

## Accepted Manuscript

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PII: S0044-8486(07)00336-5  
DOI: doi: [10.1016/j.aquaculture.2007.04.015](https://doi.org/10.1016/j.aquaculture.2007.04.015)  
Reference: AQUA 627633

To appear in: *Aquaculture*

Received date: 27 February 2007  
Revised date: 3 April 2007  
Accepted date: 10 April 2007



Please cite this article as: Palstra, Arjan, Curiel, Danilo, Fekkes, Madelon, de Bakker, Merijn, Székely, Csaba, van Ginneken, Vincent, van den Thillart, Guido, Swimming stimulates oocyte development in European eel, *Aquaculture* (2007), doi: [10.1016/j.aquaculture.2007.04.015](https://doi.org/10.1016/j.aquaculture.2007.04.015)

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**Swimming stimulates oocyte development in European eel**

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18 Running head: Swimming induced maturation in eel

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**Abstract**

20

21 In this study, we subjected eels from Lake Balaton (Hungary) to a swimming  
22 period of one, two or six weeks. Most eels were silver and were 13 - 21 years old. Time  
23 dependent changes in morphometrical parameters and developmental characteristics of  
24 the oocytes were determined. Already after one week of swimming, the gonadal mass  
25 increased and oocytes became larger, filled with large numbers of lipid droplets. After  
26 two and six weeks of swimming we found in addition a significant enlargement of the  
27 eyes, which is a sign of sexual maturation. In contrast to the resting eels, that had oocytes  
28 in the primary growth phase (stage 1 - 2); the swimming eels had oocytes in stage 3; the  
29 cortical alveolus or lipid droplet stage. The results indicate that lipid mobilisation induced  
30 by swimming is a requirement for the natural incorporation of lipid droplets in the  
31 oocytes, a crucial step in oocyte maturation. As the Balaton eels responded stronger to  
32 swimming than young farmed eels, it is suggested that older eels are more sensitive for  
33 maturation triggers.

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35 Keywords: *Anguilla anguilla* Linnaeus, reproductive migration, maturation, lipid  
36 mobilisation, silvering, vitellogenesis

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## Introduction

38

39 European eels spend their feeding stage as immature yellow eels in the fresh and  
40 brackish European waters. It appears that at the end of each growth season many eels  
41 cease feeding and metamorphose (silvering) to prepare for oceanic migration. It is  
42 suggested that fat content is a key factor in the onset of migration (Larsson et al., 1990;  
43 Svedäng and Wickström, 1997) since reserves have to suffice to fuel migration (van  
44 Ginneken and van den Thillart, 2000; van den Thillart et al., 2004; van Ginneken et al.,  
45 2005) and are required for successful development of the oocytes. Drastic changes occur  
46 during silvering, most apparent though is the enlargement of the eyes discriminating the  
47 yellow and silver phase (Pankhurst, 1982). Durif (et al., 2005) demonstrated recently that  
48 silvering and migration are closely related processes. As also the pectoral fins become  
49 longer (Durif et al., 2005) and shape changes (Tesch, 2003), Durif et al. (2005) proposed  
50 an index on basis of length, weight, eye diameter and pectoral fin length, which provides  
51 an estimate of the proportion of silver eels that are true migrants. This was needed since  
52 their abundance was overestimated as demonstrated by Svedäng and Wickström (1997)  
53 and Feunteun et al. (2000).

54 The ovaries of silver eels contain oocytes in the first developmental stages  
55 (Adachi et al., 2003) after transformation of the oogonia. Further progression requires  
56 incorporation of lipids droplets (stage 3: cortical alveolus stage or lipid droplet stage) and  
57 vitellogenin. Although separated in other fish species, these two processes occur  
58 simultaneously in artificially matured Japanese eel (Adachi et al., 2003) and European eel  
59 (Palstra et al., 2005), which suggests an unnatural situation. Untreated silver eels are in a

60 prepubertal stage, still far from sexual maturity (Larsen and Dufour, 1993; Dufour, 1994;  
61 Dufour et al., 2003) and remain as such when kept resting in aquaria. Further sexual  
62 development of silver eel appears to be blocked by dopaminergic inhibition of  
63 hypothalamus and pituitary resulting in insufficient FSH and LH levels (Dufour et al.,  
64 2003). This blockage is likely required in order to allow the long spawning migration.  
65 Obviously there must be natural conditions that lead to release of this blockage. As  
66 European eels have to swim about 5,500-km to reach their spawning site, we hypothesize  
67 that swimming is the crucial trigger for releasing the dopaminergic inhibition.

68 Exercise has never been thoroughly investigated as a stimulating factor for  
69 maturation in fish. Eels have to cover an enormous distance to their spawning grounds, it  
70 is therefore likely that swimming is at least one of the factors that triggers natural  
71 maturation. Recently, van Ginneken et al. (2007) observed increased oocyte diameters in  
72 3 year old hatchery eels after swimming 5,500-km. Significantly higher levels of pituitary  
73 LH were found in the exercised group as compared to the controls. Those results indicate  
74 that long term swimming indeed has a stimulatory effect on maturation. Also in a recent  
75 study on swim performance based on short swim trials, we found indications that  
76 swimming triggered sexual maturation in Lake Balaton silver eels (Palstra et al., 2006d);  
77 the GSI in the swim group was higher than in rest group. Apparently Balaton eels are  
78 more sensitive to swimming than the farmed eels used by van Ginneken et al. (2007), as  
79 they responded already within a week. Therefore we used Lake Balaton silver eels in this  
80 study to investigate the effect of short swim trials (1-6 weeks) on oocyte development.

