"This is the peer reviewed version of the following article: Rørvik, K. A., Dessen, J. E., Åsli, M., Thomassen, M. S., Hoås, K. G., & Mørkøre, T. (2018). Low body fat content prior to declining day length in the autumn significantly increased growth and reduced weight dispersion in farmed Atlantic salmon Salmo salar L. *Aquaculture Research*, 49(5), 1944-1956., which has been published in final form at <u>https://doi.org/10.1111/are.13650</u> This article may be used for non-commercial purposes in accordance with Wiley Terms and Conditions for Use of Self-Archived Versions."

1	Low body fat content prior to declining day length in the autumn significantly increased
2	growth and reduced weight dispersion in farmed Atlantic salmon Salmo salar L.
3	
4	Kjell-Arne Rørvik <sup>1,2</sup> , Jens-Erik. Dessen <sup>1,2*</sup> , Magnus Åsli <sup>1,2†</sup> , Magny S. Thomassen <sup>2</sup> , Kjellrun
5	G. Hoås <sup>1</sup> & Turid Mørkøre <sup>1,2</sup>
6	
7	<sup>1</sup> Nofima, NO-1432 Ås, Norway
8	<sup>2</sup> Department of Animal and Aquaculture Sciences, Norwegian University of Life Sciences,
9	NO-1432 Ås, Norway
10	
11	*Corresponding author: Jens-Erik Dessen; Nofima, 1432 Ås, Norway; Tel: +47 979 52 768;
12	Email: jens-erik.dessen@nofima.no
13	†Current address: Cermaq Group As, NO-0102 Oslo, Norway
14	
15	
16	
17	
17	
18	
19	
20	
20	
21	
22	Running head: Body fat and growth in Atlantic salmon
23	Key words: Salmon, growth response, body lipids, seasonal cues

Based on the regulatory effects of body fat on appetite and seasonal variations in fat 26 27 deposition and growth of Atlantic salmon, the present study tested the hypothesis that body fat content prior to declining day length in the autumn can significantly modulate growth rate. 28 The growth rate of salmon (mean initial body weight, BW=2.3 kg) with different muscle fat 29 content prior to autumn, subjected to natural photoperiod and temperature, during a 7-months 30 period (mean final BW=6.6 kg) was studied. In August, three fish groups (HF, LF and 0.5LF 31 group) with significantly different muscle fat content (HF=16.4%, LF=13.2% and 32 0.5LF=11.3%), individually marked with PIT-tag, were mixed into the four net pens and fed a 33 standard high-energy diet until March the following year. The muscle fat content prior to the 34 autumn had a highly significant (P < 0.0001) effect on growth during the seven month main-35 36 dietary period, even after identical fat stores among the groups were restored, indicating a more complex explanation than just a lipostatic regulation mechanism. Mean thermal growth 37 38 coefficients were HF=2.9, LF=3.4 and 0.5 LF=3.9, resulting in increased final weight gain for LF and 0.5LF of 590 g. and 980 g., respectively, compared to the HF group. The LF groups 39 obtained a significantly higher homogeneity in BW and shape than HF fed fish in March, 40 optimizing automatic gutting and filleting at slaughter. The improved growth response among 41 the LF groups by reducing lipid levels can potentially be utilized in closed and semi-closed 42 production units where photoperiod can be manipulated. 43

- 44
- 45
- 46
- 47
- 48

### 49 **INTRODUCTION:**

50

Fish that encounter setbacks induced by nutritional deficit, feed deprivation or sub-optimal 51 conditions often display increased feed consumption (hyperphagia) and compensatory growth 52 (CG) when circumstances are normalized (Ali, Nicieza, & Wootton, 2003; Foss & Imsland, 53 2002; Metcalfe & Monaghan, 2001). The degree of CG in fish vary and is often categorized 54 based on the growth catch-up ability (Ali et al., 2003). Feed restriction or deprivation induce 55 changes in body energy by depleting lipid stores, and during the course of CG and 56 hyperphagia, body weight and lipid reserves are gradually restored (Ali et al., 2003; Bull & 57 58 Metcalfe, 1997; Jobling & Miglavs, 1993; Metcalfe & Thorpe, 1992). The lipostatic model is often discussed within the circumstances of CG responses in fish (Jobling & Johansen, 1999; 59 Johansen, Ekli, Stangnes, & Jobling, 2001). The lipostatic regulation hypothesis identifies 60 61 adipose tissue and stored lipids to have an important role in governing appetite (Jobling & Johansen, 1999; Keesey & Corbett, 1984; Kennedy, 1953). The model implies that the 62 amounts of stored fat has a negative feedback control on feed intake and is important for the 63 regulation of energy homeostasis. Hence, CG is not only a response to recover body weight, 64 but also a strong response to restore lipid levels and thereof CG will cease once this is 65 achieved (Ali et al., 2003; Jobling & Johansen, 1999; Johansen, Ekli, & Jobling, 2002). 66 Johansen et al., (2002) showed that altering body lipids of juvenile salmon by dietary 67 administration of low-fat feeds yield similar growth responses as deprivation or feed 68 restriction per se. 69

70

In modern high-fat diets for salmonids, lipids of marine and vegetable origin are the main
sources of energy and support growth efficiently if essential fatty acids requirements are met
(Bell et al., 2001; Thomassen & Røsjø, 1989; Torstensen, Lie, & Frøyland, 2000). Because

salmonids have a high ability to utilize large amount of lipids efficiently for growth, high-fat 74 diets with up to 380 g kg<sup>-1</sup> of fat are commonly used in intensive salmon farming (Torrissen et 75 al., 2011). However, salmonids also have the capacity to store large amounts of excess fat as 76 77 triacylglycerols mainly in the muscle and visceral cavity (Aursand, Bleivik, Rainuzzo, Leif, & Mohr, 1994). Body lipid content of farmed salmonids correlates with fish size, dietary fat 78 level and feed intake (Aksnes, 1995; Hemre & Sandnes, 1999; Torstensen, Lie, & Hamre, 79 2001). Like other anadromous species, Atlantic salmon display seasonal changes in feed 80 intake, growth and lipid deposition during the seawater phase (Mørkøre & Rørvik, 2001). 81 Farmed Atlantic salmon display elevated deposition of lipids in muscle and increased 82 retention of lipids in whole body during declining day length in autumn, with a concomitant 83 increase in feed intake, somatic growth and condition factor (CF) (Alne, Oehme, Thomassen, 84 Terjesen, & Rørvik, 2011; Dessen, Weihe, Hatlen, Thomassen, & Rørvik, 2017; Mørkøre & 85 86 Rørvik, 2001; Rørvik et al., 2010). This is particularly pronounced for salmon reared at high latitudes that experience long winters and late spring, which results in reduced lipid levels and 87 CF prior to summer and autumn. 88

89

The recent increase in automation of fish processing at slaughter requires uniform body 90 weight (BW) and shape among the salmon for optimal efficiency and quality of products such 91 as gutted fish and fillets. Increased uniformity of BW and CF reduces the need for manual 92 gutting/filleting of very small or large individuals. Due to this, the homogeneity in body shape 93 and mass of salmonids are important parameters in salmon farming industry and low 94 dispersion in BW and CF are beneficial at time of harvest. The homogeneity of BW may be 95 strongly influenced by events occurring during the production cycle, i.e. disease outbreaks, 96 handling stress, reduced seawater tolerance or competition of feed (McLoughlin, Nelson, 97 McCormick, Rowley, & Bryson, 2002; Ryer & Olla, 1996; Taksdal et al., 2007; Usher, 98

99 Talbot, & Eddy, 1991). The dispersion in the distribution of BW, length and CF are often 100 assessed by calculating the coefficient of variation (CV). The CV of BW for farmed salmon 101 grown from 70 until 300 g. and from 60 until 500 g. fed either in excess or restrictively for 102 period followed by unrestricted feeding, are reported to vary from 9 to 13% and 16 to 21%, 103 respectively (Johansen et al., 2001). In the latter study, no significant differences was 104 observed in the CV of BW between fish fed in excess and fish fed restrictively.

