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1	Biochemical composition of copepods for evaluation of feed quality in
2	production of juvenile marine fish.
3	
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5	
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11 12 13 14	Abstract
15	To increase current knowledge on the nutritional value of natural prey organisms, the
16	biochemical components of mainly three copepods (Acartia grani, Centropages
17	hamatus, and Eurytemora affinis) from a marine pond system were analysed once a
18	week from spring until late fall, over two years. The analysed components were total
19	lipid, lipid class composition, total lipid fatty acid composition, free amino acids, total
20	protein, protein-bound amino acids, pigment (astaxanthin and ß-carotene), and
21	vitamins (A, thiamine, riboflavin, C, D ₃ , and E). Copepod dry weight (DW), dry
22	matter (% of wet weight), and ash content (% of DW) were also determined. The data
23	are unique due to the homogenous content of copepods in the samples and the long
24	time span of sampling. The copepods were characterised by moderate levels of lipids
25	(6.9-22.5% of DW), with polar lipids accounting for 37.9 to70.2% of the total lipid.
26	The most abundant fatty acids in total lipid (as % of total lipid) were 16:0 (palmitic

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27 acid, 10.8-17.1%), 20:5n-3 (EPA, 8.3-24.6%), and 22:6n-3 (DHA, 13.9-42.3%). The 28 amount of 20:4n-6 (ARA) was generally low (0-2.6%), giving an EPA/ARA range 29 between 7.5and 49.5. The DHA/EPA ratio was between 1.0 and 4.9. Free amino acids 30 (FAA) constituted between 4.3 and 8.9% of copepod DW, and varied with salinity. 31 Glycine, taurine, and arginine dominated FAA, and the fraction of indispensable 32 amino acids varied between 15.5 and 26.8%. Protein, as back-calculated from the 33 protein-bound amino acids (PAA), amounted to 32.7-53.6% of copepod DW, and 34 contained a stable fraction of indispensable amino acids (37.3-43.2% of PAA). 35 Glutamine/glutamic acid, asparagine/aspartic acid, leucine, alanine, and glycine were 36 the most abundant PAA. Astaxanthin was abundant in the copepods (413-1422 μ g/g 37 DW), while B-carotene was not found. High but variable concentrations of vitamin C 38 (38-1232 µg/g DW) and vitamin E (23-209 µg/g DW) were found, while vitamin A 39 and D₃ occurred in trace amounts or were not detected. Detectable levels were found 40 for both thiamine (3.5-46.0 μ g/g DW) and riboflavin (23.2-35.7 μ g/g DW). The data 41 may generate an important base for improvement of live feed enrichment emulsions or formulated feeds used during larval and early juvenile stages in marine fish culture. 42 43

- 44
- 45 Keywords: Lipid class composition, Fatty acids, PUFA, DHA, EPA, TAG,

46 Phospholipid, Protein content, Free amino acids, Pigments, Astaxanthin, Vitamin A,

47 Ascorbic acid, Vitamin D, Vitamin E, Thiamine, Riboflavin, Larval nutrition,

48 Essential nutrients.

49 **1. Introduction**

50

51 High survival and growth, normal pigmentation, and low frequencies of skeletal 52 deformities are characteristics of marine fish reared on natural assemblages of marine 53 zooplankton that mainly consists of copepods (Næss et al., 1995; van der Meeren and Naas, 1997; Støttrup et al., 1998; Shields et al., 1999; Finn et al., 2002; Hamre et al., 54 55 2002). This has been particularly evident for Atlantic halibut (Hippoglossus 56 hippoglossus) and Atlantic cod (Gadus morhua). In the latter case, lagoon or 57 mesocosm rearing is still superior to intensive fry production with rotifers and Artemia 58 as feed. Using copepods as feed compared to intensive rearing of cod larvae on rotifers 59 has indicated a significant nutritional influence on juvenile quality and growth 60 (Imsland et al., 2006). The superiority of copepods for larviculture of marine fish has 61 recently increased the interest for controlled culture of copepods (Støttrup, 2003; Lee 62 et al., 2005).

63

A number of beneficial effects have been linked to copepod nutrient composition in 64 65 relation to early larval nutrition. In particular, emphasis has been put on lipid 66 composition, and the content and ratio of the polyunsaturated fatty acids (PUFA) 67 docosahexaenoic acid (DHA), eicosapentaenoic acid (EPA), and arachidonic acid 68 (ARA) (Scott and Middelton, 1979; Seikai, 1985; Kanazawa, 1993; Reitan et al., 1994; Reitan et al., 1997; Nanton and Castell, 1998; Venizelos and Benetti, 1999; Bell et al., 69 70 2003). The composition of lipid classes and distribution of certain fatty acids between 71 neutral and polar lipids has also gained some attention in lipid nutrition of fish (Olsen 72 et al., 1991; Coutteau et al., 1997; Geurden et al., 1998; McEvoy et al., 1998; Sargent 73 et al., 1999).

74

Further, Nakamura et al. (1986) concluded that insufficient skin pigmentation

76 (melanin) was a result of rhodopsin deficiency, and hence deficiency in the rhodopsin

77 precursors DHA and retinol (vitamin A). In this respect, deficiencies in compounds

78 like carotenoids, thiamine, riboflavin, and cholecalciferol (vitamin D₃) may be

79 considered. Nutrients with antioxidative properties, comprising astaxanthin, ascorbic

80 acid (vitamin C), and tocopherol (vitamin E), may also be of importance. For example,

81 vitamin C appears to enhance the ability of fish larvae to resist stress and infections

82 (Merchie et al., 1997).

83

As marine fish larvae have a high growth potential, they have high dietary 84 85 requirements for protein and essential amino acids. In addition, fish larvae use of 86 amino acids for energy (Rønnestad et al., 1999b; Wright and Fyhn, 2001), which will 87 further increase the demand for dietary amino acids and protein. Consequently, some 88 essential amino acids have been suggested as limiting for larval growth (Conceição et 89 al., 1997; Aragao et al., 2004b). Thus, increased knowledge on the variation in both 90 content and composition of free amino acids and protein in the natural diet will be 91 essential in current understanding on the importance of these factors in larval 92 development and survival.

93

94 Data on biochemical composition of copepods are fragmentary, both with respect to 95 what parameters investigated, and how they vary between copepod species and 96 seasons. Most previous work has concentrated on lipid and fatty acid compositions 97 (Gatten et al., 1983; Watanabe et al., 1983; Witt et al., 1984; Sargent and Henderson, 98 1986; Fraser and Sargent, 1989; Klungsøvr et al., 1989; Olsen et al., 1991; van der 99 Meeren et al., 1993; Norsker and Støttrup, 1994; Evjemo and Olsen, 1997; Evjemo et 100 al., 2003; Morehead et al., 2005). But there are also some data on amino acids and 101 protein (Fyhn et al., 1993; 1995; Helland et al., 2003a,b,c), pigments (Rønnestad et al., 102 1998), and vitamins (Mæland et al., 2000). There are however, to our knowledge, no 103 studies describing the seasonal variation in both macro- and micronutrients in natural 104 prey organisms of fish larvae. The present work includes copepods sampled weekly 105 from a marine pond system over two years from spring to late autumn, and is an 106 attempt to establish more comprehensive database on a number of biochemical 107 components in copepods that are nutritionally important for fish larvae. The work 108 includes analyses of dry matter, ash content, lipids, fatty acids, protein content, 109 protein-bound amino acids, free amino acids, pigments, and vitamins. Such data will 110 be valuable in the on-going research to improve enrichment emulsions and nutritional

- 111 quality of live feed used in marine fish culture, as well as for development of
- 112 formulated starter or early weaning diets for marine fish larvae.
- 113
- 114

115 **2. Materials and methods**

116

117 2.1. Copepod production and collection system

118

119 Copepods were collected during 2000 and 2001 from the marine pond system 120 "Svartatjern" (Naas et al., 1991; van der Meeren, 2003), which is situated near 121 Institute of Marine Research (IMR), Austevoll Research Station at 60°N on the west coast of Norway. Svartatjern is a 20,000 m³ seawater pond, with largest depth of 3.5 122 123 m, and in which all the water can be pumped out and replaced over 3-4 weeks period. 124 A management protocol has been established since the system was started in 1984, 125 which includes draining and refilling the pond twice a year (in early February and 126 early July). Seawater was pumped from 35 m depth in the open fjord outside the pond, 127 and filtered through a UNIK-900 wheel filter (Unik Filtersystem AS, Os, Norway) 128 with 80 µm mesh size (Støttrup, 2005; van der Meeren and Naas, 1997). From March 129 to mid-October, the pond was fertilised weekly or daily depending on weather with 130 agricultural NPK 21-4-10 fertiliser (no trace elements were listed: Yara Norge AS, 131 Oslo, Norway). Fertilisation was always stopped when secci-disk readings became less 132 than 1.5 m. This would ensure a net primary production in the whole water column. 133 The pond was also gently mixed with a propeller placed at 2 m depth. This prevented 134 stratification and formation of oxygen depletion in the bottom layer. This production 135 cycle gives relatively pure populations of mainly calanoid copepods, which are the 136 dominant plankton of Norwegian coastal lagoon systems (Næss, 1996). During winter 137 and pond draining, the copepods survive in the sediments as resting or dormant eggs 138 (Næss, 1991).

139

140 In addition to filtering the incoming water, the UNIK-900 wheel filter was also used

141 for copepod collection from Svartatjern (van der Meeren, 2003). The collection and

142 concentration system was placed inside a small building on a raft in the middle of 143 Svartatjern, and consisted of a slow-impeller-pump (1250 rpm) with up to 1000 l/min 144 capacity, the filter, and six collection and settling tanks. The pump was submerged to 2 145 m depth and lifted pond water into the first compartment of the wheel filter. A rotating 146 fibreglass wheel equipped with 800 µm plankton net sorted out objects too big for 147 being copepods (e.g. hydromedusas), and the water entered the second compartment 148 which was limited by a second wheel with 250 µm plankton net. The copepods were 149 trapped on this latter wheel filter, flushed off into a funnel, and drained down into a set 150 of six 250 l round fibreglass tanks with conical bottoms. When these tanks were filled 151 to the outlet, outputs from the filter bypassed these collection tanks, enabling 152 sedimentation of dead plankton and other organic debris. A timer controlled the wheel 153 filter and pump so collection and sedimentation could take place automatically during 154 night and early morning. In this manner, the remaining live zooplankton could 155 immediately be concentrated in the morning by slowly flushing the tank content 156 through an 80 µm conical plankton net submerged in the pond water. In the tanks, an 157 inner tube with openings 15 cm above the cone prevented settled material from 158 entering the drained water. Further, air and oxygen were supplied at the bottom of the 159 submerged net to prevent the collected copepods from settling in the net cone. From 160 experience, settling would induce heavy mortality among the copepods.

161

162 In addition to collection of copepods, 60 ml water samples were taken at 2 m depth 163 and preserved in 0.6 ml of a glutaraldehyde-Lugol solution (Rousseau et al., 1990) for 164 determination and enumeration of algal species and groups in the pond. 165 Hydrographical data (Table 1) were monitored twice a week with WTW portable 166 meters (WTW LF 330 with Tetra Con 325 probe for salinity and temperature, and 167 WTW Oxi 330 with CellOx 325 electrode for oxygen; WTW GmbH, Weilheim, 168 Germany). Water samples for pH measurements and nutrient analyses were collected 169 once a week and analysed for nitrate (including nitrite), orthophosphate, and silicate, 170 using standard procedures (Koroleff, 1983). A Radiometer PHM 210 (London 171 Scientific Ltd, London Ontario, Canada) was used for pH readings, and nutrients were 172 quantified on a Shimadzu UV-160 UV-visible Recording Spectrophotometer

- 174 hydrography were always collected between at 09:00 and 10:00 h.
- 175

176 In 2001, a single sample of zooplankton was also collected from the Hyltro lagoon in 177 Austevoll, another coastal marine lagoon system previously used for copepod 178 production and juvenile marine fish rearing (Øiestad et al., 1985). However, low 179 copepod biomass prevented further collection from this system. Therefore, no 180 hydrography, nutrients, or phytoplankton samples were collected from the Hyltro 181 lagoon. Moreover, to be able to directly compare the copepod samples with intensive-182 produced live feed for marine fish larvae, one sample of the rotifer Brachionus 183 plicatilis and three samples of Artemia franciscana (Great Salt Lake strain) were 184 included during the 2000 season. The rotifers were reared at IMR with Isochrysis 185 galbana and Rotimac (Bio-Marine Aquafauna Inc., Hawthorne, CA, USA) as feed. 186 Two of the Artemia samples were 1-day old metanauplii obtained from IMR and from 187 the commercial cod and halibut fry producer Austevoll Marin Yngel AS (AMY), 188 respectively. Both these Artemia groups were enriched with DC-DHA Selco (INVE 189 Aquaculture, Dendermonde, Belgium). The third sample was 3-day old Artemia from AMY, which also used Algamac 2000 (Bio-Marine Aquafauna Inc.) as feed in 190 191 addition to the DC-DHA Selco for this on-grown Artemia group. To compare 192 biochemical components of copepod nauplii (sieved through 150 µm and retained on 193 80 µm plankton nets) and the older stages of copepods in the 250-800 µm fraction, 194 three samples of nauplii from Svartatiern were included during the 2000 season. The 195 collected nauplii biomasses were insufficient for other analyses than lipids, dry weight, 196 and content of dry matter and ash. 197 198 In the following, samples from the Svartatjern pond are referred to as copepods and

nauplii, the sample from the Hyltro lagoon as zooplankton, and the samples of the
intensive produced live feed as rotifers and <u>Artemia</u>.

