±0.6	26.4 ±2.3	43.0 ±16.3
16:1n7c	5.9 ±0.1	8.9 ±0.4	25.3 ±19.8
18:1n9	5.6 ±0.4	8.5 ±0.9	8.2 ±4.8
16:1n9	1.4 ±0.1	4.0 ±3.6	0.9 ±4.7
18:1n7	2.6 ±0.2	5.0 ±2.5	8.7 ±1.1
ΣPUFA	57.6±0.8	41.5 ±8.3	30.5 ±10.8
18:2n6	3.5 ±0.3	4.5 ±0.3	5.0 ±2.3
18:3n3	5.2±0.2	6.9 ±2.6	6.8 ±4.8
18:4n3	25.7 ±0.4	11.6 ±1.6	3.5 ±1.1
18:3n6	1.3 ±0.1	0.8 ±0.2	0.6 ±0
20:4n3	9.0 ±1.8	0.6 ±0.1	0.6 ±0.1
20:4n6	1.6 ±0.2	3.8 ±0.1	4.7 ±1.1
20:5n3	10.6 ±0.5	12.8 ±3.6	9.0 ±4.4
22:6n3	0.8 ±0.2	0.8 ±0.1	0.4 ±0.2
d) ΣSFA:PUFA	0.5 ±0.4	0.8 ±0.3	0.9 ±0.1
Σn3:n6	7.9 ±0.2	3.6 ±0.8	2.0 ±0.1

*this value is only from Lake Simoncouche, Lunzersee did not have a sample for fall body fat.

Table 3. Summary of the permutational ANOVA comparing *Daphnia* a) body fat, b) total FAME, c) FA composition, d) PUFA, e) SFA, f) SDA, g) SFA:PUFA, (h) n3:n6, and i) seston FA composition between the overwintering strategies (active, diapause, both) at the onset of winter. Except for i), all *p*-values were obtained with Monte-Carlo routine (see methods).

Source	df	MS	Pseudo-F	<i>p</i> -value
(a) Body fat				
Strategy	2	1318.9	4.27	0.182
Lake(strategy)	2	256.29	3.17	0.069
Residual	13	80.9		
Total	17			
(b) Total FAME				
Strategy	2	7.08x10⁵	17.4	0.023
Lake(strategy)	3	40852	4.69	0.008
Residual	17	8717		
Total	22			
(c) FA composition				
Strategy	2	82.67	2.67	0.068
Lake(strategy)	3	31.22	5.71	0.001
Residual	17	5.47		
Total	22			
(d) PUFA				
Strategy	2	0.16	7.53	0.061
Lake(strategy)	3	0.02	12.2	0.001
Residual	17	0.002		
Total	22			
(e) SFA				
Śtrategy	2	0.01	1.46	0.344
Lake(strategy)	3	0.009	3.17	0.058
Residual	17	0.003		
Total	22			
(f) SDA				
Strategy	2	0.10	249.3	0.001
Lake(strategy)	3	0.0004	0.283	0.279
Residual	17	0.0003		
Total	22			
(g) SFA:PUFA				
Strategy	2	0.411	2.68	0.217
Lake(strategy)	3	0.154	2.31	0.107
Residual	17	0.067		
Total	22			
(h) n3: n6				
Strategy	2	75.46	52.9	0.004
Lake(strategy)	3	1.43	1.21	0.308
Residual	17	1.18		
Total	22			
(i) FA composition seston				
Strategy	2	29.52	1.01	0.522
Lake(strategy)	3	26.82	5.02	0.001
Residual	9	5.34		
Total	14			

Figure 1. Nonmetric multidimensional scaling (nMDS) plot of fatty acid (FA) methyl ester concentrations from lakes with *Daphnia* showing a) active (black symbols), b) diapause (white symbols) or c) both (grey symbols) overwintering strategies. Data points represent samples collected in fall from the six lakes (MS = Malla South, SA = Saanajärvi, SI = Simoncouche, LU = Lunzersee, BR = Big Rock, ME = Mekkojärvi).