105

106 The majority of studies regarding growth responses related to lipid content are based on inhouse laboratory experiments with small juvenile salmonids under constant conditions. To our 107 knowledge, few have investigated grow out salmon with different lipid content subjected to 108 seasonal environmental changes in photoperiod and temperature. Due to the regulatory effects 109 of body fat on appetite and the observed fat storage in salmon linked to the seasonal cues, the 110 present study tested the hypothesis that lipid status prior to declining day length in the autumn 111 functions as a significant growth regulator. Accordingly, the growth rate for three groups of 112 salmon with different muscle fat content prior to autumn, subjected to natural photoperiod and 113 114 temperature, was studied throughout a 7-months period. About each second month, weight samplings and analysis of muscle fat content was conducted to investigate any relationship 115 between fat accumulation and periodic growth rate, and to identify the duration of a potential 116 lipostatic regulatory effect. Changes in visceral fat, CF, length, and the dispersion in BW and 117 CF were further assessed. 118

119

## 120 MATERIAL & METHODS:

121

122 This experiment was conducted in accordance with laws and regulations that control123 experiments and procedures in live animals in Norway, as overseen by the Norwegian Animal

Research Authority. Stunning and sampling of fish were performed in accordance with the Norwegian Animal Welfare act. Fish were treated as production fish up to the point of tissue sampling which was only conducted after the fish were put to death.

127

The experiment was conducted in seawater on the Norwegian west coast (Ekkilsøy, Norway 128 3° 03' N, 7° 35' E) at Nofima research center from August 2011 to March 2012. In July 2010, 129 the fish were transferred to seawater as S1 smolt, at which time the BW was 62 g. From the 130 10 to 12 of May 2011, the post-smolt were re-stocked into three net-pens (343 m<sup>3</sup>) with 650 131 fish per pen. Prior to this, all individual fish were measured for weight and length, and tagged 132 using passive integrated transmitter tags (PIT-tags) placed in the body cavity just posterior to 133 the gut. The average BW per pen was 1085 g. (SD = 79 g.) and each pen received different 134 dietary treatments: a high-fat diet (HF), a low-fat high-protein diet (LF) or half the ration of 135 136 the low-fat high-protein diet (0.5LF). The 0.5LF-group were given half the amount of the feed provided to fish administrated the LF-diet the day before. Skretting (Averøy, Norway) 137 produced the feeds and the composition of the HF diet was (wet weight, as is basis): dry 138 139 matter 93.4%, crude protein 33.5%, crude lipid 34.1%, nitrogen-free extract (NFE) 21.2%, ash 4.6% and gross energy of 25.1 MJ kg<sup>-1</sup>. The composition of the LF diet was (wet weight, 140 as is basis): dry matter 91.7%, crude protein 49.9%, crude lipid 17.5%, NFE 17.1%, ash 7.2% 141 and gross energy 21.7 MJ kg<sup>-1</sup>. The three dietary treatments were fed from 12 of May until 9 142 of August (pre-dietary phase). May 12th, the fish were sampled for analysis of initial muscle 143 fat content and biometric data. The analysis showed the following (mean  $\pm$  SE, n = 30): BW: 144  $1087 \pm 97g$ , initial muscle fat:  $12.2 \pm 1.1\%$  and initial CF:  $1.10 \pm 0.06$ . After ending the pre-145 dietary phase, the PIT-tag, BW and length of all individual fish in the three pens were 146 recorded. In addition, fish from each pen were sampled for analysis of muscle and visceral fat 147 content. The pre-dietary feeding phase generated three fish groups with significantly different 148

(P < 0.05) muscle fat content, visceral fat and visceral mass (Table 1). During the pre-dietary</li>
phase, 2.5%, 0.6% and 0.3% fish died in the HF, LF and 0.5LF group, respectively. The
majority of mortality occurred from May until mid-June and was not related to any disease
outbreak (non-specific morality).

153

154 (**Table 1**).

155

At the 10 to 11 of August, the fish were restocked from the three original pens used in pre-156 dietary phase into four new pens (125 m<sup>3</sup>). Each of the four pens contained 50 fish from each 157 of the three pre-dietary treatments (HF, LF and 0.5LF), 150 fish in total (Fig 1). During the 158 period from 11 of August until termination at 20 of March 2012 (main-dietary phase), the 159 pens were fed isonitrogenous and isoenergetic diets produced by Ewos (Bergneset, Norway) 160 161 (Table 2). The current experiment was an integrated part of a large study were potential effects of dietary oil source were investigated. Therefore, two pens in the main-dietary phase 162 were fed a diet with a marine oil profile (MO), whereas the two other pens were fed a diet 163 with a rapeseed oil profile (RO). The marine oil diet (MO) had an inclusion of 70 % South 164 American fish oil and 30 % of rapeseed oil. The rapeseed oil diet (RO) had an inclusion of 70 165 % rapeseed oil and 30 % South American fish oil. During the main-dietary phase, the pellet 166 size was changed from 7 to 9 mm in December due to the increase in BW of the fish. 167

168

169 (Fig. 1 and Table 2)

170

In both periods, feed was administrated using automatic feeders (Betten Maskinstasjon AS,
Vågland, Norway) and uneaten feed was collected as described in Einen, Mørkøre, Røra &
Thomassen (1999) and corrected for the recovery of dry matter as described by Helland,

Grisdale-Helland & Nerland (1996). The fish groups (except the 0.5LF group during the pre-174 dietary phase) were fed to satiation and the feed ration was set at 5-10 % in excess (ad libitum 175 feeding). The fish were fed four times a day until October 2011, after this, the fish were fed 176 177 three times a day until termination in March 2012. Adjustments of the feed ration was done according to the daily amount of uneaten feed collected. Due to the stocking of 50 fish from 178 each of the pre-dietary treatments into each net pen, it was not possible to determine the feed 179 intake or feed utilization of the different pre-dietary groups during the main-dietary phase. 180 The pens were checked for mortalities daily and the dead fish were collected and weighed. 181 The fish were exposed to natural variations in photoperiod and sea temperature during the 182 experiment (Fig. 2). 183

184

185 (Fig. 2)

186

Three samplings were performed during the main-dietary phase; from 9 to 11 October 2011, 187 from 6 to 9 December 2011 and the final sampling and termination of the experiment was 188 conducted from 20 to 22 March 2012. At each sampling, all fish were anaesthetized (MS-222 189 metacaine 0.1 g L<sup>-1</sup>, Alpharma, Animal Health, Hampshire, UK) and the PIT-tag, fork length 190 and weight of each individual fish were recorded. All fish were starved two days prior to the 191 samplings in August and October, and three days prior to the samplings in December and 192 March to avoid feed matter in the gastrointestinal system. At each sampling, 10 fish from each 193 pre-dietary group in all the pens were sampled. The sampled fish at each sampling point were 194 selected so that the mean weight corresponded to the mean weight of all the fish in the 195 respective fish group within each pen (as all possible fish were weighted and PIT-tag read). 196 197 After anesthetization, a blow to the head was used to kill fish sampled for analysis. Then the gill arches were cut and the fish were bled out in ice seawater. Length and weight of each 198

individual fish sampled for analysis were recorded after bleeding and the fish visually tagged. 199 The fish were then gutted and filleted by hand during the pre-rigor state. Norwegian Quality 200 Cut, NQC (NS9401, 1994) from the left fillet was photographed and the fat content was 201 202 predicted by digital image analyses (PhotoFish, AKVAgroup, Bryne, Norway), as described by Folkestad et al. (2008). The visceral mass of the sampled fish were pooled on group level, 203 homogenised and frozen at - 20°C for later analyses of total lipid content as described by 204 205 Folch, Less & Stanley (1957). The proximate composition of crude protein, lipid (acidichydrolysis method), starch and moisture of the diets were analysed according to the methods 206 207 described by Oehme et al. (2010). To determine the fatty acid (FA) composition of the diets, lipids were first extracted according to Foch et al. (1957), and a sample of 2 ml from the 208 chloroform-methanol phase was dried under N2 gas, then the residual lipid extract was trans-209 210 methylated overnight with 2',2'-dimethoxypropane, methanolic HCl and benzene at room temperature according to Mason & Waller (1964). Finally, the methyl esters were separated 211 by gas chromatography and individual FA were identified as described in Røsjø et al. (1994). 212

213

The growth rate of the fish are presented as the thermal growth coefficient (TGC), and were calculated as described by Iwama & Tautz (1981): TGC =  $[(M_1^{1/3} - M_0^{1/3}) \times (\Sigma T)^{-1}] \times 1000$ , where  $M_0$  and  $M_1$  are the initial and final BW, respectively, and  $\Sigma T$  is the sum of day degrees during the period (feeding days x average temperature, °C). The mean TGC for the total main-dietary phase was calculated as the weighted arithmetic mean of the periodical TGC to balance these values in relation to their relative contribution to the weight gain.