201

202 2.2. Sample preparation

204 The collected copepods were transported live for 10 min in a black 12-l-bucket to the 205 sample preparation laboratory. Here, the copepods were placed in a mixing column of 206 6 l volume and 9.5 cm diameter (van der Meeren, 2003), with densities between 400 207 and 900 copepods/ml. To ensure proper mixing and sufficient oxygen supply, air and 208 oxygen were mixed and bubbled gently from the tip of the cone at the bottom of the 209 column. With this arrangement, copepods could easily be kept alive for more than 4 h, 210 which was sufficient to prepare the samples for biochemical analyses. The bubbling 211 also led to a homogenous distribution of copepods in the column, as shown from a 212 biomass of 2.6 g \pm 0.12 (mean wet weight \pm SD) among 10 subsequent samples of 213 equal volume collected through a silicon tube placed 15 cm above the cone bottom. 214 Further, the relationship between sample size in ml (V) and sample wet weight in 215 grams (WW) showed high correlation among 5 replicate samples of unequal volume in the range of 50 to 500 ml (V = 258.98 WW - 26.379, $R^2 = 0.9989$). Similarly, the 216 217 relationship between actual counts of copepods from these samples (N) and V also showed high correlation (N = 138.46 V + 753.26, $R^2 = 0.9942$). In this way, 218 219 uniformity of collected biomass among repeated samples from the column was 220 demonstrated.

221

Aliquots of copepods were sampled from the column for the following biochemical 222 223 analyses: lipid classes and total lipid fatty acids, pigments, protein and free amino 224 acids, lipid-soluble vitamins, and water-soluble vitamins. In addition, one aliquot was 225 collected to determine individual copepod wet weight, followed by another aliquot for 226 determination of dry matter and ash content. Between 0.5 and 2.7 g copepod wet 227 weight were sampled for each analysis. Finally, an aliquot of 50 ml was preserved with 228 0.9 ml Lugol solution for identification of copepod species and stages, as well as other 229 zooplankton species. Copepod samples were also made available for iodine analyses 230 (published in Moren et al., 2006).

231

Wet weight was determined in all unpreserved samples by weak vacuum filtration at
680 mm Hg (van der Meeren, 2003). The unit was equipped with 52 mm diameter
filter disks of 60 µm mesh size plankton net (Sefar Nitex 03-60/35, Sefar Holding Inc.,

235 Freibach, Switzerland). To remove salt, the samples were flushed 2-3 times with 10% 236 salt water made from distilled water and 0.2 µm filtered 35‰ seawater. Salinity lower 237 than 10‰ was observed to burst the copepod exoskeleton, with subsequent loss of 238 biomass. The resulting semi-dry "cake" of copepods was further divided into subsamples by a spatula and transferred to pre-weighed Nunc cryotubes with an externally 239 240 treaded lid. The cryotubes were then quickly weighed to nearest 0.1 mg on a Mettler 241 AE200 (Mettler-Toledo Inc., Columbus, OH, USA). Lipid samples were then 242 immediately frozen in liquid nitrogen, while the samples for the other biochemical 243 components were quickly placed in an -80°C freezer. By this procedure, a short time 244 (3-5 min) was ensured from sample collection to placement in freezer. 245

246 The sample for determination of individual copepod WW was first filtered and 247 weighed as described above, then 75 to 100 ml of 10% salt water was added along 248 with a few drops of Lugol solution to improve contrast, and finally ten well-mixed 249 aliquots of 0.2-0.5 ml were collected from the sample and counted to determine the 250 total number of copepods. A Leica MS5 stereo Microscope with options for both light 251 and dark field (Leica Microsystems GmbH, Wetzlar, Germany) was used for counting. 252 Variation among the 10 counts was low, with an average coefficient of variation of 253 11%.

254

255 After freezing, the sample for determination of dry matter content was dried in a Heto 256 FD8 freeze-drier (Heto-Holten AS, Allerød, Denmark). A freeze-drying period of 72 h 257 was necessary to reach stable weight. To ensure reliable dry weight (DW) 258 measurements over a range of different sample sizes, the sample DW in g was 259 regressed on the corresponding WW for 9 replicate samples between 0.5 and 5.0 g wet 260 weight. This sample series showed high linear correlation (DW = 0.140 WW + 0.004, $R^2 = 0.999$). Amount of dry matter (% of WW) was calculated, and ash content (% of 261 262 DW) was determined by combusting at 550°C for 24 h in pre-weighed porcelain 263 crucibles.

The frozen samples for analysis of total protein, protein-bound amino acids, and free amino acids were also freeze-dried and weighed for determination of DW before being shipped in dry condition to the laboratory for analysis. All other samples were packed on dry ice and kept frozen when shipped to the analytical laboratories within 3 h. Preparation of the zooplankton, rotifer, and <u>Artemia</u> samples was in all respects similar to the copepod samples.

271

After the sample preparations were completed, copepod viability of the remaining 272 273 biomass was checked by a light-dark test. A sample of copepods was placed on a Petri 274 dish with seawater, and partly covered by aluminium foil. The cover was then moved 275 to the other half of the disk. In both cases, almost 100% of the copepods gathered 276 under the shadowed area within a short time. This was consistent throughout the 277 sampling seasons, showing no mortality during sample collection. In addition, the 278 samples were inspected under the Leica stereo microscope for damages on the 279 copepod antennae and tail, and for content of organic debris (van der Meeren, 2003).

280

281 2.3. Analytical methods

282

283 2.3.1. Lipids and fatty acids

284

285 Frozen samples were homogenized in solvent using an Ultra Turrax (IKA Werke 286 GmbH, Staufen, Germany) and total lipid extracted according to the method of Folch 287 et al. (1957). After evacuation of the solvent under nitrogen, water was evacuated 288 under vacuum over dry sodium hydroxide, and total lipid quantified gravimetrically. 289 The lipid was then stored in chloroform:methanol (2:1) under nitrogen at -80°C until 290 used for further analysis. Lipid class composition was assessed using the HPTLC 291 double development method of Olsen and Henderson (1989). For fatty acid analysis of 292 total lipid, portions of the samples were subjected to the sulphuric acid catalysed 293 transesterification method of Christie (1982), extracted into hexane, and stored at – 294 80°C until analysed. Quantitative analysis of fatty acid methyl esters were carried out 295 by gas liquid chromatography using a HP 5890 gas chromatograph (Hewlett Packard

Labs Inc., Palo Alto, CA, USA) equipped with a J&N Scientific Inc DB-23 fused silica
column (30 m x 0.25 mm i.d.) as described by Olsen et al. (2004). Abbreviations for
lipid classes and fatty acids used in the text are given in Table 2.

299

300 2.3.2. Protein and amino acids

301

Sub-samples (15-25 mg) of the freeze-dried samples were extracted in Eppendorf 302 303 tubes in 1 ml 6% tri-chloro-acetic acid (TCA) under rotation (Heto Rota-Mix) for 24 h at 4°C. After centrifugation (15000 x g, 10 min, 4°C), the supernatant was used for 304 305 free amino acid (FAA) analysis after appropriate dilution in borate buffer (100 mM, 306 pH 10.4). The precipitate was washed once in 6% TCA, re-centrifuged, and transferred 307 to a 10 ml tube and dissolved in 4 ml of 1 M NaOH by rotation for 48 h at room 308 temperature for analysis of total protein and protein-bound amino acids (PAA). After 309 centrifugation (15000 x g, 10 min, 20°C), the supernatant was collected and 310 appropriately diluted to 0.5 M NaOH with distilled water, and used for determination 311 of total protein by the method of Lowry et al. (1951), using the micro-modification of 312 Rutter (1967) with bovine serum albumin (BSA, Sigma A-7638) in 0.5 M NaOH as 313 standard and 0.5M NaOH as blank. The colour was allowed to develop in darkness for 314 30 min and, after an additional mixing, the sample absorbance was read on a Perkin 315 Elmer Biolambda spectrophotometer (PerkinElmer Inc., Waltham, MA, USA) at 750 316 nm. Preliminary tests showed no increase in the protein or FAA contents of the freeze-317 dried copepod, Artemia, or rotifer material by Potter-Elvehjem glass-glass 318 homogenisation, so direct extraction of the freeze-dried material in TCA or NaOH was 319 routinely used in this study. 320 321 An aliquot (200 µl) of the NaOH supernatant was added concentrated HCl to reach 322 final concentration of 6 M HCl to allow acid protein hydrolysis (106°C, 24 h) in N₂-323 flushed stoppered glass vials. Samples of 6 M HCl were included in the hydrolysis as

324 blank controls. The hydrolysed samples were neutralised by addition of equal volume

325 of 6 M NaOH and appropriately diluted in the borate buffer before analysis. All

326 reagents used in the analyses were prepared from glass-distilled, ion-exchanged

327 (Millipore Milli-Q) water with a resistance of $18 \text{ M}\Omega$.

328

329 Amino acid analysis was performed by reversed-phase chromatography using a Gilson 330 HPLC (Gilson Medical Electronics Inc., Middleton, WI, USA) with fluorometric 331 detection (OPA and FMOC reagents) and connected to an ASTED (Automated 332 Sequential Trace Enrichment of Dialysates) sample robot and a 3 x 150 mm, 3 µm 333 particle size Inertsil ODS-3 column from Varian (Varian Inc., Palo Alto, CA, USA). 334 The analytical reproducibility based on repetitive analyses of standards was <1% for 335 all amino acids except proline (4%). The applied HPLC procedure did not separate 336 phosphoserine and aspartic acid. In the analysis of FAA of the 2001 samples, the 337 glycine peak dominated the following threonine peak so it could not be resolved or 338 quantified. Protein-bound tryptophan is difficult to quantify after acid hydrolysis since 339 it is partly destroyed by the treatment. Gilson Unipoint 715 Software, version 2.10 was 340 used for peak analysis and sample integration.

341

342 The PAA values (umoles/mg DW of analysed material) were converted to the 343 equivalent protein content and expressed both in molar terms of the various amino 344 acids (µmoles/mg DW), and in weight-specific terms as an equivalent to protein 345 content (μ g/mg DW). Abbreviations for the amino acids used in the text are the lower 346 case equivalents to abbreviations used in Tables 3 and 4. The terminology of 347 dispensable (DAA) and indispensable (IAA) amino acids are used according to Harper 348 (1983) and the following 10 amino acids are termed IAA for fishes according to 349 Wilson (1985): arg, his, ile, leu, lys, met, phe, thr, trp, and val. The inclusion of arg 350 and tyr among the IAA in this study of the natural feed organisms of fish larvae is in 351 agreement with results on embryonic and neonatal vertebrate nutrition which 352 document their strong dependency on amino acids (e.g. Rønnestad et al., 2003; Wu et 353 al., 2004; Dabrowski et al., 2005; Urschel et al., 2006, 2007). 354

355 2.3.3. Pigments

357 The frozen samples were added acetone and homogenized on ice using an Ultra Turrax 358 homogenizer. Moisture was removed by means of Na₂SO₄ and samples stored at -80°C 359 until analysed. Astaxanthin and β -carotene were quantified using a HP automated 360 sample injector (G1329A ALS), a G1315A DAD diode array detector and G1316A 361 ColComp column temperature controller, maintained at a constant temperature of 4°C. 362 Separation was performed using tandem installed Chromspher 5 mm C18 columns 363 (100 mm x 3 mm i.d.) with a guard column of C18 material (Chromsep guard column 364 SS) preceding the main column. The mobile phase was acetonitrile:dichlormethane:methanol:propionic acid:water (61:20:7.6:5.7:5.7), which 365 366 was filtered before use. Vitamin C (263 mg/l) was added to the mobile phase as an 367 antioxidant. The flow rate was isocratic at 1 ml/min. Both column and auto injector 368 temperatures were maintained at 1°C. Peaks were detected at 476 nm for astaxanthin 369 and β -carotene, and subsequently quantified with reference to authentic standards.

Each sample was analysed in triplicates. Data were stored and processed using HPChemstation software.

372

373 *2.3.4. Vitamins*

374

All analyses of vitamins were performed on thawed samples and related to wet sample
weight. After analysis, data were converted relative to DW by dividing with the dry
matter fraction obtained from separate samples as described above in section 2.2.
Whenever vitamin concentration was between the detection and quantification limits,
it was denoted as trace amounts. However, to reduce error and variation, particularly at
low vitamin concentrations, the trace values were included in the calculations of
average vitamin levels.

382

Samples for analyses of the lipid soluble vitamins were homogenised and weighed into screw-capped glass tubes, saponified, and extracted with hexane. Vitamin D was upconcentrated by passage over a preparative normal phase HPLC column, where the isomeres D_2 and D_3 eluted as one peak, which was collected. The collected fraction was then subjected to analytical reverse phase HPLC with UV detection at 275 nm,

388 which separates the vitamin D isomers. Vitamin D_3 was quantified by using vitamin 389 D_2 as internal standard and vice versa. Vitamin D_2 was not detected at all in the 390 samples. The method and instrumentation are described in detail in Horvli and Lie 391 (1994) and CEN (1999a).

392

393 Vitamin A was subjected to normal phase HPLC with UV detection at 325 nm and 394 quantified by external standards according to method and instrumentation described in 395 Moren et al. (2004a). This method gives a large peak with similar retention time as all 396 trans retinol in samples from Artemia. However, later work has shown, by the use of a 397 diode array detector, which produces UV spectra of the peaks, that this compound is 398 not vitamin A (Moren et al., 2005). The tocopherols (vitamin E isomers) were also 399 analysed by normal phase HPLC, detected by fluorescence at 295 nm excitation and 400 330 nm emission and quantified using external standards (CEN 1999b). Given relative 401 to wet weight of the sample, the detection and quantification limits of the analytical 402 methods are 6 and 20 ng/g for vitamin D, 8 and 28 ng/g for vitamin A, 11 and 38 ng/g for α -tocopherol, and 8 and 28 ng/g for the other tocopherols, respectively. 403

404

405 The samples for ascorbic acid (vitamin C) were homogenised and extracted in meta-406 phosphoric acid with dithiothreitol, which reduces de-hydro ascorbic acid to ascorbic 407 acid. Compounds in the extract were separated by reverse phase HPLC, and ascorbic 408 acid was detected by amperiometrically at 0.6 V and quantified using external 409 standards (Mæland and Waagbø, 1998). The B vitamins, thiamine and riboflavin, were 410 analysed by semi-automated microbiological methods which are detailed in Mæland et 411 al. (2000). Detection and quantification limits of the methods relative to wet weight of 412 the sample are 0.35 and 1.1 μ g/g for vitamin C, 1.3 and 4.3 μ g/g for riboflavin, and 413 0.02 and 0.2 μ g/g for thiamine, respectively.