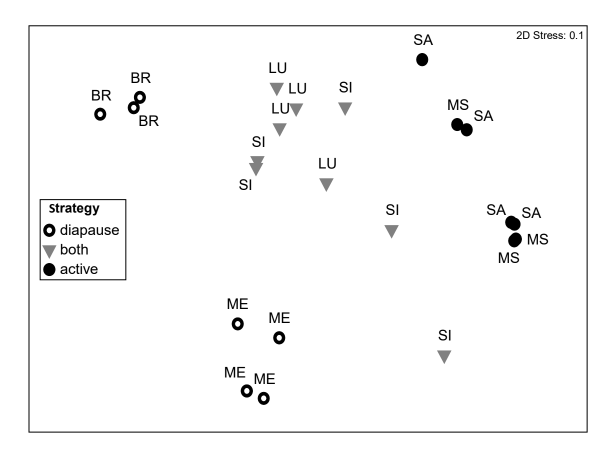
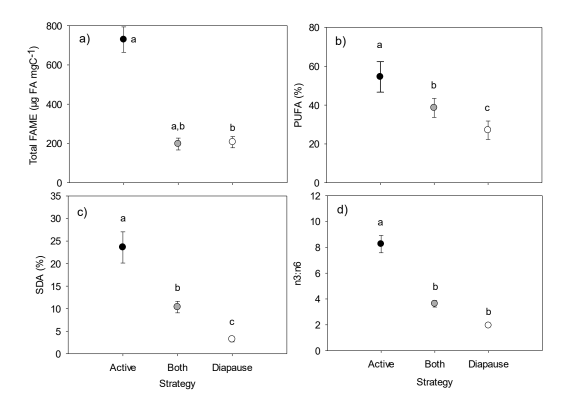


Figure 2. *Daphnia* a) total FAME concentration (µg FA mgC⁻¹), b) percentage of PUFA of total FA, c) percentage of SDA (18:4n3) of total FA and d) n3:n6 ratio in communities showing active, diapause or both overwintering strategies at the onset of winter. Error bars are SE. Different letters indicate statistically different values.



Supplementary Information

SI Table 1. Seston a) total FAME concentration (μ gFA mgC⁻¹), b) fatty acid composition (mole % of total fatty lipids) for saturated (Σ SFA), monounsaturated (Σ MUFA), and polyunsaturated (Σ PUFA) fatty acids, and c) ratios of Σ SFA:PUFA and Σ n3:n6 during the fall. All values are mean ±SD, no value is shown when SD <0.1. n = number of replicates, nd = no data.

	Act	ive	Both st	trategies	Diapause		
	Malla South	Saana-järvi	Simon- couche	Lunzersee	Big Rock	Mekkojärvi	
	n=3	n=3	n=1	n=4	n=1	n=3	
a) ΣFAME	28.9 ±9,5	33.8 ±4,8	19.1	65.3 ±29	80.2	42.2 ±19	
b) ΣSFA	67,2 ±0.3	48.4	43.8	41.3 ±0.05	47.9	88.6 ±0.1	
14:0	41±0.4	6.7	8.2	9.7	3.4	80.0	
15:0	1,6	4.0	2.2	2.1	11.1	0.0	
16:0	21.8 ±0.2	33.9	29.4	23.1	22.5	6.8	
18:0	2.6	2.2	4.0	6.5	10.9	1.8	
ΣMUFA	24.5 ±0.2	44.0	31.9	27.1	39.2	7.5	
16:1n7	13.6	29.4	15.0	19.1	18.5	4.2	
18:1n9	8,3	10.9	11.5	4.1	15.7	3.3	
16:1n9	2.7	3.7	5.1	3.6	5.1	0.0	
18:1n7	0.0	0.0	0.4	0.2	0.0	0.0	
ΣPUFA	8.3	7.6	24.3	31.5	12.9	3.9	
18:2n6	1.0	2.9	4.2	6.9	2.9	0.0	
18:3n3	2.9	0.0	8.2	9.3	1.0	0.0	
18:4n3	0.0	0.0	1.4	0.7	1.0	0.0	
18:3n6	1.6	0.0	0.0	0.2	0.0	0.0	
20:4n3	0.0	2.1	4.9	5.8	1.9	2.3	
20:4n6	0.0	0.4	2.8	4.3	0.0	0.0	
20:5n3	2.9	2.1	2.8	2.3	1.5	1.5	
22:6n3	1.5	nd	nd	1.9	nd	0.0	
c) ΣSFA:PUFA	8.1 ±1.2	6.4 ±0.4	1.8	1.3 ±0.3	3.7	22.8±9.2	
Σn3:n6	7.1 ±0.2	1.3 ±0.1	2.5	1.7	3.3	0.0	