220

All fish sampled and killed for analysis were starved and bled. The calculation of visceralsomatic index is therefore based on BW with minimal blood content and no feed material in the gastrointestinal system. Visceral-somatic index (VSI), was calculated as:  $Y(g) \ge BW(g)^{-1}$  x 100, where *Y* is the measured visceral mass. The visceral mass was defined as all mass in the abdominal cavity except liver, heart, kidney and swim bladder. The CF was calculated as:  $100 \times BW$  (g) x fork length (cm) <sup>-3</sup>. The dispersion in the distribution of BW, length and CF were assessed by calculating the CV: (standard deviation x mean value<sup>-1</sup>) x 100.

228

The results were analysed by the General Linear Model (GLM) procedure in the SAS 9.4 229 computer software (SAS Institute Inc., Cary, NC, USA). Mean results per fish group in each 230 pen were initially subjected to a two-way analysis of variance (ANOVA) to evaluate the 231 effects of muscle fat content due to the pre-dietary phase (0.5LF, LF and HF), main-dietary 232 treatment (oil source; MO and RO-diet) and their interaction (pre-diet x main-diet). As the 233 statistical analysis showed that neither oil source nor the interaction term has significant 234 effects on the traits studied, the data was analysed using pre-dietary treatment as the only 235 236 experimental factor (one-way ANOVA). Significant differences among experimental groups within treatments were indicated by Duncan's multiple range test. Least-square means 237 (lsmeans) comparison were also used to identify differences among variables within 238 treatments. The Pearson product-moment correlation coefficient was used to describe the 239 association between two variables. Linear regression analysis were conducted using Microsoft 240 excel. The proportion of total variance explained by the model was expressed by  $R^2$  and the 241 level of significance was chosen at P < 0.05. Tendencies was identified at P = 0.05 - 0.1. The 242 results are presented as means  $\pm$  SEM, if not otherwise stated. 243

244

No significant effects of the main-dietary treatment (RO and MO-diet) or interaction term (main x pre-diet) *per se* were detected on the traits examined during the main-dietary phase. Thus, only the effects of body fat content due to the pre-dietary treatment are presented in the results. No significant differences in mortality among the pre-dietary groups were observed
during the main-dietary phase (24 out of 650 fish, 3.6%).

250

### 251 **RESULTS:**

252

The muscle fat content increased by 8.1% for 0.5LF fish, 5.6% for the LF group and 3.6% for 253 HF group from August to October (Fig 4A1). Thus, during an 8-week period of declining day 254 length, the initial significant differences in muscle fat content was equilibrated. TGC was 255 highest for the 0.5LF group, intermediate for the LF group and lowest for the HF group (Fig 256 5A). The growth rate and the increase in muscle fat content from August to October showed a 257 significant positive linear relationship, and the increase in muscle fat explained 81% of the 258 variation in growth (Fig 3). From August to October, the growth rates were therefore highly 259 affected by the pre-dietary treatment (ANOVA:  $R^2 = 0.97$ , P < 0.001). The muscle fat did not 260 differ significantly between the pre-dietary treatments in October or December (Fig. 4A1), but 261 pre-diet still significantly influenced the growth rates (ANOVA: P < 0.05,  $R^2 = 0.51$ ) and the 262 TGCs were similar, relatively, to the period from August to October (0.5LF > LF > HF), 263 although no significant differences was found between LF and HF group. In the period 264 December to March, the TGC for the 0.5LF and LF group were significantly higher (P <265 0.05) than the HF group (Fig 5A). At the end of the main-dietary phase, the muscle fat content 266 of the LF group was significantly lower (P < 0.05) than the 0.5LF group, and tended to be 267 lower (P < 0.1) than the HF group (Fig 4A2). 268

269

270 (Fig. 3 and 4)

The BW of the LF group reached a similar BW as the HF fish in October, whereas the 0.5LF 272 group reached a similar BW as the HF group in December (Fig 4B1). At the end of the trial in 273 March, the LF group (6.87  $\pm$  0.07 kg.) had a significantly higher (P < 0.05) BW than the HF 274 group (6.40  $\pm$  0.16 kg.) (Fig 4B2). The 0.5LF group (6.62  $\pm$  0.12 kg.) had numerical higher 275 BW than the HF group, however, no statistically significant difference was detected. From 276 August 2011 to March 2012, the 0.5LF group gained 980 g. and the LF group gained 590 g. 277 more relative to the BW of the HF group (Fig 5B). The overall weighted mean TGC during 278 the main-dietary phase were 3.9 for the 0.5LF group, 3.4 for the LF group and 2.9 for the HF 279 group. Hence, the pre-dietary treatment and consequently the fat status in August 2011 had a 280 clear and significant effect on growth, weight gain and the changes in BW throughout the 281

whole main-dietary phase, with a total duration of seven months.

283

```
284 (Fig. 5 and 6)
```

285

No significant differences in length between LF and HF group were detected during the trial 286 (Fig 6B1). The strong growth spurt of the 0.5 LF group resulted in no significant differences 287 in length between the 0.5 LF (75.9  $\pm$  0.2 cm.) and HF group (76.4  $\pm$  0.8 cm.) at the trial 288 termination in March. However, the LF (77.9  $\pm$  0.1 cm.) group was significantly longer (P < 289 290 0.05) than the 0.5LF group (Fig 6B2). The 0.5LF group that had the lowest CF in August, ended up having the significantly highest CF at termination (Fig. 6A1 and A2). The overall 291 development in CF correlated well with the changes in muscle fat during the study (r = 0.98, 292 P < 0.01). Significant positive overall correlations were also observed between the final CF 293 and mean TGC (r = 0.88; P < 0.001), and between the final CF and total weight gain (r = 294 295 0.88; *P* < 0.001).

The visceral fat content of the HF group was consistently highest, although only significant in October (Fig 7). The VSI of the LF group ( $8.5 \pm 0.1$ ) was significantly lower (ANOVA: *P* < 0.01) than the HF group ( $9.0 \pm 0.1$ ) in October, whereas the VSI of the 0.5LF group ( $8.7 \pm$ 0.1) was not different from the LF or HF group. No significant differences in VSI were detected in December (overall mean; VSI:  $8.8 \pm 0.1$ ) or March (overall mean; VSI:  $9.8 \pm 0.2$ ).