414

415 2.4. Statistical analysis

416

417 Differences in biochemical indices were tested by Students t-test after checking for

418 normal distribution by Kolmogorov-Smirnov tests for normality (goodness of fit,

419 Lilliefors P-values). Student t-tests were carried out for copepods between the two 420 years, and between copepods and copepod nauplii in 2001. Whenever the biochemical 421 indices were percentages, arcsine transformation was carried out before statistical 422 testing as suggested by Sokal and Rohlf (1995). Differences among means were 423 considered statistically significant at P < 0.05. 500 424 425 426 **3. Results** 427 428 3.1. Hydrography and phytoplankton 429 430 Temperature in Svartatjern during sample collection (Table 1) typically started 431 between 7-9°C in the spring, rising in May to around 15-16°C with a peak of 18-19°C 432 before emptying the pond in mid-summer. After refilling in late July, temperature was 433 in the range of 17-18°C until early September, and dropped gradually to 7-6°C at early 434 December. Salinity started in the range of 30-31% every time the pond was, but

435 dropped slowly over time due to precipitation run-off. At salinities below 24‰, new 436 salt water was pumped into the system. Average salinity was 25.2 and 26.3% for 2000 437 and 2001, respectively (Table 1). Oxygen saturation fluctuated with algal production, 438 being highest during periods of net primary production at good light conditions (March 439 to October). During intensive primary production in May and June, water became 440 supersaturated with oxygen (up to 160% saturation) and with corresponding high pH 441 level up to 9.1 (Table 1). Average Secci disc readings were 1.4 and 1.7 m in 2000 and 442 2001, respectively. Algal nutrients (Table 1) were low during the seasons of net 443 primary production, but increased quickly from mid-October when light intensity and 444 photoperiod declined.

445

446 Many of the phytoplankton species present in the pond were small (3-5 μ m) single-447 celled specimens that were not possible to identify. This confined between 81.9 and 448 99.9% of monads and flagellates, which overall was the most abundant phytoplankton 449 group (Fig.1), with densities in the range of 21 to 378 cells/ μ l (2000), and 1 to 269

450 cells/µl (2001). Both years, cell densities of monads and flagellates fell below 30 451 cells/µl at end of October. Similarly, all other phytoplankton groups also quickly 452 declined in late autumn (Fig. 1). Considering abundances above 5 cells/µl, 453 Rhizosolenia fragilissima was initially the most abundant diatom (Bacillariophyceae) 454 with 19 cells/µl during late May of the 2000 season. This was followed by the green 455 algae (Chlorophyceae) Gloeocystis sp (5 cells/ul) and Oocystis sp (11 cells/ul) in last 456 half of June, with late September appearance of the diatoms Skeletonema costatum (10 457 cells/ μ l) and a small Chaetoceros sp (93 cells/ μ l) in October. 458 In 2001, the green alga Nephrocytium sp (11 cells/ul) was abundant in April and first 459 half of May, followed by Gloeocystis sp (22 cells/µl) and Oocystis sp (16 cells/µl) that 460 461 lasted until end of August. R. fragilissima peaked at 9 cells/µl in late May, but was 462 abundant until late July. Among the diatoms, a small Thalassiosira sp bloomed to 12

463 cells/µl in late July and lasted to mid-October, while N<u>itzschia closterium</u> went up to a
 464 maximum of 43 cells/µl during it's blooming period in September and October. Other

465 algae just exceeding 5 cells/µl in 2001 were Katodinium sp (Dinophyceae) in mid-

466 June and <u>Emiliania huxleyi</u> (Haptophyceae) in late July. Ciliates were often dominated

467 by <u>Strombidium</u> sp, and reached high levels of more than 100 cells/ml several times

468 during late spring and autumn both years (Fig.1).

469

470 *3.2. Copepod species and stages*

471

472 Three species of copepods dominated the samples from Svartatjern: Eurytemora 473 affinis, Centropages hamatus, and Acartia grani (Fig. 2). These copepods typically 474 occurred in single or paired dominance, and a substantial fraction of all three species 475 together was therefore rarely observed and only during short transitions. In 2001, the 476 common succession pattern previously observed in Svartatiern from spring to autumn 477 (Eurytemora-Centropages-Acartia-Centropages-Eurytemora) was shifted, as A. grani 478 had its main season before the pond was emptied at mid-summer, and therefore 479 overlapped with E. affinis in May. In this sense, the seasonal succession pattern 480 diverged the two years of copepod collection. Other copepod species constituted

maxima of 2.4% (2000) and 3.4% (2001) of the total zooplankton items in the samples
(Fig. 2). Of non-copepod zooplankton species in Svartatjern, the cladoceran Podon sp
occurred only during short periods and contributed up to 13.1% (2000) and 20.5%
(2001) of single samples (Fig. 2). Podon sp was most abundant during September both
years. The other brief contributor to the non-copepod zooplankton was young medusa
stages of Sarsia sp, with 11.3% of the plankton numbers and only found in the 18May-sample of 2001.

488

489 The nauplii sample from April 2001 contained both copepod nauplii (55%) and first

490 copepodid stages (45%). In this sample, 32% was <u>A. grani</u>, while <u>C. hamatus</u> and <u>E.</u>

491 <u>affinis</u> constituted the rest. Copepodids were not found in the other two nauplii

492 samples from late July and mid-September 2001, in which <u>A. grani</u> comprised 65 and

493 39%, respectively. In the Hyltro lagoon sample, <u>E. affinis</u> constituted 43.3% of

494 enumerated zooplankton, while other observed zooplankton species or groups were the

495 copepod <u>Paracalanus parvus</u> (2.7%), copepod nauplii (16.7%), decapod zoeae (32.4%),

496 and <u>Sarsia</u> sp medusae (4.8%).

497

498 3.3. Zooplankton size, dry matter and ash content

499

500 Individual copepod DW (Fig. 3, Table 2) was in the ranges of 5.3-13.7 µg (2000) and 501 4.2-13.9 µg (2001). In 2000, DW increased with time and reached maximum values in 502 late June, and another maximum in October. In contrast, the 2001 copepods were 503 biggest in late May, and smallest in November. The DW of individual zooplankton 504 from the Hyltro lagoon was 9.9 µg. Copepod nauplii (Table 2) had low DW in two of 505 the samples (0.18 and 0.25 μ g per nauplius at end of July and mid-September, 506 respectively), while DW was 1.46 µg per nauplius in the late-April sample, reflecting a 507 higher content of young copepodid stages observed in this latter sample. The rotifers 508 weighed 0.61 µg per individual (Table 2), while 1-day-old Artemia was 2.12 and 2.14 509 μg and 3-day-old Artemia was 2.48 μg.

511	Dry matter content in the copepods was quite stable and averaged 14.9 and 15.3% of
512	WW for 2000 and 2001, respectively (Fig. 3, Table 2). Dry matter content of the
513	nauplii was quite similar to the copepods (Table 2). In contrast, the zooplankton
514	contained more dry matter (17.7%). Rotifers contained less dry matter (13.2%) than
515	copepods, and <u>Artemia</u> even less (8.9-10.8%).
516	\mathcal{Q}
517	Average ash content was quite constant both years, and within 9.5 and 10.5% of DW
518	for the copepods, nauplii, rotifers, and Artemia (Table 2, Fig.3). This contrasted the
519	zooplankton sample, which contained 14.3% ash.
520	
521	3.4. Lipids and fatty acids
522	
523	The total lipid content (TL) in the copepods was relatively low and stable, with the
524	exception of one sample that was 220 μ g/mg DW, corresponding to 22% of DW
525	(Table 2, Fig. 4). Average copepod TL in 2000 and 2001 was close (108 and 111
526	μ g/mg DW), while TL of the copepod nauplii (86 μ g/mg DW) was significantly lower
527	than in the copepods. The zooplankton had higher TL than that found in copepods and
528	was more similar to the rotifers (Table 2), while Artemia was loaded up with a lipid
529	content of approximately 25% of DW.

530

531 Regarding lipid class composition, the main components of the copepod neutral lipids 532 were TAG and cholesterol. TAG averaged 2.6 and 2.2% of copepod DW in 2000 and 533 2001, respectively, which corresponded to 21.9 and 20.2% of TL for the two years 534 (Table 2, Fig. 4). Similarly, mean cholesterol levels were 1.5 and 1.3% of copepod 535 DW, equivalent to 13.2 and 12.4 % of TL in 2000 and 2001, respectively. Copepod 536 nauplii had lower fractions of TAG and cholesterol than the average values of the 537 copepods samples, but only statistically significant for cholesterol. TAG showed a 538 large variation among both copepod and nauplii samples. The zooplankton displayed 539 almost twice the amount of TAG (4.2% of DW and 29.4% of TL) compared to copepods, and rotifers had even more TAG (6.1% of DW and 39.4% of TL). In 540 541 Artemia, TAG constituted as much as 16.8-19.6% of DW (69.0-77.1% of TL). It

should also be noted that the algae-derived neutral glycoglycerolipids (galactocides) in
combination with neutral glycosphingolipids (cerebrocides) or sulfoglycoglycerolipids
(sulfolipids) were more or less absent in rotifers and <u>Artemia</u> (Table 2: MGDG+CB
and DGDG+SL). Significant differences in copepod neutral lipids between the two
years were only detected for MGDG+CB.

547

548 Amounts of polar lipids in the copepods averaged 6.2 and 6.3% of copepod DW for

549 2000 and 2001, respectively, with a relatively stable fraction averaging 57.1% (2000)

and 58.2% (2001) of TL (Table 2, Fig. 4). Polar lipid content in rotifers was more

similar to copepods, constituting 6.1% of DW but corresponding only to 39.8% of TL.

552 Copepod nauplii and the zooplankton sample had somewhat lower content of polar

553 lipids (5.4 and 5.1% of DW, equivalent to 63.1 and 36.0% of TL, respectively). In

554 contrast, polar lipids in <u>Artemia</u> were lower and between 3.9 and 5.0% of DW (15.4-

555 20.5% of TL). The major phospholipids in copepods and copepod nauplii were PC and

556 PE, each having average levels between 1.5 and 2.0% of DW and 17.6-20.5% of TL

557 (Table 2, Fig. 4). Significant difference in copepod phospholipid class composition

558 between the two years was only found for PS. PC and PE also dominated

phospholipids in the zooplankton and the rotifer samples (1.5-2.1% of DW and 10.6-

560 13.7% of TL), as well as in the <u>Artemia</u> samples (1.2-1.9% of DW and 4.8-7.6% of

561 TL).

562

563 In the copepods, PUFA dominated the TL fatty acid composition, accounting for 63.3 564 and 64.2% of TL in 2000 and 2001, respectively (Table 2). Variation in PUFA was 565 low between the samples within each year. Although not significantly different from 566 the copepods, PUFA fraction in copepod nauplii was even higher (69.4% of TL), on 567 the expense of MUFA. Zooplankton was more similar to rotifers and Artemia, with 568 PUFA levels ranging between 43.6 and 48.5% of TL. Compared to copepods, these 569 reduced levels of PUFA were balanced by increased fractions of MUFA (20.1-34.8% 570 of TL).

572 Among the single fatty acids, DHA was abundant in the copepod samples, averaging 573 34.4 and 32.9% of TL for 2000 and 2001, respectively (Table 2, Fig. 5). The copepod 574 nauplii averaged 40.5% DHA, which was significantly higher than for the copepod 575 samples in 2001. These high levels contrasted the DHA fraction of 17.3% found in the zooplankton sample. In the intensively produced live feed, DHA was between 10.6 576 577 and 20.0%, with highest level in the 3-day on-grown Artemia. In the copepods, 578 averages of EPA were between 16.2 and 17.4% of TL, including copepod nauplii and 579 zooplankton. However, in rotifers and Artemia EPA was lower, ranging between 7.1 580 and 9.2%, respectively. Another abundant fatty acid was palmitic acid (16:0), which 581 was between 13.7 to 19.7% of TL in all groups (Table 2). Among other important fatty 582 acids, ARA was very low in the copepod and copepod nauplii samples and even below 583 detection limit in many samples. This contrasted that of zooplankton, rotifers, and 584 Artemia where ARA was more abundant, ranging between 1.6 and 3.2% of TL. 585 Significant differences in fatty acids composition between the copepod samples from 586 2000 and 2001 were mainly found among the fatty acids with 18 carbon atoms (C18), along with myristic acid (14:0). Similarly, significant lower fractions among C18 fatty 587 acids were also found for copepod nauplii when compared with the copepod samples 588 589 from the same year (Table 2).

590

591 The average DHA/EPA ratio was 2.1 and 2.2 for copepods in 2000 and 2001, 592 respectively (Table 2, Fig. 5). Copepod nauplii had somewhat higher DHA/EPA ratio, 593 but not significantly different from the 2001 copepods. The zooplankton had the 594 lowest DHA/EPA ratio (1.1), while intensively reared live feed varied between 1.4 and 595 2.2, the latter belonging to 3-day on-grown Artemia. The EPA/ARA ratio was in 596 general very high in copepods and copepod nauplii (on average between 23.2 and 597 27.7), and also relatively high in the zooplankton sample (10.3). This contrasted the 598 EPA/ARA ratios in rotifers (3.7) and Artemia (2.9-4.0). A similar pattern was seen for 599 the (n-3)/(n-6) ratio, which was highest in copepods and lowest in the rotifers (Table 2, 600 Fig. 5).