SI Table 2. Site-specific average (±SD) values of *Daphnia* a) body fat (% of dry weight), b) total FAME concentration (μ gFA mgC⁻¹), c) fatty acid composition (mole % of total fatty lipids) for saturated (Σ SFA), monounsaturated (Σ MUFA), and polyunsaturated (Σ PUFA) fatty acids, d) ratios of Σ SFA:PUFA and Σ n3:n6 during the fall when *Daphnia* are either preparing to enter diapause or remain active during winter ice cover. Error values not shown when SD <0.1. n = number of replicates, nd = no data.

	Act	ive	Both strat	tegies	Diapause		
	Malla South	Saanajärvi	Simon- couche <i>D. galeata,</i>	Lunzer- see	Big Rock	Mekko- järvi	
	D. umbra	D. umbra	D. dubia,	D. Iongispina	D. Iongispina	D. Iongispina	
			D. longiremis		longispina	longispina	
	n=3	n=4	n=5	n=4	n=3	n=4	
a) Body fat	59.5 ±9.2	57.7 ±12.8	30.9 ±3.3	nd	23.9 ±5.3	41.2 ±11.1	
b) ΣFAME	838.9±83.7	646.3±181	258.4 ±65.6	121±49	270.2±43	159.2 ±58	
c) ΣSFA	26.9	27.0	36.4 ± 0.1	27.8	30.4	22.6	
14:0	9.9	9.5 ±0.1	16.7 ± 0.1	6.5	2.1	5.3	
15:0	1.0	0.8	0.7	0.9	1.0	1.1	
16:0	10.5	10.9	15.2	15.9	19.1	10.7	
17:0	0.2	0.2	0.6	0.9	0.8	0.7	
18:0	5.3	5.8	3.2	3.5	7.4	4.7	
ΣMUFA	15.0	15.8	28.0	24.8	31.5	54.5	
16:1n7	5.9	5.8	9.2	8.6	11.4	39.3	
18:1n9	5.3	5.9	9.1	8.0	11.5	4.8	
16:1n9	1.3	1.4	6.5 ±0.1	1.4	0.7	1.1	
18:1n7	2.5	2.7	3.2	6.8	7.9	9.4	
ΣPUFA	58.1	57.0	35.6 ±0.1	47.4	38.1	22.9	
18:2n6	3.3	3.7	4.2	4.7	6.6	3.3	
18:3n3	5.0	5.3	5.0	8.7	10.2	3.4	
18:4n3	26.0	25.4	10.4	12.7	2.7	4.3	
18:3n6	1.2	1.3	0.9	0.6	0.6	0.6	
20:4n3	10.2 ±0.1	7.8 ±0.1	0.5	0.7	0.5	0.7	
20:4n6	1.5	1.7	3.7	3.9	5.4	4.0	
20:5n3	10.3	10.9	10.2	15.3	11.9	6.1	
22:6n3	0.6	0.9	0.7	0.9	0.2	0.5	
d) ΣSFA:PUFA	0.5	0.5 ±0.1	1.0 ±0.5	0.6 ±0.1	0.8	1.0 ±0.1	
Σn3:n6	8.7 ±1.5	7.5 ±2.1	3.0 ±0.6	4.2 ±0.4	2.0 ± 0.1	1.9 ±0.2	