302

303 (Fig. 7)

304

The 0.5LF group had the highest CV<sub>BW</sub> at the end of the pre-dietary phase (Fig 8A). From 305 August to October, the CV<sub>BW</sub> of the 0.5LF group decreased and no significant difference in 306 CV<sub>BW</sub> was observed at the samplings in October and December. However, at termination in 307 March, the HF group had a significantly (P < 0.05) higher CV<sub>BW</sub> compared to both LF and 308 309 0.5LF group. The CV<sub>CF</sub> was lowest for the LF group and similar for the HF and 0.5LF group at the end of the pre-dietary phase (Fig 8B). At the sampling in October, after the large 310 311 increase in fat deposition, growth and CF, the 0.5LF group had the highest CV<sub>CF</sub>. The 312 variation within the CV of CF for the 0.5LF group was at this time very high and no significant differences between the groups was detected. The CV<sub>CF</sub> for the HF group 313 increased gradually from October to March. In line with the CV<sub>BW</sub>, the HF group had a 314 significantly (P < 0.05) higher CV<sub>CF</sub> compared to the 0.5LF and LF group at termination. No 315 significant differences in the CV<sub>LENGHT</sub> was detected during the experiment (results not 316 shown). 317

318

319 (Fig. 8)

320

The coinciding increase in fat and improved growth shown by the 0.5LF and LF group 324 325 compared to the HF group in the beginning of main-dietary phase (August and September), seem to reflect a growth response similar to CG and lipostatic regulation observed in previous 326 studies in the field and laboratory (Ali et al., 2003; Jobling & Johansen, 1999; Johansen et al., 327 328 2002, 2001). The obtained growth rates, fat increase and weight gain from August to October, together with the high feed intake (on pen basis), indicate that the 0.5LF and LF group had 329 increased feed consumption and hyperphagic behaviour. In addition to the high growth rate of 330 331 the 0.5LF and LF groups, the increase in muscle and visceral fat content during August and September were substantial for these two groups. However, the muscle fat of the HF group 332 also increased during this period (16.4%  $\rightarrow$  20.0%). The TGC of the HF group had an average 333 334 of 3.0, which is regarded as a normal and sufficient growth rate (Austreng, Storebakken, & Åsgård, 1987; Thorarensen & Farrell, 2011). Thus, improved growth in the LF groups from 335 August to October, compared to the HF group, is not a result of impaired growth due to 336 337 adiposity in the latter group, but rather a stronger response among the fish in the LF and 0.5LF group. The growth responses from August to October differ from the observations of 338 339 Johansen, Sveier, & Jobling (2003), where Atlantic salmon fed a high fat diet during both the build-up and main phase, maintained their body fat levels after the build-up phase, at the same 340 time as feed intake was down-regulated and growth impaired. In the present study, the salmon 341 were exposed to natural photoperiod, as opposed to the study by Johansen et al. (2003), where 342 the salmon were held under continuous light (24L:0D). It has been suggested that reduction in 343 day length is an important environmental factor that trigger the salmon to assess its current 344 mass during this time of the year (Maclean & Metcalfe, 2001). It may also apply for energy 345 status and body condition (Kadri, Mitchell, Metcalfe, Huntingford, & Thorpe, 1996). In 346

addition, high retention of dietary lipid, elevated fat deposition, increased CF and rapid
growth are observed during the autumn period (Alne et al., 2011; Dessen et al., 2017; Kadri et
al., 1996; Mørkøre & Rørvik, 2001). Hence, the influence of natural seasonal cues might be
the main reason for the observed differences in growth between the present study and the one
of Johansen et al. (2003).

352

In October, two months after the start of the main-dietary phase, muscle fat and CF were 353 restored in both the LF and 0.5LF group compared to the HF group. This observation shows 354 that Atlantic salmon is able to rapidly replenish lipid stores and body condition during the 355 autumn following a feeding period of a low-fat diet or restricted ration of this diet. In contrast, 356 the visceral fat content among the groups maintained about the same pattern thought out the 357 study. The level or severity of restricting lipid deposition during pre-dietary phase was highly 358 359 negatively related with the magnitude of the subsequent growth response from August to October. This was particularly linked to the relative muscle fat content at termination of the 360 pre-dietary phase prior to autumn. The degree of CG response seem also related to the level of 361 deviance in body condition, length and mass in the restricted or deprived fish groups 362 compared to their non-treated counter-specifics (Alvarez & Nicieza, 2005; Johansen et al., 363 2001; Johnsson & Bohlin, 2005; Johnsson & Bohlin, 2006). Although the deviance in mass 364 and length may have contributed to the growth response in the present study, the small 365 difference between the LF and HF groups in August and the strong correlation between 366 muscle fat and growth, indicate that fat/energy status seem to be the most important growth 367 regulator during August and September. The increased growth and rapidly replenishment of 368 lipid stores suggest a robust mechanism for the regulation of body fat, and are in line with the 369 observation of Silverstein, Shearer, Dickhoff & Plisetskaya (1999). 370

Several studies have indicated that animals displaying CG prioritise the restoration of body 372 373 condition and fat stores before more resources are allocated to support structural and skeletal growth (Broekhuizen, Gurney, Jones, & Bryant, 1994; Johnsson & Bohlin, 2006). In part, the 374 375 results of the present study support these observations, as both the relative muscle fat content and CF were quickly restored in the 0.5LF group, but not that quickly restored for BW and 376 length. Some studies have also suggested that structural restoration can be delayed due to the 377 effects of food deprivation or restriction on the endocrine system, involved in the regulation 378 of growth (Björnsson, 1997; Johnsson, Jönsson, & Björnsson, 1996). There is evidence that 379 skeletal and muscle growth are independent processes and that the relationship between 380 length and weight is approximately cubic (Einen, Waagan, & Thomassen, 1998; Jobling, 381 2002; Mørkøre & Rørvik, 2001). Thus, changes in weight are relatively greater than in length, 382 and the rapid increase in BW and fat content observed among the 0.5LF group in the autumn, 383 384 may be a factor explaining why length are restored later than body shape and fat content.

385

The stabilisation of the muscle fat in October coincides with the study of Mørkøre & Rørvik 386 (2001). This may suggest that the capacity of muscle fat deposition has reached an upper limit 387 at this time point. There is documentation that CG responses will cease as lipid stores and 388 body condition are restored to similar levels as the non-affected conspecifics (Johansen et al., 389 2001; Ali et al., 2003; Alvarez & Nicieza, 2005; Johnsson & Bohlin, 2005). In the present 390 study, the LF and 0.5LF groups continued to grow faster than the HF group both during the 391 periods October to December and December to March. The improved growth of the LF 392 groups from December to March was evident although the relative muscle fat content, CF and 393 BW were restored prior to this period and not significantly different from the HF group. 394 395 Hence, the observed growth response in this period is not directly related to restoration of fat or BW. The sexual maturation process in Atlantic salmon requires, in addition to photoperiod, 396

sufficient fat and energy reserves (Kadri et al., 1996; Rowe & Thorpe, 1990; Taranger et al., 397 2010). The production of gonads are energetically expensive and acquire high-energy 398 investment (Fleming, 1996; Jonsson, Jonsson, & Hansen, 1997). Appropriate and available 399 400 energy and fat reserves during the spring period seem to be a major factor controlling initiation and proceeding of the maturation process (Thorpe, 1994; Thorpe, Mangel, Metcalfe, 401 & Huntingford, 1998; Wright, 2007). Too low energy and fat levels may arrest the maturation 402 process and postpone reproduction (Duston & Saunders, 1999; Rowe & Thorpe, 1990; Rowe, 403 404 Thorpe, & Shanks, 1991; Thorpe, 1994; Thorpe, Talbot, Miles, & Keay, 1990). Hence, well growing salmon with a high and stable fat content are more likely to adopt the development 405 pathway of becoming sexual mature (Thorpe, 1994). Following this line of arguments, the 406 stronger growth response observed in both LF groups compared to the HF group prior to the 407 spring period in the present study, may have been triggered by the salmon reproductive life 408 409 strategy. However, to verify this, the groups of salmon must be studied for a longer period (during late spring, summer and autumn) and measurements of relevant plasma hormones, 410 411 gonad-somatic index and gene expression of e.g. myosin should be conducted. Unfortunately, 412 this was not possible in the present study. Anyhow, observation of a long-term improved growth response is important for a further development of a dynamic seasonal feeding 413 concept in salmon farming. Not only for traditional sea cage farming, but also in closed and 414 semi-closed production units where photoperiod may be manipulated. Taken into 415 consideration that the initial BW of the 0.5LF group was 738 g. less than the HF group, a 416 relative increase in weight gain of 950 g. more than the HF group is impressive. 417