603

604 The protein content determined by the Lowry method using BSA as reference standard 605 averaged 38.3 and 56.5% of copepod DW for 2000 and 2001, respectively (given as 606 µg/mg DW in Table 3). This difference was significant, but did not correspond to a 607 similar magnitude in the protein calculated from weight-specific protein-bound amino 608 acids (PAA_w). Although still significantly different, the average PAA_w values in 609 copepods from the two years were more similar, and corresponded to 44.4 and 41.3% 610 of copepod DW in 2000 and 2001, respectively. Variation in PAA_w over time was low 611 (Table 3, Fig. 6) as indicated by a coefficient of variation close to 10%. No significant 612 correlations were observed between protein determined by the Lowry method and 613 protein calculated as PAA_w for any of the two years with copepod samples. Some 614 discrepancy also occurred between the two methods of protein content determination 615 in the zooplankton sample (36.6 vs. 30.3% for the Lowry vs. PAA_w method), while 616 protein contents determined by the two methods were more in agreement for rotifers 617 and Artemia samples (Table 3). Rotifers were lowest in PAA_w-calculated protein content (24.8% of DW), followed by 1-day-old and 3-day-old Artemia (27.8 to 36.8% 618 619 of DW). The reasons for the discrepancies in protein determination between the Lowry 620 and the PAA_w methods for zooplankton and copepods were not clarified. 621

The concentration of protein-bound amino acids (PAA_c) was lowest in rotifers (2.3

 μ moles/mg DW), being almost half of that in copepods in 2000 (4.1 μ moles/mg DW)

624 (Table 3). All concentration-specific PAA and IAA indices applied on the copepod

samples were significantly different between 2000 and 2001, but with low variation

626 within each of the years (Table 3, Fig. 6). Considering all prey types sampled, the

627 concentration-specific IAA fraction of PAA (IAA_c/PAA_c) was between 40.4 and

43.7%. Similarly, the IAA_c/DAA_c ratio of the hydrolysed protein averaged 0.68 and

629 0.70 in the copepod samples from 2000 and 2001, respectively (Table 3), while for the

630 rotifers and Artemia it was higher (between 0.75 and 0.78). In contrast, the

631 IAA_c/DAA_c ratio in the zooplankton sample was 0.71, and more in accordance with

632 the copepods.

In the PAA_c, leu, val, lys, and ile were the most dominant IAA in all samples, followed 634 635 by arg, phe, and thr (Table 3). Among DAA, glu+gln, asp+asn, ala, and gly were the 636 most abundant amino acids. Concentrations of all amino acids, except lys and asp+asn, 637 were significantly different between the copepod samples of the two years (Table 3). 638 In absolute values, amino acid concentrations were generally lower in the zooplankton, 639 rotifers, and Artemia, compared to the copepods (Table 3). However, regarding the 640 amino acid profiles expressed as percentage of the hydrolysed copepod protein, they 641 were similar the two years of sampling (Fig. 6), with no significant differences found 642 for major IAA as thr, leu, lys, and ile. Also the zooplankton, rotifers, and Artemia 643 PAA profiles showed similarities with the copepods. The observed differences can be 644 attributed to very low variation in fractions of single amino acids in the hydrolysed 645 protein (Fig. 6), typically displaying coefficients of variation between 3 and 15%.

646

647 *3.6. Free amino acids*

648

The weight-specific content of free amino acids (FAA_w) in the copepod samples from 649 650 Svartatjern varied between 4.3 and 8.9% of copepod DW, averaging 5.6 and 6.5% for 2000 and 2001, respectively (given as $\mu g/mg$ DW in Table 4). The average FAA_w 651 652 content of the copepods was significantly different between the two years. In the 653 zooplankton sample, FAA_w was in the upper range of the levels observed in the 654 copepods and composed 8.6% of the zooplankton DW, while in the intensive reared 655 live feed FAA_w was considerably lower than in copepods and corresponded to 1.7% in 656 rotifers and 2.6 to 3.4% in Artemia.

657

Concentration of free amino acids (FAA_c) was lowest in rotifers and <u>Artemia</u>, higher
in copepods, and highest in the zooplankton (Table 4). The absolute levels of
indispensable free amino acid concentration (IAA_c) in copepods were not significantly
different between 2000 and 2001. However, significant differences among copepods
occurred between the two years when other concentration-specific IAA indices like
IAA_c/FAA_c and IAA_c/DAA_c ratios were considered, and among concentrations of
most individual FAA (Table 4). Only the rotifers had a higher IAA_c/FAA_c fraction

665 (30.6%) than the copepods (19.1-24.3%), with <u>Artemia</u> and zooplankton displaying the 666 lowest IAA_c/FAA_c fractions (10.0-15.6%). A similar pattern was demonstrated for the 667 IAA_c/DAA_c ratio. Variation in all IAA_c indices was low among the copepod samples 668 each year (Table 4, Fig. 7).

669

670 Assuming similar levels of thr in 2001 as in 2000, the averaged copepod FAA_c profiles

- 671 expressed as percentage (relative abundance) were dominated in decreasing order by 672 gly, tau, arg, and ala (26.9-9.0%, totalling 70.6% of FAA_c in 2000, and 39.0-6.1%,
- totalling 76.9% of FAA_c in 2001). In the zooplankton sample, the four most abundant

amino acids were in decreasing order gly, tau, pro, and arg (30.3-8.8%, totalling 72.8%)

675 of FAA_c), with also ala being abundant (8.8%). In rotifers, the FAA_c profile was more

676 diverse, and the four most abundant amino acids included ser, glu, arg, and tyr (13.2-

8.5%, adding up to 44.3% of FAA_c). The four most abundant FAA_c in the <u>Artemia</u>

samples were all DAA and comprised tau, ala, pro, and glu (averaged to 24.3-12.6%

679 which summed up to 68.0% of total FAA_c). Relative abundance of single amino acids

680 in the FAA_c profiles throughout the sampling season was more variable compared to

681 the PAA_c profiles (Fig. 6, 7).

682

683 Considering all copepod samples of both years, total FAA concentration correlated 684 significantly with salinity ($R^2 = 0.379$, P < 0.0001), where increased salinity elevated 685 the total FAA_c level. Among individual amino acids of the FAA_c pool, significant 686 positive correlation with salinity was found for of gly ($R^2 = 0.466$, P < 0.0001), pro 687 ($R^2 = 0.174$, P = 0.0013), and arg ($R^2 = 0.131$, P = 0.0061), while asn had a weak but 688 significant negative correlation ($R^2 = 0.122$, P = 0.0083).

688 689

690 *3.7. Pigments and vitamins*

691

692 Astaxanthin was abundant in the copepods, and the levels were relatively similar

693 between 2000 and 2001 (Table 5, Fig. 8). The copepod astaxanthin content was lowest

- during the two weeks after mid-summer, with minimums of 321 and 362 μ g/g DW in
- 2000 and 2001, respectively. In 2000, the copepod astaxanthin level reached $832 \mu g/g$

696DW in mid-October, while in 2001 the levels continued to rise and peaked in mid-697November at 1422 μ g/g DW. In the zooplankton sample, astaxanthin was about 25%698of the average copepod pigment content in the corresponding year, while the rotifers699similarly contained 3.8% of the copepod astaxanthin content. In all cases, only free700astaxanthin was found, and no esters were observed. All Artemia samples were devoid701of astaxanthin, but contained canthaxanthin in the same ranges as copepod astaxanthin702(Table 5). Further, β-carotene was not detected in any of the samples.

703

Of the lipid-soluble vitamins, vitamin D₃ was either not detected in the copepod 704 705 samples or found in trace amounts (three of the samples). On average, it was therefore 706 considered below the detection limits of the method (Table 5). The zooplankton 707 sample was also free of vitamin D_3 , while levels in rotifers and Artemia were 0.9 and 708 0.7-1.8 µg/g DW, respectively. Further, vitamin A was found in low levels or beyond 709 quantification limits in the copepods. In many samples, vitamin A was even below 710 detection limit, particularly in 2001 (Table 5). Zooplankton and rotifers were also low 711 in Vitamin A (0.2 µg/g DW), and in Artemia realistic values for vitamin A were not 712 possible to quantify due to analytical problems (see section 2.3.4.). Vitamin E was 713 abundant in all samples (Fig. 8) and was dominated by the isomer, E_{α} (Table 5), 714 constituting between 90 and 100% of total vitamin E. No other isomers were detected in the zooplankton sample, while the remaining vitamin E in the copepods was E_{γ} and 715 E_{β} , the latter only observed in 2000. Both rotifers and <u>Artemia</u> showed low levels of 716 717 vitamin E_{γ} , and in addition <u>Artemia</u> displayed low but consistent levels of vitamin E_{δ} , 718 not found in the other feed types.

719

In the water-soluble vitamins, copepods showed high but variable levels of vitamin C
(Table 5, Fig. 8). Vitamin C in zooplankton, rotifers, and <u>Artemia</u> was within the range
of one standard deviation of the average values observed in the copepods. In copepods,
levels of thiamine was consistent and well above the quantification limit of the
method, with some variation between the years at different seasons (Fig. 8). Thiamine
was also abundant in zooplankton, rotifers and Artemia (Table 5). In contrast,

70(rite flering and have a seried amount fleretion limit of the mothed in series do (Fig. 8)
/26	riboriavin values varied around quantification limit of the method in copepods (Fig. 8),
727	zooplankton and rotifers, while <u>Artemia</u> had slightly higher levels (Table 5).
728	
729	
730	4. Discussion
731	
732	The biochemical composition of the copepods from Svartatjern was generally
733	characterised by substantial amounts of polar lipids, high levels of n-3 PUFA
734	(particularly DHA and EPA), protein with a diverse amino acid contribution in the
735	PAA profile (both for IAA and DAA, FAA dominated by few amino acids (gly, tau in
736	DAA and arg in IAA), high levels of astaxanthin, and considerable amounts of vitamin
737	C and vitamin E. In addition, compounds like β -carotene and vitamin D ₃ were
738	virtually absent in the copepods, while vitamin A and riboflavin were in the range of
739	trace limit concentrations. Further, the biochemical composition showed surprisingly
740	high stability between years or seasons within a year, despite large changes in copepod
741	species composition and environmental conditions (e.g. photoperiod, temperature and
742	salinity). However, the zooplankton sample from the Hyltro lagoon contrasts that of
743	Svartatjern copepods in containing more lipids with less PUFA and DHA. In addition,
744	the zooplankton had less protein, somewhat different FAA profile with more
745	dispensable FAA, along with lower astaxanthin and vitamin C content. These
746	discrepancies may most likely be explained by differences in phytoplankton
747	communities and densities, but also by a different composition of crustacean taxa,
748	since decapod larvae contributed to 32.4% of enumerated plankton in the zooplankton
749	sample. This may also clarify occurrence of slightly heavier individuals with higher
750	fraction of dry matter and ash in the zooplankton, probably because decapod zoeae are
751	more heavily armoured with carapace spines than copepods.
752	
752	An important question is to what extent Swartation represents natural acceptations and

An important question is to what extent Svartatjern represents natural ecosystems, and how this pond-like system may affect the biochemical composition of copepods? The Svartatjern pond system is managed by a specific protocol that implies fertilisation to boost primary production, mixing to prevent stratification, and emptying and refilling

according to renewal of copepod plankton from resting eggs (Naas et al., 1991; Næss, 757 758 1991). In this sense, copepods from Svartatiern may be regarded as "reared" copepods, 759 although reared on a diverse and natural assemblage of phytoplankton in a large 760 outdoor ecosystem. However, regarding dry matter, ash content, total lipids, and FAA 761 content the Svartatjern copepods were close to or within the mode values for other 762 copepods (reviewed by Båmstedt, 1986), but lower in protein content which on the 763 other hand was in accordance with data reported by Mæland et al. (2000). Protein 764 content may depend on the analytical method, and at present back calculation based on 765 PAA is regarded to be the most precise method for other larval prev (Hamre et al., 766 2007). Analyses of lipid class composition in copepods are mostly from high-latitude 767 oceanic calanoids (e.g. Calanus sp), which normally are rich in wax esters used as 768 energy source during overwintering and reproduction (Lee et al., 1971; Sargent and Falk-Petersen, 1988; Fraser et al., 1989). The copepod species included in the present 769 770 investigation are neritic calanoid species that do not overwinter as adults in the pond 771 system. Instead, they use resting eggs as a reproductive mode to ensure survival from 772 one generation to another during unfavourable conditions, e.g. during the seasonable 773 disruption of the production cycles (Næss, 1996). Storage of wax esters may therefore 774 not be required to the same extent as in the larger Calanus sp. The Svartatiern 775 copepods rather resembled naupliar and early copepodid stages of Calanus sp, which 776 are rich in structural phospholipids and contain TAG as main storage lipid (Sargent 777 and Henderson, 1986; Sargent and Falk-Petersen, 1988). In this respect, it should be 778 noted that nauplif and the young copepodid stages of Calanus sp are the primary prev 779 for larvae of many fish species.