418

When feed availability is restricted, competition for the feed often increase and dominant
individuals may try to monopolize the feeding area to obtain larger amounts of feed that is
supplied (Maclean & Metcalfe, 2001; Ryer & Olla, 1996). High competition for feed may

therefore lead to increased disparities in feed intake and growth that consequently will give 422 higher variation in BW. To minimize such effects, the 0.5LF group was administrated all 423 daily feed in only one ration during the pre-dietary phase. The high dispersion in BW and CF 424 among the HF group at termination of the main-dietary phase indicates that the 0.5LF and LF 425 group had an increase in weight and CF that was more homogeneous than the HF group. This 426 was probably due to the increased growth of LF groups in latter stages of the trial. The 427 possibility that fish among the LF groups displayed aggressive behaviour and tried to 428 429 monopolize food in this period seem unlikely due to three main factors: i) the HF group showed a normal and satisfying growth with mean TGC of 3.2, ii) feed was administered in 430 excess during the main-dietary phase to ensure ad libitum feeding and iii) no or little fin 431 damage were observed at termination. 432

433

In summary, salmon with low body fat levels (LF groups) prior to declining day lengths in the 434 autumn displayed significantly higher growth rate and weight gain compared to the control 435 436 fish (HF group). The initial differences in muscle fat and CF were restored after only two 437 months, displaying rapid replenishment of lipid stores and body condition. Differences in body fat content prior to autumn had significant effect on growth throughout the whole seven-438 month main-dietary phase, even after similar body fat stores among the groups were obtained, 439 indicating a more complex explanation than just a lipostatic regulation mechanism. The LF 440 and 0.5LF fed fish obtained a significantly lower variation in BW and CF than the HF fed fish 441 at trial termination. This increased uniformity of BW and CF may reduce the amount of 442 manual gutting and filleting of large and small individuals, which optimizes the efficiency of 443 automatic gutting and filleting of salmon at the time of slaughter. 444

445

# 446 ACKNOWLEDGMENTS:

448	The authors appreciate the excellent technical assistance and rearing of fish that was provided
449	by the staff from the former Nofima research station at Averøy (now Marine Harvest research
450	station), with special thanks to Sissel Nergaard. This work was supported by a project grant
451	from the Norwegian Seafood Research Fund (FHF, grant number 900653). The authors
452	declare that there are no potential sources of conflict of interest with this work.
453	
454	<b>REFERENCES:</b>
455	
456	Aksnes, A. (1995). Growth, feed efficiency and slaughter quality of salmon, Salmo salar L.,
457	given feeds with different ratios of carbohydrate and protein. Aquaculture Nutrition, 1,
458	241-248. https://doi.org/10.1111/j.1365-2095.1995.tb00050.x
459	Ali, M., Nicieza, A., & Wootton, R. J. (2003). Compensatory growth in fishes: A response to
460	growth depression. Fish and Fisheries, 4, 147–190. https://doi.org/10.1046/j.1467-
461	2979.2003.00120.x
462	Alne, H., Oehme, M., Thomassen, M., Terjesen, B., & Rørvik, K. A. (2011). Reduced growth,
463	condition factor and body energy levels in Atlantic salmon Salmo salar L. during their
464	first spring in the sea. Aquaculture Research, 42, 248–259.
465	https://doi.org/10.1111/j.1365-2109.2010.02618.x
466	Alvarez, D., & Nicieza, A. G. (2005). Compensatory response "defends" energy levels but not
467	growth trajectories in brown trout, Salmo trutta L. Proceedings of the Royal Society B:
468	Biological Sciences, 272, 601–7. https://doi.org/10.1098/rspb.2004.2991
469	Aursand, M., Bleivik, B., Rainuzzo, J. R., Leif, J., & Mohr, V. (1994). Lipid distribution and
470	composition of commercially farmed Atlantic salmon (salmo salar). Journal of the
471	Science of Food and Agriculture, 64, 239–248. https://doi.org/10.1002/jsfa.2740640214

- 472 Austreng, E., Storebakken, T., & Åsgård, T. (1987). Growth rate estimates for cultured
- 473 Atlantic salmon and rainbow trout. *Aquaculture*, **60**, 157–160.
- 474 https://doi.org/10.1016/0044-8486(87)90307-3
- Bell, J. G., McEvoy, J., Tocher, D. R., McGhee, F., Campbell, P. J., & Sargent, J. R. (2001).
- 476 Replacement of fish oil with rapeseed oil in diets of Atlantic salmon (*Salmo salar*)
- 477 affects tissue lipid compositions and hepatocyte fatty acid metabolism. *The Journal of*
- 478 *Nutrition*, **131**, 1535–43. Retrieved from
- 479 http://www.ncbi.nlm.nih.gov/pubmed/11340112
- 480 Björnsson, B. T. (1997). The biology of salmon growth hormone: from daylight to
- dominance. *Fish Physiology and Biochemistry*, **17**, 9–24.
- 482 https://doi.org/10.1023/A:1007712413908
- 483 Broekhuizen, N., Gurney, W. S. C., Jones, A, & Bryant, A.D. (1994). Modeling
- 484 Compensatory Growth. *Functional Ecology*, **8**, 770–782. Retrieved from
- 485 isi:A1994PW40300013
- 486 Bull, C. D., & Metcalfe, N. B. (1997). Regulation of hyperphagia in response to varying
- 487 energy deficits in overwintering juvenile Atlantic salmon. *Journal of Fish Biology*, **50**,

488 498–510. https://doi.org/10.1111/j.1095-8649.1997.tb01945.x

- 489 Dessen, J. E., Weihe, R., Hatlen, B., Thomassen, M. S., & Rørvik, K. A. (2017). Different
- 490 growth performance, lipid deposition, and nutrient utilization in in-season (S1) Atlantic
- 491 salmon post-smolt fed isoenergetic diets differing in protein-to-lipid ratio. *Aquaculture*,
- **473**, 345–354. https://doi.org/10.1016/j.aquaculture.2017.02.006
- 493 Duston, J., & Saunders, R. L. (1999). Effect of winter food deprivation on growth and sexual
- 494 maturity of Atlantic salmon (Salmo salar) in sea water. Canadian Journal of Fisheries
- 495 *and Aquatic Sciences*, **56**, 201–207. https://doi.org/10.1139/f98-165
- Einen, O., Mørkøre, T., Røra, A. M. B., & Thomassen, M. S. (1999). Feed ration prior to

- 498 *salar*). *Aquaculture*, **178**(1–2), 149–169. https://doi.org/10.1016/S0044-8486(99)00126-
- 499

Х

- 500 Einen, O., Waagan, B., & Thomassen, M. S. (1998). Starvation prior to slaughter in Atlantic
- salmon (*Salmo salar*): I. Effects on weight loss, body shape, slaughter- and fillet-yield,
- 502 proximate and fatty acid composition. *Aquaculture*, **166**, 85–104.
- 503 https://doi.org/10.1016/S0044-8486(98)00279-8
- 504 Fleming, I. a. (1996). Reproductive strategies of Atlantic salmon: ecology and evolution.
- 505 *Fisheries, Reviews in Fish Biology and Fisheries*, **6**, 349–416.
- 506 https://doi.org/10.1007/BF00164323
- 507 Folch, J., Lees, M., & Stanley, G. H. S. (1957). A simple method for the isolation and
- 508 purification of total lipides from animal tissues. *Journal of Biological Chemistry*, **226**,
- 509 497–509. https://doi.org/10.1007/s10858-011-9570-9
- 510 Folkestad, A., Wold, J. P., Rørvik, K. A., Tschudi, J., Haugholt, K. H., Kolstad, K., &
- 511 Mørkøre, T. (2008). Rapid and non-invasive measurements of fat and pigment
- 512 concentrations in live and slaughtered Atlantic salmon (*Salmo salar* L.). *Aquaculture*,
- **280**, 129–135. https://doi.org/10.1016/j.aquaculture.2008.04.037
- 514 Foss, A., & Imsland, A. K. (2002). Compensatory growth in the spotted wolffish Anarhichas
- 515 *minor* (Olafsen) after a period of limited oxygen supply. *Aquaculture Research*, **33**,
- 516 1097–1101. https://doi.org/10.1046/j.1365-2109.2002.00768.x
- 517 Helland, S. J., Grisdale-Helland, B., & Nerland, S. (1996). A simple method for the
- 518 measurement of daily feed intake of groups of fish in tanks. *Aquaculture*, **139**, 157–163.
- 519 https://doi.org/10.1016/0044-8486(95)01145-5
- 520 Hemre, G. I., & Sandnes, K. (1999). Effect of dietary lipid level on muscle composition in
- 521 Atlantic salmon *Salmo salar*. *Aquaculture Nutrition*, **5**, 9–16.