780

Lipid content and composition in copepods have been found to be relatively diverse, and to vary with developmental stage, species, feed preference, latitude, season, and life cycle strategy (Båmstedt, 1986; Sargent and Falk-Petersen, 1988; Fraser et al., 1989; Norrbin et al., 1990; Støttrup, 2003). The Svartatjern copepod lipid composition may therefore be regarded as within the natural variation among copepods. Supporting this is also the high levels of certain fatty acids like 16:0, EPA, and DHA, which are in concordance with several other studies on neritic calanoid copepod species (Evjemo

788 and Olsen, 1997; Evejemo et al., 2003; Sørensen et al., 2007). Further, FAA in the 789 Svartatiern copepods was dominated by gly, tau, arg, ala, and pro, in a similar order 790 and magnitude as in other calanoid copepods (Båmstedt, 1986; Helland et al., 791 2003a,c). Astaxanthin, thiamine, riboflavin, vitamin C, and vitamin E were within the 792 ranges previously reported for copepods (Fisher et al., 1964; Hapette and Poulet, 1990; 793 Rønnestad et al., 1999a; Mæland et al., 2000). It may therefore be concluded that in 794 most biochemical indices, the Svartatiern copepods fell well within the variation 795 observed for copepods collected elsewhere. Thus, despite the manipulations imposed 796 for enhancement of primary production in the Svartatjern pond system, the copepods 797 preserved their similarities with wild copepods. Similar preservation of nutritional 798 composition has been reported from other zooplankton production systems (Mischke 799 et al., 2003). This indicates that the diverse phytoplankton and protozoan communities 800 in Svartatjern were conserved, preventing extreme lipid and fatty acid profiles which 801 can appear when one or two sub-optimal algal species are used in intensive copepod 802 culture systems (McKinnon et al., 2003). Copepods from Svartatjern have been used in 803 a several larval finfish studies, and have shown to support very high growth and 804 survival rates, and good juvenile quality (van der Meeren et al., 1993, 1994; Næss et 805 al., 1995; Conceição et al., 1997; McEvov et al., 1998; van der Meeren and Lønøv, 806 1998; Finn et al., 2002; Hamre et al., 2002; van der Meeren and Moksness, 2003). 807 Consequently, these copepods should represent a nutritionally adequate feed for many 808 larval fish species, and the data on biochemical composition may therefore serve as a 809 base for nutritional improvements of enrichment media used in culture of intensive 810 produced live feed for marine fish larvae, as well as for nutritional optimisation of 811 early weaning formulated diets.

812

Inadequate nutritional composition of intensive produced live prey has been
considered an important bottleneck in the production of high-quality juvenile marine
fish, and a substantial effort has been put into development of adequate live feed
enrichments (Støttrup, 2003; Marcus, 2005). Comparison between copepods, rotifers,
and <u>Artemia</u> data of the present study suggests a considerable potential for
improvement of enrichment emulsions. Recent advances in knowledge about lipid and

819 fatty acid requirements of marine fish larvae have pointed out the importance of 820 phospholipids, DHA, EPA, ARA, and the ratios of such PUFA for optimal lipid 821 digestion, normal larval development, larval survival and growth, and stress tolerance 822 (Olsen et al., 1991; Coutteau, 1997; Kanazawa, 1997; Sargent et al., 1999; Shields et 823 al., 1999; Izquierdo et al., 2001; Bell et al., 2003; Cahu et al., 2003; Hadas et al., 2003; 824 Støttrup, 2003). Compared to rotifers and Artemia, the Svartatjern copepods were 825 loaded with EPA and DHA. DHA was particularly abundant in the copepod nauplii, 826 indicating the importance of this fatty acid in the nutrition of young fish larvae whose 827 initial exogenous feed would be such prey. The high EPA/ARA ratio in the copepods 828 should be noted, as successful pigmentation during metamorphosis in flatfish larvae 829 may be dependent on this (Hamre et al., 2007). Considering the fraction of 830 phospholipids relative to total lipid, copepods were rich in phospholipids (57-63%) 831 compared to rotifers (40%) and particularly to Artemia (15-20%). However, taking 832 into account phospholipids per mg live prey biomass, differences were lesser (Table 833 2), probably due to the higher lipid content of the intensive prey types from 834 enrichment. Most enrichment oils for rotifers and Artemia are usually TAG, and 835 enhancing the phospholipid content of the prey by enrichment has turned out to be difficult (Rainuzzo et al., 1997; Harel et al., 1999). This is expressed as accumulation 836 837 of TAG with increasing lipid levels, with the potential for imbalances in both lipid 838 class and PUFA composition. Dietary phospholipids may enhance larval ingestion 839 (Koven et al., 1998), and phospholipids seem to be necessary for optimal lipid 840 transport and synthesis in the larval digestive system, as well as a number of cell 841 membrane and signalling functions (Bell et al., 2003; Cahu et al., 2003). Also the 842 relative abundance of different phospholipid classes may be of importance for larval 843 growth and development (Geurden et al., 1998). In the present data, both rotifers and 844 Artemia displayed many similarities with copepods when the relative composition of 845 the phospholipid profile was compared, indicating that structural lipids in the marine 846 food web are to some extent conservative. Quantitative deviations from the copepod 847 phospholipids were however evident, particularly in Artemia. More focus on 848 phospholipid enrichment of live feed and phospholipid supplement in formulated feed

849 is therefore necessary, with the goal to reach balanced levels of lipid classes and

- 850 PUFA as observed in copepods.
- 851

852 The gut system of young fish larvae has initially high assimilation capability of FAA 853 and low protein digestibility, with a gradual maturation of the proteolytic capacity 854 throughout ontogenesis (Cahu and Zambonino Infante, 2001; Rønnestad and 855 Conceição, 2005, Kvåle et al., 2007). FAA may serve as both energy substrate and 856 sustain protein synthesis in marine fish larvae (Rønnestad et al., 1999b; Wright and Fyhn, 2001; Rønnestad et al., 2003). The Svartatjern copepods were rich in FAA, and 857 858 the FAA concentration relative to DW was found to correlate with salinity. This 859 correlation may be explained by the need for copepods to use FAA in osmoregulation 860 (Båmstedt, 1986; Fyhn et al., 1993). Fish larvae may be very efficient in retaining and 861 absorbing FAA from the gut lumen, in particularly IAA (Conceição et al., 2002). 862 However, larval growth potential is in most cases very high, and daily weight gain 863 may exceed 20% even in coldwater species (van der Meeren et al., 1994; Finn et al., 864 2003). The observed FAA levels alone in larval live prey cannot sustain the amino 865 acid requirements surged by the protein deposition rate necessary to maintain such 866 high growth rates, and protein digestion must play a significant role in total amino acid 867 availability, absorption, and subsequent protein synthesis. Concordantly, recent studies 868 have shown that young marine fish larvae also are able to utilize peptide chains in 869 protein hydrolysates (Zambonino Infante et al., 1997; Cahu et al., 1999; Hamre et al., 870 2001), and that amino acids supplied in the diet in this form may reduce larval spinal 871 malformations (Cahu et al., 2003). Peptide digestion may be aided by high activity of 872 peptidases in young fish larvae (Cahu and Zambonino Infante, 2001). Although young 873 fish larvae have limited proteolytic capacity, access to peptide chains and amino acids 874 from dietary protein may be facilitated by autolysis of the ingested prey (Fyhn et al., 875 1993; Kolkovski, 2001). In this respect, Luizi et al. (1999) noted that copepods were 876 much more readily digested in Atlantic halibut larvae than Artemia. Furthermore, in 877 vitro digestibility studies with pancreatic enzymes chosen to mimic the conditions in 878 the larval intestine, show that water-soluble protein is more digestible than insoluble 879 protein (Tonheim et al., 2007). Both in intensive live feed and in copepods there are a

large fraction (approximately 50%) of water-soluble protein which has been suggested
to be highly bioavailable (Carvallo et al., 2003; Srivastava et al., 2006; Kvåle et al.,

- 882 2007; Tonheim et al., 2007).
- 883

Due to the high growth rate of fish larvae the demand for dietary amino acids for 884 885 protein accretion is especially high, and the supply of all amino acids, IAA as well as 886 DAA, may become critical for sustaining optimal growth. Thus, in juvenile rainbow 887 trout (Oncorhynchus mykiss) addition of crystalline DAAs (gln, gly, glu) to an 888 otherwise complete diet significantly increased growth rate and feed efficiency 889 (Schuhmacher et al., 1995). Such experiments have not been performed with marine 890 fish larvae although the suggestions have been made (Wright and Fyhn, 2001). Total 891 amino acid requirements may be related to larval growth rate which again may be 892 affected by a number of physical and biological factors (e.g. temperature, species, 893 larval size and age, and diet characteristics). In salmonids, deficiency in a single amino 894 acid (trp) during the first 4 weeks of exogenous feeding induced scoliosis (Akiyama et al., 1986). Other amino acids (thr, leu, arg, met, lys, and his) have been suggested as 895 896 limiting when rotifers or Artemia is used as feed for marine fish larvae (Conceição et 897 al, 1997, 2003; Aragão et al., 2004b). Deficiencies in these amino acids are mostly 898 inferred from imbalances between larval fish and prev profiles. However, these amino 899 acids were abundant in the Svartajern copepods, either in PAA, FAA, or both. With 900 some exceptions in rotifers, relative composition of amino acid profiles in protein 901 seems to be conserved between different plankton taxa. Since a substantial part of 902 dietary amino acids are in the form of PAA, amino acid deficiency may rather be a 903 matter of protein content in the feed, and how much of this protein that is digestible 904 and thereby available to absorption in the larval gut. In this respect, protein content in 905 rotifers and Artemia was lower compared to copepods. Further, dissimilarities in FAA 906 profiles of copepods, rotifers and Artemia were more pronounced, and FAA content 907 was highest in copepods. Use of live algae versus commercial enrichment products has 908 induced considerable variation in total amino acid profiles of rotifers and Artemia 909 (Aragão et al., 2004a). In absence of more detailed knowledge about specific amino

- 910 acid requirements, copepods might therefore be regarded as a baseline recipe for
- 911 protein, PAA, and FAA contents and profiles in feed for marine fish larvae.
- 912

913 Requirements for dietary micronutrients like pigments, vitamins, minerals, and trace 914 elements are little investigated in marine fish, and such studies are particularly scarce 915 for larval and early juvenile stages. Regarding minerals and trace elements, only iodine 916 was analysed from the Svartatiern copepods, as presented elsewhere (Moren et al., 917 2006). Compared to adult fish, the high growth rates and rapid organogenesis may 918 account for elevated micronutrient requirements and turnover during early developmental in fish (Lie et al., 1997), and recommendations suggested for adult fish 919 920 (e.g. in NRC, 1993) may therefore not be valid for younger life stages (Mæland et al., 921 2000). Levels of micronutrients found in copepods that sustain growth and normal 922 development, may be better indices for requirements in larval and juvenile marine fish, 923 and the present study is an attempt to provide such baseline data. 924

925 Regarding pigments, the consistent high levels of astaxanthin in the copepods suggest 926 that this compound should receive more attention in larval fish nutrition. Together 927 with canthaxanthin commonly found in Artemia, astaxanthin, lutein, and β -carotene 928 belong to the carotenoid family that may serve as precursors for vitamin A in fish 929 (Bendich and Olson, 1989; Christiansen and Torrissen, 1996; Moren et al., 2005; 930 Palace and Werner, 2006). Since β -carotene was not detected in the copepods, 931 astaxanthin and possibly also lutein may be important provitamin A compounds in 932 such plankton (Rønnestad et al., 1998). Astaxanthin have also demonstrated profound 933 antioxidant properties, particularly as a coantioxidant working synergistically with 934 vitamin E in suppressing lipid peroxidation (Bell et al., 2000). Antioxidant action on 935 active oxygen radicals in marine organisms has also been suggested for a number of 936 other carotenoids, including canthaxanthin (Shimidzu et al., 1996), and carotenoids 937 enhanced survival and reduced lipid peroxidation in Japanese flounder larvae 938 (Okimasu et al., 1992). Carotenoids may therefore assist the enzymatic antioxidant 939 system in fish, which is already functional during early larval stages (Peters and Livingstone, 1996; Mourente et al., 1999a; Martínez-Álvarez et al., 2005). Data on 940

biological activities of pigments in fish are scarce, but effects of astaxanthin on skin
and muscle coloration are well documented (Torrissen et al., 1989; Chatzifotis et al.,
2005). Low intake of astaxanthin may reduce growth in salmonids (Christiansen and
Torrissen, 1996), and maternal deficiency may significantly reduce transfer of
astaxanthin to the fish eggs and possibly erode survival in the larval stages (Pickova et
al., 1998).