- 522 https://doi.org/10.1046/j.1365-2095.1999.00081.x
- 523 Iwama, G. K., & Tautz, A. F. (1981). A Simple Growth Model for Salmonids in Hatcheries.
- 524 *Canadian Journal of Fisheries and Aquatic Sciences*, **38**, 649–656.
- 525 https://doi.org/10.1139/f81-087
- 526 Jobling, M. (2002). Environmental factors and rates of development and growth. In *Handbook*
- 527 *of fish biology and fisheries. Volume 1: Fish biology* (pp. 97–122).
- 528 https://doi.org/10.1002/9780470693803
- Jobling, M., & Johansen, S. J. S. (1999). The lipostat, hyperphagia and catch-up growth.
- 530 *Aquaculture Research*, **30**, 473–478. https://doi.org/10.1046/j.1365-2109.1999.00358.x
- Jobling, M., & Miglavs, I. (1993). The size of lipid depots a factor contributing to the
- control of food intake in Arctic charr, *Salvelinus alpinus? Journal of Fish Biology*, 43,
  487–489.
- Johansen, S. J. S., Ekli, M., & Jobling, M. (2002). Is there lipostatic regulation of feed intake
- in Atlantic salmon *Salmo salar* L.? *Aquaculture Research*, **33**, 515–524.
- 536 https://doi.org/10.1046/j.1365-2109.2002.00736.x
- Johansen, S. J. S., Ekli, M., Stangnes, B., & Jobling, M. (2001). Weight gain and lipid
- deposition in Atlantic salmon, *Salmo salar*, during compensatory growth: Evidence for
- 539 lipostatic regulation? *Aquaculture Research*, **32**, 963–974.
- 540 https://doi.org/10.1046/j.1365-2109.2001.00632.x
- Johansen, S. J. S., Sveier, H., & Jobling, M. (2003). Lipostatic regulation of feed intake in
  Atlantic salmon *Salmo salar* L. defending adiposity at the expense of growth?
- 543 *Aquaculture Research*, **34**, 317-331. https://doi.org/10.1046/j.1365-2109.2003.00821.x
- Johnsson, J. I., & Bohlin, T. (2005). Compensatory growth for free? A field experiment on
- 545 brown trout, *Salmo trutta*. *Oikos*, **111**, 31–38. https://doi.org/10.1111/j.0030-
- 546 1299.2005.13972.x

- Johnsson, J. I., & Bohlin, T. (2006). The cost of catching up: increased winter mortality
- 548 following structural growth compensation in the wild. *Proceedings. Biological Sciences* /
- 549 *The Royal Society*, **273**, 1281–6. https://doi.org/10.1098/rspb.2005.3437
- Johnsson, J. I., Jönsson, E., & Björnsson, B. T. (1996). Dominance, nutritional state, and
- growth hormone levels in rainbow trout (*Oncorhynchus mykiss*). *Hormones and*
- 552 *Behavior*, **30**, 13–21. https://doi.org/10.1006/hbeh.1996.0003
- Jonsson, N., Jonsson, B., & Hansen, L. P. (1997). Changes in proximate composition and
  estimates of energetic costs during upstream migration and spawning in Atlantic salmon
- 555 Salmo salar. Journal of Animal Ecology, **66**, 425–436. https://doi.org/10.2307/5987
- 556 Kadri, S., Mitchell, D. F., Metcalfe, N. B., Huntingford, F. A., & Thorpe, J. E. (1996).
- 557 Differential patterns of feeding and resource accumulation in maturing and immature
- Atlantic salmon, *Salmo salar*. *Aquaculture*, **142**, 245–257. https://doi.org/10.1016/00448486(96)01258-6
- 560 Keesey, R. E., & Corbett, S. W. (1984). Metabolic defense of the body weight set-point.
- 561 *Research Publications Association for Research in Nervous and Mental Disease*, **62**,
- 562 87–96. Retrieved from http://europepmc.org/abstract/MED/6695117
- 563 Kennedy, G. C. (1953). The Role of Depot Fat in the Hypothalamic Control of Food Intake in
- the Rat. *Proceedings of the Royal Society B: Biological Sciences*, **140**, 578–592.
- 565 https://doi.org/10.1098/rspb.1953.0009
- 566 Maclean, A., & Metcalfe, N. B. (2001). Social status, access to food, and compensatory
- 567 growth in juvenile Atlantic salmon. *Journal of Fish Biology*, **58**, 1331–1346.
- 568 https://doi.org/10.1111/j.1095-8649.2001.tb02290.x
- 569 Mason, M. E., & Waller, G. R. (1964). Dimethoxypropane Induced Transesterification of Fats
- and Oils in Preparation of Methyl Esters for Gas Chromatographic Analysis. *Analytical*
- 571 *Chemistry*, **36**, 583–586. https://doi.org/10.1021/ac60209a008

- 572 McLoughlin, M. F., Nelson, R. N., McCormick, J. I., Rowley, H. M., & Bryson, D. B. (2002).
- 573 Clinical and histopathological features of naturally occurring pancreas disease in farmed
- 574 Atlantic salmon, *Salmo salar* L. *Journal of Fish Diseases*, **25**, 33–43.
- 575 https://doi.org/10.1046/j.1365-2761.2002.00334.x
- Metcalfe, N. B., & Monaghan, P. (2001). Compensation for a bad start: Grow now, pay later? *Trends in Ecology and Evolution*, 16, 254-260. https://doi.org/10.1016/S0169-

578 5347(01)02124-3

579 Metcalfe, N. B., & Thorpe, J. E. (1992). Anorexia and defended energy levels in over-

580 wintering juvenile salmon. *Journal of Animal Ecology*, **61**, 175–181.

- 581 https://doi.org/10.2307/5520
- 582 Mørkøre, T., & Rørvik, K. -A. (2001). Seasonal variations in growth, feed utilisation and
- product quality of farmed Atlantic salmon (*Salmo salar*) transferred to seawater as 0 +

smolts or 1 + smolts. *Aquaculture*, **199**, 145–157. https://doi.org/10.1016/S0044-

- 585 8486(01)00524-5
- 586 Norwegian Standards Association, 1994. Norwegian Quality Cut [NQC] NS 9401 (Oslo).
- 587 Oehme, M., Grammes, F., Takle, H., Zambonino-Infante, J. L., Refstie, S., Thomassen, M. S.,

588 Rørvik, K.-A & Terjesen, B. F. (2010). Dietary supplementation of glutamate and

- arginine to Atlantic salmon (*Salmo salar* L.) increases growth during the first autumn in
- sea. *Aquaculture*, **310**, 156–163. https://doi.org/10.1016/j.aquaculture.2010.09.043
- 591 Rowe, D. K., & Thorpe, J. E. (1990). Suppression of maturation in male Atlantic salmon
- 592 (*Salmo salar* L.) parr by reduction in feeding and growth during spring months.
- 593 *Aquaculture*, **86**, 291–313. https://doi.org/10.1016/0044-8486(90)90121-3
- Rowe, D. K., Thorpe, J. E., & Shanks, A. M. (1991). Role of Fat Stores in the Maturation of
- 595 Male Atlantic Salmon (*Salmo salar*) Parr. *Canadian Journal of Fisheries and Aquatic*
- 596 *Sciences*, **48**, 405–413. https://doi.org/10.1139/f91-052