947

Use of dietary carotenoids may be a safe way to provide vitamin A in larval fish, as 948 949 dietary surplus of vitamin A or its derivates (retinoids) may have detrimental effects on normal bone development (Dedi et al., 1995; Cahu et al., 2003). Retinol and other 950 951 retinoids seem to be very low or absent in copepods, and the hidden source of vitamin 952 A in larval fish is probably carotenoids, which are enzymatically cleaved to form 953 retinoids in fish (Moren et al., 2005). In this way, carotenoids may be converted to 954 vitamin A, depending on the retinoid and protein status of the animal (Bendich and 955 Olson, 1989). Similarly to retinoids, β -carotene also seems to be very low or absent in 956 copepods, which may explain why fish, compared to land vertebrates, display less 957 specificity for this carotenoid as a vitamin A source (Palace and Werner, 2006). 958 However, conversion of β -carotene to retinols at a higher rate than with other 959 carotenoids has been demonstrated in juveniles of Atlantic halibut (Moren et al., 960 2004a), although quantification of this conversion remains to be determined for larval 961 fish. Under the assumption that larval halibut has a vitamin A requirement in the same 962 range as juvenile halibut, astaxanthin levels in copepods or canthaxanthin in Artemia 963 could cover the need for this vitamin (Moren et al., 2004a,b). Alternatively, covering 964 vitamin A requirements for larval fish in terms of dietary retinoids needs more 965 attention, since certain retinoids may inflict disruptive actions on fish physiology, 966 development, growth, and survival (Woodward, 1994; Dedi et al., 1995; Furuita et al., 967 2001; Haga et al., 2002; Moren et al., 2004a; Palace and Werner, 2006), including 968 teratogenic effects on bone development at the level of gene expression (Cahu et al., 969 2003; Hamre et al., 2007; Lall and Lewis-McCrea, 2007). 970

971 Another vitamin not found in the copepods was cholecalcipherol (vitamin D_3). This 972 was unexpected, as this vitamin may play important roles in calcium and phosphorous 973 metabolism and affect bone formation and remodelling in vertebrates. Vitamin D_3 is 974 the main storage form in the liver of marine teleosts, and may be converted to 25-975 hydroxyl vitamin D₃ isomers in various fish tissues (Takeuchi et al., 1991; Graff et al., 976 1999; Holick, 2003; Lall and Lewis-McCrea, 2007). However, data on effects of 977 dietary vitamin D₃ in larval fish are very scarce. In a recent study of young juvenile 978 Japanese flounder, hypermelanosis on the blind side and vertebral deformities have 979 been reported when dietary levels exceed 5 μ g/g vitamin D₃ or 0.5 μ g/g 1,25(OH)₂ 980 vitamin D₃ (Haga et al., 2004). Copepods may contain vitamin D₃ levels below the 981 analytical detection and quantification limits, or they may contain precursors as the 7-982 dehydrocholesterol (7-DCH), which is the provitamin responsible for vitamin D_3 983 production in the skin of terrestrial vertebrates under UV light irradiation. In this 984 respect, several studies agree on that copepods displays either lack of vitamin D_2 and D₃ while 7-DCH is detected in reasonable amounts (Geiger, 1958; Takeuchi et al., 985 1991; Kenny et al., 2004). Approximately 1.4% of the Svartatjern copepod DW was 986 987 cholesterol and sterol esters, but 7-DCH was not specifically analysed for. In adult 988 fish, both photo-conversion in the skin and enzymatic dark-transfer of 7-DCH in the 989 liver to vitamin D₃ has been reported (Holick, 2003; Blondin et al., 1967), but also 990 disputed (Takeuchi et al., 1991). No data have been presented on this matter for fish 991 larvae, and this calls for further exploration. If fish is able to convert this provitamin to 992 vitamin D₃ it may account for 7-DCH as a potential important vitamin D source in 993 most stages of planktivorous fish, and explain the paradox of vitamin D_3 enrichment at 994 this trophic level in the marine food web. Photo-conversion implies that such fish has 995 to reside close to daylight at the surface, which e.g. fish larvae or pelagic schooling 996 fish often do. It also means that indoor rearing of larval fish in absence of UV-light 997 might require dietary vitamin D_3 , which is actually supplied in rotifers and Artemia 998 due to use of fish oils in the enrichment emulsions (Table 5). Since 7-DCH occurs 999 naturally in fish liver (Takeuchi et al., 1987), enzymatic dark-conversion is an 1000 intriguing aspect that also needs further investigation. However, with the enormous 1001 potential of prey ingestion in larval fish, bioaccumulation from ingested zooplankton

1002 containing traces of vitamin D_3 may not be ruled out as a sufficient source. Analogue 1003 to retinoic acid, vitamin D isomers may be involved in regulation of gene transcription 1004 in a ligand-dependent manner through their interaction with specific DNA sequences 1005 (Crisp et al., 1998; Hamre et al., 2007), and should therefore be added to the larval diet 1006 with care as long as larval storage capabilities and metabolic pathways are unknown. 1007

The high fraction of phospholipids and PUFA in the copepods may require substantial 1008 1009 protection against oxidation by free radicals. The main function of vitamin E is to 1010 reduce peroxyl radicals in membrane lipids and prevent the chain reaction leading to 1011 lipid peroxidation, and vitamin E is therefore crucial for normal development of 1012 tissues, including bone and cartilage (Lall and Lewis-McCrea, 2007). Vitamin E may 1013 also inhibit the oxidations induced by the electronically excited singlet oxygen, and 1014 have a number of other effects as reviewed by Kamal-Eldin and Appelqvist (1996) and 1015 Azzi and Stocker (2000). Due to the lipid protective activity, it is not surprising that 1016 the copepods were rich in vitamin E and other synergists like carotenoids and vitamin 1017 C, the latter being important in regenerating the antioxidative properties of vitamin E 1018 by converting the oxidised form (α -tocopheroxyl) to α -tocopherol (Hamre et al., 1997) 1019 which is the most abundant and bioactive form of the vitamin E isomers (Kamal-Eldin 1020 and Appelqvist, 1996, Hamre et al., 1998). Rapid growth and formation of cell 1021 membranes in larval fish count for high PUFA requirements, with the risk of high 1022 oxidative stress. Dietary vitamin E in larval fish should therefore relate to PUFA 1023 intake (Martínez-Álvarez et al., 2005), and will as a free radical scavenger support the 1024 antioxidation enzyme systems encountered in fish larvae (Mourente et al., 1999b; 1025 Tocher et al., 2002). Due to the high metabolic turnover in larval fish, the specific 1026 vitamin E requirements suggested by NRC (1993) for older stages may not be 1027 appropriate, and higher levels have been suggested (Lie et al., 1997). However, 1028 restoration of vitamin E by other micronutrients implies that body contents of 1029 regenerative compounds and their dietary intake, together with restoration rates need 1030 to be accounted for in study of larval vitamin E deficiency. The fact that other 1031 micronutrients also effectively contribute as antioxidants makes assessment of specific 1032 larval α -tocopherol requirements even more challenging. Vitamin E levels in copepods

were low compared to rotifers and <u>Artemia</u>, but high levels of other synergistic
compounds like astaxanthin and vitamin C are suggesting that the copepods might
provide sufficient antioxidant potential for fish larvae. The high levels of vitamin E
provided through enrichment emulsions may therefore not be necessary, but more
research should be carried out to determine requirements and metabolic pathways of
tocopherols in larval fish.

1039

The vitamin C in the copepods was high but variable. Copepod vitamin C content 1040 1041 originates from dietary phytoplankton since biosynthesis of ascorbic acid does not 1042 occur in copepods (Hapette and Poulet, 1990). The omnivorous nature of many 1043 copepod species may explain some of the variation in copepod vitamin C levels, 1044 induced by variations in algal vitamin C content, copepod gracing, and food selection. Most fish cannot synthesise vitamin C, which is a strong reducing agent that can be 1045 1046 restored enzymatically, and that acts as a cofactor in production of procollagen, a 1047 precursor of collagen (NRC, 1993). Vitamin C is therefore important for development 1048 of connective tissue, wound repair, and formation of bone matrix. Vitamin C may also 1049 enhance immune function (Woodward, 1994), and deficiencies may affect hepatocyte 1050 cellular compartmentation (Merchie et al., 1997). Vitamin C requirements in larval and 1051 early juvenile fish have been indicated in the range of 20 and 500 µg/g DW, while in 1052 some species enhanced growth, increased stress tolerance, and reduced incidence of 1053 opercular deformities occurred at levels up to 1750 µg/g DW (Merchie et al., 1997; 1054 Gapasin et al., 1998). Opercular abnormalities are distortion of gill filament cartilages 1055 resulting from de-calcification, and are characteristic of scorbutic fish (Cahu et al., 1056 2003). The above-mentioned differences in dietary requirements between species or 1057 stage of development may be explained by metabolic activity (Merchie et al., 1997). 1058 The Svartatjern copepods should therefore have no problem in supporting dietary 1059 needs of vitamin C for both temperate and cold-water larval and juvenile marine fish 1060 species. 1061

1062 The copepod thiamine content resembled rotifer levels reached after more than 10 h of

1063 enrichment on the algae <u>Isochrysis galbana</u> (Lie et al., 1997), and corresponded to
levels in other copepods (Mæland et al., 2000). No data on thiamine deficiency or 1064 1065 requirements of larval marine fish have to our knowledge been reported. Thiamine 1066 combines with pyrophosphate in a coenzyme used for oxidative decarboxylation of α -1067 keto acids and transketolase reaction in the pentose shunt, and therefore relate closely 1068 to energy production (NRC, 1993; Woodward, 1994). In fish, deficiency in thiamine 1069 has been associated with the M74 and Cayuga syndromes in salmonids, leading to high 1070 mortality during early life stages in wild fish (Fisher et al., 1996; Åkerman et al., 1998; 1071 Pickova et al., 1998; Ketola et al., 2000). In thiamine deficient farmed fish, 1072 malfunctioning of the nervous system, including loss of equilibrium accompanied by 1073 whirling, melanotic appearance, inability to feed, progressive weakness, and paralysis 1074 were described by Woodbury (1943). In other vertebrates, thiamine deprivation causes 1075 pan-necrosis affecting the nuclei of the brain stem and diencephalons (Dreyfus and 1076 Victor, 1961). The use of Svartatjern copepods for successful rearing of cold-water 1077 species accounts for satisfactory thiamine levels in these copepods, which is above the 1078 levels suggested for adult fish by NRC (1993), but research is needed to verify larval 1079 requirements.

1080

1081 The observed riboflavin levels in the copepods exceeded the recommended minimum 1082 requirements for fish, including for juveniles that do not seem to have elevated needs 1083 for riboflavin compared to older fish (NRC, 1993; Serrini et al., 1996; Bjørnstad et al., 1084 2002; Deng and Wilson, 2003). However, most of these data are collected from studies 1085 of freshwater or anadromous fish species, and no investigations on riboflavin 1086 requirements of marine fish larvae have been published. Through its involvement in 1087 two coenzymes, riboflavin functions as electron mediator in oxidation-reduction 1088 reactions involved in metabolism of keto-acids, fatty acids, and amino acids in the 1089 mitochondrial electron system (NRC, 1993). Symptoms of riboflavin deficiency may 1090 be species-specific, and include elevated mortality, reduced weight gain, rapid 1091 opercular movements, aneroxia, lethargy, dark or light body colour, severe fin erosion, 1092 cataracts, photophobia, reduced hepatic D-amino acid oxidase activity, and 1093 haemorrhages (Woodward, 1984; NRC, 1993; Serrini et al., 1996; Deng and Wilson, 1094 2003). The riboflavin levels in the Svartatjern copepods were lower than in the rotifers

and <u>Artemia</u>, but slightly above the levels presented for the copepod <u>Temora</u>
 <u>longicornis</u> by Mæland et al. (2000). Since no riboflavin-related deficiency symptoms

1097 have been observed when feeding the copepods to larval coldwater fish, use of rotifers

- 1098 and <u>Artemia</u> should therefore assumingly cover the requirements. But controlled
- 1099 experiments to verify riboflavin requirements in marine fish larvae are still lacking.
- 1100
- 1101

1102 **Conclusions**

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1104 From present knowledge about nutritional requirements of marine fish larvae, small 1105 neritic calanoid copepods display a macronutrient composition that seems to satisfy 1106 the demands of the larvae. In particular, this comprises medium protein and high FAA 1107 contents with balanced amino acid profiles, medium to low lipid content, high 1108 fractions of phospholipids, DHA, and EPA, with optimal ratios regarding DHA/EPA 1109 and EPA/ARA. The low content of wax esters resembles nauplii and young copepodid 1110 stages of Calanus sp, which are a major component of the larval feed in many marine 1111 ecosystems. Among the micronutrients, copepods are rich in pigment, and particularly 1112 astaxanthin, which may be an important source of retinoids for larval fish since β -1113 carotene and vitamin A are scarce in copepods. Absence of vitamin D_3 in the copepods 1114 may indicate dietary precursors as source of cholecalcipherol in larval fish, but data on 1115 potential precursors are lacking. In contrast, copepods are rich in vitamin E and 1116 ascorbic acid, which together with astaxanthin are pointing to high antioxidative 1117 capacity needed to protect against peroxidation of membrane lipids. Vitamin C was 1118 most abundant, making the copepods particularly suitable for fish larvae with a high 1119 growth potential. The copepod content of thiamine and riboflavin may be sufficient to 1120 sustain larval development in marine fish, but data on larval requirements are absent in 1121 the literature. High metabolism linked to the rapid growth rates often displayed by 1122 young marine fish larvae may account for elevated micronutrient needs beyond what 1123 are suggested for older fish. Determination of optimal larval requirements are lacking 1124 for many of the micronutrients, and such data should be collected since insufficient 1125 dietary supply of some micronutrients already has demonstrated impairment of normal

larval development. Copepods have successfully been applied as feed for marine fish 1126 1127 larvae, also in intensive rearing systems. Since copepods are the principal prey of 1128 marine fish larvae, this suggests specific larval adaptations to universal traits of 1129 copepod biochemical composition. Thus, evolution of the larval digestive and 1130 metabolic systems may have set limits to tolerance of nutritional variability in the 1131 larval prey, limits that were surpassed when Artemia and rotifers were introduced in 1132 intensive production of marine fish juveniles. Alteration of nutritional composition of 1133 rotifers, Artemia, and formulated feed should therefore be made in the direction of 1134 copepods, and the present data provide a comprehensive outline of this direction. 1135 M/L 1136 1137 **Aknowledgements** 1138 We want to thank Tore Håkon Næss for identification of copepod species and stages, 1139 and Nils Bernt Andersen for phytoplankton enumeration and determination. Also 1140 thanks to Svanhild Lohne Gokstad for nutrient analysis, Maria Sula Evjen for protein 1141 1142 and amino acid analyses, Vibecke Asphaug for organising the sample distribution at 1143 NIFES and analyses of the B vitamins, Kjersti Ask, and Idun Kallestad for analyses of 1144 the lipid soluble vitamins. The work was financially supported by the Norwegian 1145 Research Council (project no.138379/120) as a co-operative project between Institute 1146 of Marine Research and the Atlantic halibut fry producer Norsk Kveite AS (named 1147 Austevoll Marin Yngel AS in 2000). 1148 1149 1150 References 1151 1152 Akiyama, T., Murai, T., Nose, T., 1986. Oral administration of serotonin against spinal 1153 deformity of chum salmon fry induced tryptophan deficiency. Bull. Jpn. Soc. Sci. 1154 Fish. 52, 1249-1254. 1155 Aragao, C., Conceicao, L.E.C., Dinis, M.T., Fyhn, H.J., 2004a. Amino acid pools of

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1607 <u>Figure legends</u>:

- 1608 Figure 1. Densities of major phytoplankton groups and protozoans in the Svartatjern
- 1609 pond during collection of copepods in 2000 and 2001. The grey areas indicate when
- 1610 the pond was drained and refilled during the summer season. Note that the ordinate is
- 1611 logarithmic. Values indicated by lines below the abscissa equal zero.
- 1612
- 1613 Figure 2. Relative abundance (percent of numbers) of the copepods <u>Eurytemora</u>
- 1614 <u>affinis</u>, <u>Centropagus hamatus</u>, and <u>Acartia grani</u>, the cladoceran <u>Podon</u> sp., and
- 1615 miscellaneous zooplankton (including other copepods) in the samples collected from

1616 the Svartatjern pond in 2000 and 2001.

1617

1618 Figure 3. Dry weight of individual copepods, dry matter (% of wet weight), and ash

1619 content (% of dry weight) from the 2000 and 2001 samples collected in the Svartatjern

1620 pond. Note that the ordinate is broken in the lower panel.

1621

Figure 4. Total lipid content relative to dry weight (DW) and relative abundance of
major lipid classes in the copepod samples from the Svartatjern pond in 2000 and
2001. See table 2 for explanation of abbreviations.

1625

Figure 5. Fatty acid ratios and major fatty acids extracted from total lipids of copepods
samples from the Svartatjern pond in 2000 and 2001. See table 2 for explanation of
abbreviations.

1629

1630 Figure 6. Protein fraction relative to dry weight (DW) and calculated from protein-

bound amino acids (PAA), fraction of indispensable PAA, and relative abundance of

amino acids with a major contribution to PAA in the copepod samples from the

1633 Svartatjern pond in 2000 and 2001. See table 3 for explanation of abbreviations. Note

1634 that the right ordinate is broken in the upper panel.

1635

1636 Figure 7. Fraction of free amino acids (FAA) relative to dry weight (DW), fraction of

1637 indispensable FAA, and relative abundance of amino acids with a major contribution

- to FAA in the copepod samples from the Svartatjern pond in 2000 and 2001. See table 1638
- 1639 4 for explanation of abbreviations.
- 1640
- 1641 Figure 8. Content of pigments and vitamins relative to dry weight (DW) in the 2000
- 1642 and 2001 copepod samples from the Svartatjern pond. Dotted line in lower panel
- 1643 indicates quantification limit for riboflavin at the present analytical method.

μ^κ the pre.

1644 Table 1

1645 Hydrographical data from the Svartatjern pond during collection of copepods.

1646

	20	000		20			
	Mean \pm SD	Min.	Max.	$Mean \pm SD$	Min.	Max.	
Temperature (°C)	14.2 ± 2.9	8.5	19.1	14.1 ± 3.7	7.1	19.3	
Salinity (‰)	25.2 ± 2.9	21.1	31.2	26.3 ± 3.2	19.9	31.4	
Oxygen (% saturation)	105 ± 19	77	145	98 ± 43	15	160	
pH ^a				8.2 ± 0.5	7.4	9.1	
Secci depth (m)	1.4 ± 0.4	1.0	2.2	1.7 ± 0.6	1.0	3.3	
Nitrate (µM)	1.8 ± 1.2	0.0	4.5	4.5 ± 5.0	0.5	15.3	
Phosphate (µM)	0.6 ± 0.3	0.2	1.1	0.7 ± 0.3	0.2	1.3	
Silicate ^a (µM)				3.4 ± 4.9	0.3	19.0	

1647

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SCI R Not

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1648 Table 2

1649 Individual size (dry weight: DW), dry matter content (% of wet weight: WW), ash 1650 content, and lipid components from copepods, copepod nauplii, zooplankton 1651 (copepods and decapod zoeae), rotifers, and <u>Artemia</u> (1-day or 3-day after hatching).

- 1652 Data are given as mean \pm SD when number of samples >1. Values below detection
- 1653 limits of the analytical method are denoted n.d.
- 1654

	_		Svartatjern		Hyltro Zoopl.	Intensive live feed			
	Abbr.	Copepods	Copepods	Cop. nauplii		Rotifers	Artemia		
		2000	2001	2001	2001	IMR ^a	1-day ^b	1-day ^c	3-day ^c
Individual size, Dry matter	•		C)					
and Ash									
Number of samples	N	30	26	3	1	1	1	1	1
Dry weight (µg/individual)	DW	9.4 ^A ± 2.5	8.1 ± 2.7	$0.63^{\rm B}{\pm}~0.7$	9.9	0.61	2.1	2.1	2.5
Dry matter (% of WW)	DM	14.9±1.1	15.3 ± 1.5	15.2 ± 1.9	17.7	13.2	10.2	10.8	8.9
Ash content (% of DW)	ASH	10.3±1.2	10.5 ± 1.3	9.9 ± 0.5	15.3	9.6	10.4	9.6	9.5
Total lipid (µg/mg DW)	TL	111±35	108 ± 21	$86^{B} \pm 12$	143	154	254	243	249
Neutral lipids (µg/mg DW)	NL	49.4±23.4	45.4±13.3	32.6±13.5	91.5	92.5	215.0	193.4	206.0
Sterol esters+Wax esters	SE+WE	1.5 ± 1.5	1.1 ± 1.2	1.3 ± 0.5	22.6	11.6	n.d.	1.3	n.d.
Triacylglycerol	TAG	26.3±19.5	22.1 ± 13.1	14.0 ± 13.3	42.1	60.6	195.9	167.9	178.4
Free fatty acids	FFA	3.3 ± 2.1	3.3 ± 1.6	3.9 ± 2.1	6.6	6.9	4.4	5.8	8.9
Cholesterol	С	14.5 ± 6.3	13.3 ± 3.5	$9.5^{\mathrm{B}} \pm 3.6$	15.5	11.6	14.6	18.4	18.7
Monogalactosides+Cerebrocides	MGDG+CB	$1.6^{A} \pm 1.0$	2.7 ± 2.6	2.1±1.3	1.3	n.d.	n.d.	n.d.	n.d.
Digalactosides+Sulfolipids	DGDG+SL	2.3 ± 1.1	2.9±1.5	1.8 ± 0.8	3.4	1.9	n.d.	n.d.	n.d.
Polar lipids (µg/mg DW)	PL	61.9±16.8	62.6 ± 14.4	53.7±2.7	51.5	61.1	39.2	49.9	43.3
Phosphatidylethanolamine	PE	19.9 ± 5.8	20.4 ± 4.8	17.3 ± 2.1	15.2	21.0	12.3	16.8	14.3
Cardiolipin	CL	5.7 ± 2.0	5.8 ± 1.6	5.2 ± 0.3	2.5	3.1	2.2	3.0	2.5
Phosphatidylglycerol	PG	2.2 ± 2.3	1.5 ± 1.0	0.8 ± 0.2	1.4	1.5	0.5	0.8	0.5
Phosphatidylinositol	PI	3.9 ± 1.7	4.2 ± 1.4	5.1 ± 2.1	5.6	10.6	3.8	5.1	4.2
Phosphatidylserine	PS	$5.5^{A} \pm 1.5$	6.6 ± 2.0	6.4 ± 0.7	4.8	5.3	3.1	4.1	3.2
Phosphatidylcholine	PC	$20.0{\pm}6.0$	19.4 ± 5.3	15.0 ± 1.3	19.0	18.5	16.4	18.6	17.4
Lysophosphatidylcholine+Sphingomyelin	LPC+SM	4.7±1.6	4.8 ± 1.6	3.9 ± 0.2	2.9	1.1	0.9	1.4	1.1
Fatty acids (% of total lipid)									
Myristic acid	14:0	$3.4^{A} \pm 1.7$	1.7 ± 1.1	1.3 ± 0.8	3.8	6.7	1.7	1.5	2.4
Palmitic acid	16:0	14.5 ± 1.9	14.4 ± 1.4	13.7 ± 2.5	14.1	19.7	14.9	14.4	15.8
Palmitoleic acid	16:1(n-7)	3.4 ± 1.8	4.4 ± 4.7	1.8 ± 1.4	7.6	9.2	4.8	1.0	3.0
Stearic acid	18:0	3.5 ± 1.0	3.7 ± 0.7	3.9 ± 1.0	4.1	3.9	5.0	5.0	5.4

Oleic acid	18:1(n-9)	2.3 ± 1.1	2.6 ± 1.4	$1.3^{\rm B}{\pm}~0.7$	7.3	7.8	23.3	22.8	17.8
Vaccenic (Asclepic) acid	18:1(n-7)	$2.7{\pm}~0.6$	2.9 ± 0.7	$2.0^{\rm B}{\pm}~0.5$	3.1	4.9	5.5	6.3	5.4
Linoleic acid	18:2(n-6)	$1.5^{\rm A}{\pm}~0.5$	2.3 ± 0.7	$1.5^{\rm B}{\pm}~0.5$	2.2	15.3	6.6	5.8	4.2
α-Linolenic acid	18:3(n-3)	$1.9^{\mathrm{A}} \pm 1.0$	2.4 ± 1.1	$1.5^{\rm B}{\pm}~0.9$	1.4	1.2	12.2	16.2	10.2
Stearidonic acid	18:4(n-3)	$2.3^{A} \pm 1.4$	4.1 ± 2.9	4.5±5.7	5.2	2.0	2.8	3.2	1.7
Arachidonic acid (ARA)	20:4(n-6)	0.8 ± 0.5	0.9 ± 0.7	0.6 ± 0.3	1.6	1.9	2.0	2.0	3.2
Eicosapentaenoic acid (EPA)	20:5(n-3)	17.4 ± 3.1	16.2 ± 3.4	16.3 ± 6.4	16.4	7.1	7.8	7.8	9.2
Docosahexaenoic acid (DHA)	22:6(n-3)	34.4 ± 4.6	32.9 ± 6.8	$40.5^{\mathrm{B}} \pm 2.4$	17.3	12.4	10.6	11.1	20.0
Other Saturated fatty acids		3.1 ± 1.1	3.3 ± 1.2	3.9 ± 2.7	9.7	n.d.	n.d.	n.d.	n.d.
Other Monounsaturated fatty acids		$3.7^{A} \pm 1.0$	2.8 ± 0.9	2.7±1.2	2.0	4.2	1.2	1.5	1.7
Other Polyunsaturated fatty acids		5.1±1.2	5.4±1.5	4.6±1.0	4.2	3.8	1.6	1.4	n.d.
Total amounts of fatty acid groups (%)									
Saturated fatty acids	SFA	$24.6^{A} \pm 2.9$	23.1 ± 2.2	$22.7{\pm}2.9$	31.7	30.3	21.6	20.9	23.7
Monounsaturated fatty acids	MUFA	12.1±2.1	12.7 ± 6.1	7.8 ± 3.2	20.1	26.1	34.8	31.6	27.8
Polyunsaturated fatty acids	PUFA	63.3 ± 3.7	64.2 ± 6.8	$69.4{\pm}~5.8$	48.3	43.7	43.6	47.5	48.5
Highly unsaturated (n-3) fatty acids	DHA+EPA	$51.8^{A} \pm 4.5$	49.1±6.8	$56.8^B{\pm}6.8$	33.6	19.4	18.4	19.0	29.2
Fatty acid ratios	(n-3)/(n-6)	$11.3^{A} \pm 2.7$	9.8±2.5	$12.5^{\mathrm{B}} \pm 3.0$	7.0	1.5	3.9	4.2	5.5
	DHA/EPA	2.1 ± 0.5	2.2 ± 1.0	2.8 ± 1.3	1.1	1.7	1.4	1.4	2.2
	EPA/ARA	24.7±9.2	23.2 ± 10.1	27.7±4.0	10.3	3.7	4.0	4.0	2.9

^a Institute of Marine Research: rotifers grown on Rotimac and <u>Isochrysis galbana</u> algae.

^b Institute of Marine Research: <u>Artemia</u> enriched with DC-DHA Selco.

1657 ^c Austevoll Marin Yngel AS: <u>Artemia</u> fed DC-DHA Selco and Algamac 2000.

1658 ^A Significant difference between copepods from 2000 and 2001.

¹⁶⁵⁹ ^B Significant difference between copepod nauplii and copepods from 2001.

1660 Table 3

1661 Content of protein (P) and protein-bound amino acids (PAA) in copepods, zooplankton

1662 (copepods and decapod zoeae), rotifers, and <u>Artemia</u> (1-day or 3-day after hatching).

1663 Values are relative to dry weight (DW) and are given as mean \pm SD when number of

1664 samples >1. Values below detection limits of the analytical method are denoted n.d.