- coho salmon: the effect of food distribution and ration size. *Journal of Fish Biology*, **48**,
- 599 686–694. https://doi.org/10.1111/j.1095-8649.1996.tb01464.x
- 600 Rørvik, K.-A., Ytrestøyl, T., Lundberg, E., Jakobsen, F. A., Jakobsen, A. A., & Bjerkeng, B.
- 601 (2010). How Apparent Digestibility of Carotenoids, Macronutrients, and Minerals are
- 602 Differently Affected by Ration Level in Atlantic Salmon, *Salmo Salar. Journal of*

603 *Applied Aquaculture*, **22**, 123–139. https://doi.org/10.1080/10454431003736227

- 604 Røsjø, C., Berg, T., Manum, K., Gjøen, T., Magnusson, S., & Thomassen, M. S. (1994).
- Effects of temperature and dietary n-3 and n-6 fatty acids on endocytic processes in
- 606 isolated rainbow trout (Oncorhynchus mykiss, Walbaum) hepatocytes. Fish Physiology
- 607 *and Biochemistry*, **13**, 119–32. https://doi.org/10.1007/BF00004337
- Silverstein, J. T., Shearer, K. D., Dickhoff, W. W., & Plisetskaya, E. M. (1999). Regulation of
  nutrient intake and energy balance in salmon. *Aquaculture*, **177**, 161-169.
- 610 https://doi.org/10.1016/S0044-8486(99)00076-9
- Taksdal, T., Olsen, A. B., Bjerkås, I., Hjortaas, M. J., Dannevig, B. H., Graham, D. A., &
- 612 McLoughlin, M. F. (2007). Pancreas disease in farmed Atlantic salmon, Salmo salar L.,
- and rainbow trout, Oncorhynchus mykiss (Walbaum), in Norway. Journal of Fish
- 614 *Diseases*, **30**, 545–558. https://doi.org/10.1111/j.1365-2761.2007.00845.x
- Taranger, G. L., Carrillo, M., Schulz, R. W., Fontaine, P., Zanuy, S., Felip, A., Weltzien, F.-
- A., Dufour, S., Karlsen, Ø., Norberg, B., Andersson, E., & Hansen, T. (2010). Control of
- 617 puberty in farmed fish. *General and Comparative Endocrinology*, **165**, 483–515.
- 618 https://doi.org/10.1016/j.ygcen.2009.05.004
- Thomassen, M. S., & Røsjø, C. (1989). Different fats in feed for salmon: Influence on sensory
- parameters, growth rate and fatty acids in muscle and heart. *Aquaculture*, **79**, 129–135.
- 621 https://doi.org/10.1016/0044-8486(89)90453-5

- salmon in closed-containment systems. *Aquaculture*. **312**, 1-14.
- 624 https://doi.org/10.1016/j.aquaculture.2010.11.043
- Thorpe, J. E. (1994). Reproductive strategies in Atlantic salmon, *Salmo salar* L. *Aquaculture Research*, 25, 77–87. https://doi.org/10.1111/j.1365-2109.1994.tb00668.x
- 627 Thorpe, J. E., Mangel, M., Metcalfe, N. B., & Huntingford, F. A. (1998). Modelling the
- 628 proximate basis of salmonid life-history variation, with application to Atlantic salmon,
- 629 *Salmo salar* L. *Evolutionary Ecology*, **12**, 581–599.
- 630 https://doi.org/10.1023/A:1022351814644
- Thorpe, J. E., Talbot, C., Miles, M. S., & Keay, D. S. (1990). Control of maturation in
- 632 cultured Atlantic salmon, *Salmo salar*, in pumped seawater tanks, by restricting food
- 633 intake. *Aquaculture*, **86**, 315–326. https://doi.org/10.1016/0044-8486(90)90122-4
- Torrissen, O., Olsen, R. E., Toresen, R., Hemre, G. I., Tacon, A. G. J., Asche, F., Hardy, R.W.
- 635 & Lall, S. (2011). Atlantic Salmon (*Salmo salar*): The "Super-Chicken" of the Sea?
- 636 *Reviews in Fisheries Science*, **19**, 257–278.
- 637 https://doi.org/10.1080/10641262.2011.597890
- Torstensen, B. E., Lie, Ø., & Frøyland, L. (2000). Lipid metabolism and tissue composition in
- 639 Atlantic salmon (*Salmo salar* L.)- Effects of capelin oil, palm oil, and oleic scid-enriched
- sunflower oil as dietary lipid sources. *Lipids*, **35**, 653–664.
- 641 https://doi.org/10.1007/s11745-000-0570-6
- 642 Torstensen, B. E., Lie, & Hamre, K. (2001). A factorial experimental design for investigation
- of effects of dietary lipid content and pro- and antioxidants on lipid composition in
- 644 Atlantic salmon (*Salmo salar*) tissues and lipoproteins. *Aquaculture Nutrition*, 7, 265–
- 645 276. https://doi.org/10.1046/j.1365-2095.2001.00184.x
- Usher, M. L., Talbot, C., & Eddy, F. B. (1991). Effects of transfer to seawater on growth and

647	feeding in Atlantic salmon smolts (Salmo salar L.). Aquaculture, 94, 309–326.
648	https://doi.org/10.1016/0044-8486(91)90176-8
649	Wright, P. J. (2007). Understanding the maturation process for field investigations of
650	fisheries-induced evolution. Marine Ecology Progress Series, 335, 279-284.
651	https://doi.org/10.3354/meps335279
652	
653	
654	
655	
656	
657	
658	
659	
cc0	
660	
661	
662	
663	
664	
001	
665	
666	

**Table 1.** Biometrics and fat content of Atlantic salmon in August 2011 fed a diet high-fat diet (HF), low-fat high-protein diet (LF) or half ration of the low fat diet high-protein diet (0.5LF) from May until August 2011, referred to as pre-dietary feeding phase. Biometric parameters for all fish are presented as means  $\pm$  SD, whereas biometric parameters and fat content for sampled fish are presented as means  $\pm$  SEM together with indications of significant differences.

Dietary treatment	HF	LF	0.5LF
Biometric parameters, all fish			
Number of fish, n	584	584	602
Bodyweight, g	$2651 \pm 335$	$2506 \pm 287$	$1865 \pm 253$
Fork length, cm	$59.1 \pm 2.3$	$59.1 \pm 2.1$	$55.8 \pm 2.3$
CF	$1.28\pm0.09$	$1.21\pm0.07$	$1.07\pm0.08$
Biometric parameters, sampled fish, $n = 20$			
Bodyweight, g	$2619\pm70^{a}$	$2515\pm 63^a$	$1881\pm47^b$
Fork length, cm	$59.0\pm0.5^{a}$	$59.0\pm0.4^{a}$	$55.7\pm0.5^{b}$
CF	$1.22\pm0.02^{a}$	$1.18\pm0.02^{a}$	$1.03\pm0.01^{b}$
VSI	$11.3\pm0.4^{a}$	$9.6\pm0.2^{b}$	$8.5\pm0.1^{c}$
Fat content, sampled fish, $n = 20$			
Muscle fat, %	$16.4 \pm 0.3^{a}$	$13.1\pm0.2^{b}$	$11.3 \pm 0.3^{c}$
Visceral fat <sup>†</sup> , %	39.0	29.0	26.6

676 CF; condition factor, VSI; Visceral somatic index

<sup>†</sup>The analysis of visceral fat content was conducted on pooled samples in August 2011 (n=1)

678 Values in the same row with different letters are significantly different ( $P \le 0.05$ ) determined by one-way

679 ANOVA followed by Duncan's multiple range test.

680

681

682

**Table 2**. Chemical compositions (wet weight, as is basis) and fatty acid composition (% of

total fatty acids) of the diets used in the main-dietary phase.

686

	7 mm	pellet	9 mm	pellet
Diet code	MO	RO	MO	RO
Chemical composition (wet weight, as is basis)				
Dry matter, %	93.2	94.0	93.8	93.9
Crude protein (N x 6.25), %	41.2	41.7	34.4	34.6
Crude Lipid, %	31.2	31.4	37.0	35.7
Starch, %	6.2	6.1	6.7	6.8
Ash, %	4.8	4.8	5.1	5.1
NFE <sup>†</sup> , %	16.0	16.1	17.3	18.5
Crude protein/lipid ratio	1.32	1.33	0.93	0.97
Calculated values <sup>‡</sup>				
Gross energy, MJ kg <sup>-1</sup>	24.8	25.1	25.7	25.5
DP, g kg <sup>-1</sup>	354	359	296	298
DE, MJ kg <sup>-1</sup>	21.4	21.5	22.2	21.9
DP/DE ratio, g MJ kg <sup>-1</sup>	16.6	16.6	13.3	13.6
Fatty acid composition (% of total fatty acids)				
C 16:0	12.7	8.5	14.3	9.3
C 18:0	3.2	2.7	3.7	2.9
∑SFA <sup>§</sup>	22.6	15.1	24.0	15.9
C 18:1 n-9	26.8	42.1	23.3	42.5
ΣMUFA¶	38.1	49.8	36.2	52.8
C 18:2 n-6	8.1	13.9	7.4	13.9
C 18:3 n-3	3.4	6.5	2.9	6.0
C 20:5 n-3	10.1	4.6	11.1	4.0
C 22:5 n-3	1.3	0.6	1.4	0.5
	72	3.5	7.5	3.6
C 22:6 n-3	1.2			
C 22:6 n-3 ∑PUFA <sup>¥</sup>	34.3	30.4	32.7	29.0
C 22:6 n-3 ΣPUFA <sup>¥</sup> SUM EPA+DHA	34.3 17.4	30.4 8.1	32.7 18.6	29.0 7.5

687 MO; Marine oil profile, RO: Rapeseed oil profile, N; Nitrogen, NFE; Nitrogen-free extracts, DP; digestible 688 protein, DE; digestible energy, MJ; Mega joule, SFA; Saturated fatty acids, MUFA; monounsaturated fatty acids,

689 PUFA; polyunsaturated fatty acids.

690 <sup>†</sup>NFE was calculated as = 100 - (protein+lipids+ash+water)

<sup>\*</sup>Gross energy, DP and DE were estimated assuming 23.7, 39.5 and 17.2 MJ kg<sup>-1</sup> as the gross energy content of

692 protein, lipids and carbohydrates, respectively. The apparent digestibility coefficients (ADCs) for protein and

693 lipids used were 0.86 and 0.94, respectively (Einen & Roem 1997), whereas 0.50 was used for NFE (Arnesen &694 Krogdahl 1993).

695 <sup>§</sup>SFA; C14:0. C15:0, C16:0, C18:0 and 22:0.

696	<sup>1</sup> MUFA; C16:1n-9, C16:1n-7, C17:1n-7,C18:1n-7, C:18:1n-9, C20:1n-7, C20:1n-9,C20:1n-11, C22:1n-
697	9,C22:1n-11,C24:1n-9
698	<sup>¥</sup> PUFA; C16:2n-3, C16:3n-4, C18:2n-6,C18:3n-6, C18:3n-3, C18:4n-3, C20:4n-3, C20:2n-6, C20:3n-6, C20:4n-
c	

6,C20:5n-3, C22:5-n-3, C22:6n-3.

_		
7	$\Omega \Omega$	
'	00	

### 718 FIGURE CAPTIONS:

719

**Fig. 1** Schematic overview of the experimental design during the pre- and the main-dietary phase. The squares during the pre-dietary phase represent net-pens fed different diets; HF =high fat diet (black filled square), LF = low fat diet (grey filled square), 0.5LF = half ration of the low-fat diet (white filled square). The large squares in the main-dietary phase represents the net-pens and the squares within the net-pens are the pre-dietary groups.

725

Fig. 2 Ambient daily sea temperature (°C, y-axis) and hours of daylight (hours, z-axis) during
the pre-dietary phase (May to August 2011) and the main-dietary phase (August 2011 to
March 2012). The length of the different periods are indicate by the different grey colours
(light grey = pre-dietary phase, dark grey = main-dietary phase).

730

**Fig. 3** Regression line between thermal growth coefficients (TGC) and the increase in muscle fat (%) from August to October in Atlantic salmon fed three different pre-dietary treatment from May to August 2011; high fat diet (black filled squares) = HF, low fat diet (grey filled triangles) = LF, half ration of the low-fat diet (white filled circles) = 0.5LF. Each point represents average per fish group/experimental unit (n = 12).

736

**Fig. 4** Muscle fat content (**A1**) and body weight (**B1**) development of Atlantic salmon fed three different pre-dietary treatment from May to August 2011. Values are means  $\pm$  SEM, n = 4 (n = 1 at termination of the pre-dietary phase). Values not sharing common superscript letters within each sampling period are significantly different ( $P \le 0.05$ ). **A2** and **B2**, present the final muscle fat and BW of the groups, respectively. The values 11.3%, 13.2% and 16.4% represent the obtained fat content at the beginning of the main-dietary phase (August 2011)
for the 0.5LF, LF and HF group, respectively. ns; not significant, \*; trend (P < 0.1).</li>

744

745

**Fig. 5** Thermal growth coefficients (TGC) (**A**) and weight gain (kg) (**B**) of Atlantic salmon fed three different pre-dietary treatment from May to August 2011. Values are means  $\pm$  SEM, n = 4. Values not sharing common superscript letters within each sampling period are significantly different ( $P \le 0.05$ ). The values 11.3%, 13.2% and 16.4% represent the obtained fat content at the beginning of the main-dietary phase (August 2011) for the 0.5LF, LF and HF group, respectively.

752

**Fig. 6** Condition factor (CF) (**A1**) and fork length (cm) (**B1**) development of Atlantic salmon. fed three different pre-dietary treatment from May to August 2011. Values are means  $\pm$  SEM, n = 4 (n = 1 at termination of the pre-dietary phase). Values not sharing common superscript letters within each sampling period are significantly different ( $P \le 0.05$ ). **A2** and **B2**, present the final CF and fork length of the groups, respectively. The values 11.3%, 13.2% and 16.4% represent the obtained fat content at the beginning of the main-dietary phase (August 2011) for the 0.5LF, LF and HF group, respectively.

760

**Fig. 7** Visceral fat development of Atlantic salmon fed three different pre-dietary treatment from May to August 2011. Values are means  $\pm$  SEM, n = 4 (n = 1 at termination of the predietary phase). Values not sharing common superscript letters within each sampling period are significantly different ( $P \le 0.05$ ). The values 11.3%, 13.2% and 16.4% represent the obtained fat content at the beginning of the main-dietary phase (August 2011) for the 0.5LF, LF and HF group, respectively.

767	
768	Fig. 8 Variation in body weight (gram) (A) and condition factor (CF) (B) assessed using
769	coefficient of variation (CV; mean x SD. <sup>-1</sup> ) among Atlantic salmon fed three different pre-
770	dietary treatment from May to August 2011. Values are means $\pm$ SEM, n = 4 (n = 1 at
771	termination of the pre-dietary phase). Values not sharing common superscript letters within
772	each sampling period are significantly different ( $P \le 0.05$ ). ns; not significant
773	
774	
775	
776	
777	
778	
779	
780	
781	
782	
783	
784	
785	



















□11.3 % (0.5LF) ■13.2 % (LF) ■16.4 % (HF)