1665 The subscripts "w" and "c" indicate data given as weight and concentration,

- 1666 respectively.
- 1667

	_	Svarta	Hyltro	Intensive live feed				
	Abbr.	Copepods	Copepods	Zoopl.	Rotifers	I	Artemia	
		2000	2001	2001	IMR ^a	1-day ^b	1-day ^c	3-day ^c
Number of samples	Ν	30	26	1	1	1	1	1
Protein ^d (µg/mg DW)	Р	$382.6^{A} \pm 25.5$	565.4 ± 40.0	366.3	243.4	287.9	309.2	326.2
PAA in weight ^e (μ g/mg DW)	PAAw	$443.6^{A} \pm 41.6$	412.6 ± 41.0	302.5	247.7	277.5	293.8	367.6
Indispensable amino acids (µg/mg DW)	IAA _w	$201.3^{\text{A}} {\pm}~16.8$	$189.6{\pm}20.8$	141.8	120.0	133.3	140.3	175.2
Indispensable amino acids (%)	IAA _W /PAA _W	45.4±1.5	$45.9{\pm}0.9$	46.9	48.4	48.0	47.8	47.7
Indispensable to dispensable ratio	IAA _w /DAA _w	0.83 ± 0.05	$0.85{\pm}0.03$	0.88	0.94	0.92	0.91	0.91
PAA concentration (µmoles/mg DW)	PAAc	$4.1^{A} \pm 0.4$	3.8 ± 0.4	2.8	2.3	2.5	2.7	3.4
Indispensable amino acids (μ moles/mg DW)	IAAc	$1.7^{\mathrm{A}} \pm 0.1$	1.6 ± 0.2	1.2	1.0	1.1	1.2	1.4
Indispensable amino acids (%)	IAA _c /PAA _c	$40.4^{\rm A}{\pm}~1.5$	$41.3 {\pm}~0.9$	41.5	43.7	43.3	42.8	42.7
Indispensable to dispensable ratio	IAA _c /DAA _c	$0.68^{\rm A}{\pm}~0.04$	$0.70{\pm}0.03$	0.71	0.78	0.76	0.75	0.75
Indispensable amino acids (nmoles/mg DW)							
Leucine	LEU	$349.0^{A} \pm 38.5$	$320.5{\pm}33.8$	246.4	230.1	225.6	295.2	237.0
Valine	VAL	$291.8^{A} \pm 36.4$	$253.0{\pm}24.9$	200.7	160.1	175.1	233.6	183.9
Lysine	LYS	241.3 ± 43.0	231.1 ± 34.7	163.8	136.6	149.0	222.6	166.7
Isoleucine	ILE	$209.6^{A} \pm 26.3$	$187.3\!\pm20.1$	146.5	143.3	137.6	186.8	148.7
Arginine	ARG	$121.7^{\text{A}} \pm 27.6$	161.7 ± 14.1	126.4	83.7	115.7	149.4	108.8
Phenylalanine	PHE	$154.4^{\text{A}} \pm 18.7$	$143.4{\pm}15.7$	112.1	114.9	105.7	138.8	111.8
Threonine	THR	$128.7^{\text{A}} \pm 13.1$	$120.0{\pm}~14.1$	95.1	70.7	86.3	114.6	89.1
Methionine	MET	$122.3^{A} \pm 13.4$	77.7 ± 38.1	69.4	47.6	56.8	40.5	63.2
Histidine	HIS	$53.7^{A} \pm 26.2$	63.7 ± 10.1	10.5	6.9	43.2	60.1	45.9
Tryptophan	TRP	44.6 ^A ± 84.1	0.7 ± 2.7	n.d.	4.4	2.3	7.0	3.5
Dispensable amino acids (nmoles/mg DW)								
Glutamic acid+Glutamine	GLU+GLN	$577.8^{\rm A} \pm 66.4$	$505.2{\pm}~52.7$	384.7	325.0	325.4	427.4	328.2
Aspartic acid+Asparagine	ASP+ASN	411.1 ± 43.4	$432.3\!\pm54.0$	335.8	293.0	271.9	363.5	282.4
Alanine	ALA	$463.4^{A} \pm 54.8$	$392.3{\pm}38.2$	284.7	189.0	230.2	306.9	252.4
Glycine	GLY	$441.2^{\text{A}} \pm 94.7$	352.1 ± 49.7	286.1	181.0	224.7	321.8	245.8

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Serine	SER	$204.9^{\text{A}} \pm 22.3$	190.7±19.2	152.0	136.8	136.7	186.3	143.7
Proline	PRO	252.0 ^A ± 59.9	$186.8{\pm}24.8$	164.4	134.4	157.4	217.4	200.8
Tyrosine	TYR	$122.2^{\text{A}} \pm 36.1$	154.5 ± 16.9	40.5	19.9	88.7	109.5	89.5

^a Institute of Marine Research: rotifers grown on Rotimac and <u>Isochrysis galbana</u> algae.

^b Institute of Marine Research: <u>Artemia</u> enriched with DC-DHA Selco.

1670 ^c Austevoll Marin Yngel AS: <u>Artemia</u> fed DC-DHA Selco and Algamac 2000,

^d Protein determined with the Bovine serum albumin method of Lowry et al. (1951) and Rutter (1967).

1672 ^e PAA in weight are calculated as protein (i.e. from the amino acid mole weight subtracted by the mole weight

- 1673 of a water molecule, which resembles the PAA before hydrolysis).
- ^A Significant difference between copepods from 2000 and 2001.

CCC CCC

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1677 Table 4

1678 Free amino acids (FAA) in copepods, zooplankton (copepods and decapod zoeae),

1679 rotifers, and Artemia (1-day or 3-day after hatching). Values are relative to dry weight

1680 (DW) and are given as mean \pm SD when number of samples >1. Values below

1681 detection limits of the analytical method are denoted n.d. The subscripts "w" and "c"

1682 indicate data given as weight and concentration, respectively.

		Svartatjern		Hyltro	Intensive live feed			
	Abbr.	Copepods	Copepods	Zoopl.	Rotifers	ł	Artemia	
		2000	2001	2001	IMR ^a	1-day ^b	1-day ^c	3-day ^c
Number of samples	Ν	30	26	1	1	1	1	1
FAA in weight (µg/mg DW)	FAA _W	$56.1^{A} \pm 9.7$	64.7 ± 9.8	86.0	16.6	33.7	32.1	27.5
Indispensable amino acids (µg/mg DW)	IAA _w	18.4 ± 3.0	18.2 ± 1.8	19.3	5.8	4.4	5.5	5.1
Indispensable amino acids (%)	IAA _w /FAA _w	$32.9^{A} \pm 2.7$	28.5 ± 3.3	22.4	34.7	12.9	17.0	18.7
Indispensable to dispensable ratio	IAA _w /DAA _w	$0.49^{\text{A}} {\pm}~0.06$	0.40 ± 0.07	0.29	0.53	0.15	0.20	0.23
FAA concentration (nmoles/mg DW)	FAAc	$471.7^{A} \pm 89.8$	580.1±95.1	766.6	124.5	277.6	254.0	219.0
Indispensable amino acids (nmoles/mg DW)	IAA _c	$113.6{\pm}\ 18.2$	$109.0{\pm}~10.7$	119.2	38.0	27.7	36.4	34.3
Indispensable amino acids (%)	IAA _c /FAA _c	$24.3^{\rm A}{\pm}2.0$	19.1 ± 2.2	15.5	30.6	10.0	14.3	15.6
Indispensable to dispensable ratio	IAA _c /DAA _c	$0.32^{\rm A}{\pm}~0.03$	$0.24{\pm}\ 0.03$	0.18	0.44	0.11	0.17	0.19
Indispensable amiono acids (nmoles/mg DW	7)							
Arginine	ARG	$79.6{\pm}15.8$	83.1±13.7	68.3	13.6	13.7	12.3	13.4
Threonine	THR	10.2 ± 2.5	_ d	_ d	3.0	0.9	2.9	3.1
Valine	VAL	$5.8^{A} \pm 1.9$	4.6±1.5	7.5	4.0	1.9	5.0	4.6
Histidine	HIS	$5.1^{A} \pm 1.7$	9.3 ± 5.7	21.3	4.9	2.9	6.2	3.2
Leucine	LEU	$3.6^{A} \pm 1.4$	3.0 ± 1.0	5.0	2.0	1.9	2.9	2.9
Lysine	LYS	3.3±1.1	3.9 ± 1.8	6.7	5.2	4.4	3.1	3.4
Isoleucine	ILE	$2.5^{A} \pm 1.1$	1.9 ± 0.8	3.2	3.0	1.1	1.8	2.0
Phenylalanine	PHE	$2.0^{A}{\pm}\ 0.9$	1.5 ± 0.5	2.6	1.6	0.8	1.1	1.0
Methionine	MET	1.3 ± 0.5	1.4 ± 0.9	4.4	0.6	0.1	0.9	0.5
Tryptophan	TRP	0.3 ± 0.2	0.3 ± 0.1	0.3	0.2	n.d.	0.2	0.1
Dispensable amiono acids (nmoles/mg DW)								
Glycine	GLY	$126.5^{\text{A}}{\pm}37.1$	$231.4^{d} \pm 58.2$	235.3 ^d	8.8	5.4	8.2	9.7
Taurine	TAU	$84.3^{\mathrm{A}}{\pm}~16.7$	101.1 ± 23.3	136.0	2.9	65.5	57.8	58.2
Alanine	ALA	43.5 ± 18.1	36.4 ± 16.0	68.0	8.9	65.7	34.2	28.0
Glutamic acid	GLU	$33.5^{A} \pm 7.1$	24.5 ± 7.0	45.0	14.6	27.0	35.2	31.2

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Proline	PRO	$24.3^{\text{A}} \pm 19.7$	38.3 ± 38.9	125.9	3.9	50.7	34.6	25.0
Aspartic acid+Phosphoserine	ASP+PHS	$17.9^{A} \pm 3.1$	13.7 ± 3.7	9.3	4.6	6.2	6.9	5.3
Glutamine	GLN	10.3 ± 1.7	10.4 ± 3.0	7.6	6.2	11.7	17.0	9.4
Serine	SER	$8.6^{A} \pm 2.3$	7.0 ± 2.0	9.3	16.4	2.1	5.0	6.3
Gamma-amino butyric acid	GABA	3.6±1.5	3.8±1.1	4.4	0.8	1.5	2.9	1.9
Tyrosine	TYR	$3.1^{A} \pm 1.1$	2.5 ± 0.7	3.9	10.6	7.3	5.8	3.5
Asparagine	ASN	$2.7^{A} \pm 0.8$	2.0 ± 0.6	2.9	8.7	6.8	10.0	6.2

^a Institute of Marine Research: rotifers grown on Rotimac and <u>Isochrysis galbana</u> algae.

^b Institute of Marine Research: <u>Artemia</u> enriched with DC-DHA Selco.

1686 ^c Austevoll Marin Yngel AS: <u>Artemia</u> fed DC-DHA Selco and Algamac 2000.

1687 ^d In the 2001 samples, high glycine content caused masking of threonine (next eluated top in the

1688 chromatogram).

1689 ^A Significant difference between copepods from 2000 and 2001.

1690 Table 5

Pigments and vitamins in copepods, zooplankton (copepods and decapod zoeae),
rotifers, and <u>Artemia</u> (1-day or 3-day after hatching). Values are relative to dry weight

- 1693 (DW) and are given as mean \pm SD when number of samples >1. Values below
- 1694 detection limits (n.d.) and trace amounts (tr.) between detection and quantification
- 1695 limits of the analytical method are indicated.
- 1696

		Svartatjern Hy			Intensive live feed				
	Abbr.	Copepods	Copepods	Zoopl.	Rotifers	:	Artemia		
		2000	2001	2001	IMR ^a	1-day ^b	1-day ^c	3-day ^c	
Pigments (µg/g DW)			S						
Number of samples	Ν	30	26	1	1	1	1	1	
Astaxanthin		626.9±139.1	747.7 ± 296.8	197.9	24.0	n.d.	n.d.	n.d.	
Canthaxanthin		n.d.	n.d.	n.d.	n.d.	752.4	744.7	654.0	
β-Carotene		n.d.	n.d.	n.d.	n.d.	n.d.	n.d.	n.d.	
Lipid-soluble vitami (µg/g DW)	ins	$\langle \rangle$							
Number of samples	N	16	19	1	1	1	1	1	
Retinol	Vitamin A	tr.	n.d.	0.2	0.2	_ d	_ ^d	_ ^d	
Cholecalcipherol	Vitamin D ₃	n.d.	n.d.	n.d.	0.9	0.7	1.8	1.0	
Total Tocopherol	Vitamin E _{tot}	$112.0{\pm}\ 28.1$	$114.0{\pm}61.3$	114.0	513.1	571.8	340.2	465.3	
α-Tocopherol	Vitamin E_{α}	$108.0{\pm}\ 28.5$	113.5 ± 61.1	114.0	509.0	562.0	327.8	424.3	
β-Tocopherol	Vitamin E_{β}	0.5 ± 0.2	n.d.	n.d.	n.d.	n.d.	n.d.	n.d.	
γ-Tocopherol	$Vitamin \ E_{\gamma}$	$3.5^{A} \pm 2.3$	0.4 ± 1.4	n.d.	4.1	7.4	9.4	32.9	
δ-Tocopherol	Vitamin E_{δ}	n.d.	n.d.	n.d.	n.d.	2.4	3.0	8.1	
Water-soluble vitami	ins								
$(\mu g/g DW)$									
Number of samples	Ν	16	19	1	1	1	1	1	
Thiamine	Vitamin B ₁	$23.1{\pm}4.7$	22.7 ± 11.7	9.2	48.6	18.2	13.0	20.9	
Riboflavin	Vitamin B ₂	tr.	$28.0{\pm}3.6$	28.9	30.7	53.1	52.1	51.9	
Ascorbic acid	Vitamin C	$476.6{\pm}224.6$	$552.9{\pm}360.2$	271.1	220.1	530.6	361.3	372.6	

^a Institute of Marine Research: rotifers grown on Rotimac and <u>Isochrysis galbana</u> algae.

1698 ^b Institute of Marine Research: <u>Artemia</u> enriched with DC-DHA Selco.

1699 ^c Austevoll Marin Yngel AS: <u>Artemia</u> fed DC-DHA Selco and Algamac 2000.

^d Interactions in the analytical method caused too high retinol readings for <u>Artemia</u>, see section 2.3.4.

1701 ^A Significant difference between copepods from 2000 and 2001.

















1709 Figure 7



1710 Figure 8