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## Materials & Methods

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### *Experimental animals*

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### *Swim-flumes & conditions*

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At the end of August 2003 and end of September 2004, a total of 120 eels were caught by electrofishing in Lake Balaton, Hungary, in the region of Keszthely and Tihany. They were transported to the laboratory in oxygen-filled plastic bags and marked individually by subcutaneous injection of PIT-TAGS (TROVAN, Aalten, The Netherlands) just behind the head. Eels were then packed into large oxygen-inflated nylon bags in Perspex and cardboard boxes after which they were sent to Leiden (The Netherlands) in early September 2003 and early October 2004 respectively.

Swim experiments were performed in 22 Blazka-type calibrated swim-flumes described in detail by van den Thillart et al. (2004). Swim-flumes were oriented towards the Sargasso Sea (WNW) in a climatized room of about 100 m<sup>2</sup>. The total water content of about 7000 l was recirculated continuously over a bio-filter. The illumination in the climatized room was switched to 670 nm light (bandwidth 20 nm). Based on eye pigment changes during silvering, it can be assumed that this far-red light is invisible for eels (Pankhurst and Lythgoe 1983). Indeed, the animals did not respond to movements of the experimenter during red light illumination. Experiments were performed in air-saturated (>75%) fresh water at 18 ± 0.5°C. The swim-flumes were 2 m long with a volume of 127 l, with an animal compartment of 110 cm suited for 1 individual eel.

106 *Protocol experiment 1*

107 Protocol 1 was performed in 2003 as part of a study on the influence of the swim-  
108 bladder parasite *Anguillicola crassus* on the swim performance (Palstra et al., 2006d). As  
109 part of the swim performance protocol, fish were exposed to up to 12 h swimming per  
110 day over a 7 day period that cumulatively examined the effect of a week of heightened  
111 and sustained but non-continuous swimming activity. The results are included here as  
112 they provided the basis for longer term experiments described in the following protocol.

113 For the control group 10 eels were upon arrival randomly chosen, then  
114 anaesthetized, measured, sacrificed and sampled. Before handling eels were anaesthetized  
115 with 1 - 1.5 ml diluted clove oil per 1 water (oil of cloves: ethanol = 1:10). Forty eels in  
116 two groups of 20 were subjected to a swim fitness test as described by Palstra et al.  
117 (2006c). In short, eels were anaesthetized, measured and introduced into swim-flumes  
118 two days before the experiment started.

119 The swim fitness protocol consisted of 7 daily experimental trials: 2 speed tests  
120 and 5 endurance tests. On day 1, eels were subjected to a first speed test. Eels started to  
121 swim at a speed ( $U$ ) of  $0.5 \text{ m s}^{-1}$  for 2 h. After these 2 h at  $0.5 \text{ m s}^{-1}$ ,  $U$  was raised with  $0.1$   
122  $\text{m s}^{-1}$  to  $0.6 \text{ m s}^{-1}$  for 2 h. Subsequently, this was repeated with steps of  $0.1 \text{ m s}^{-1}$  for a  $U$   
123 up to  $1.0 \text{ m s}^{-1}$ . On day 2 the eels were subjected to endurance tests starting at  $0.5 \text{ m s}^{-1}$   
124 for 12 h. On the following days the eels were swum at respectively 0.6, 0.7, 0.8, and 0.9  
125  $\text{m s}^{-1}$ . On day 7, the protocol was finished with a second speed test. When fish fatigued  
126 during trials, the flow was lowered to  $0.1 \text{ m s}^{-1}$ .

127 A group of 10 eels were kept resting during the same period in a 1500 l tank  
128 connected to a 2400 l recirculation system under dark conditions. PVC pipes were added

129 as shelter. At the end of the swim test both swimming and resting eels were  
130 anaesthetized, measured, sacrificed and sampled.

131

### 132 *Protocol experiment 2*

133 After arrival, 10 randomly chosen eels were anaesthetized, measured, sacrificed  
134 and sampled as control group. Twenty-one randomly chosen eels were measured and  
135 introduced in the swim-flumes. They were allowed to swim continuously at a speed of  
136  $0.5 \text{ BL s}^{-1}$ . Six eels showed problems with swimming during the experiment and were  
137 stopped. We investigated the swim-bladder of these eels and found that three out of six  
138 eels were infected, and that five out of six eels had heavily damaged swim-bladders  
139 confirming the negative interference of the parasite with swimming (Palstra et al.,  
140 2006d). Only one out of twenty-one eels showed problems with swimming that could not  
141 be connected with swim bladder dysfunction. Data of these eels were not included for  
142 analysis. Resting eels were kept as described above. After two weeks, 6 randomly chosen  
143 swimming eels were stopped, anaesthetized, measured, sacrificed and sampled as well as  
144 a group of 10 resting eels. After six weeks, the remaining 9 eels were stopped,  
145 anaesthetized, measured, sacrificed and sampled as well as a group of 6 remaining resting  
146 eels.

147

### 148 *Measurements & sampling*

149 Morphometric parameters included body-length (BL), body-weight (BW), eye  
150 diameter horizontal (EDh), eye diameter vertical (EDv), pectoral fin length (PFL) and  
151 pectoral fin width (PFW; see for abbreviations also Table 1). The following indices were

152 calculated according to the formulae below: condition factor (K), eye index (EI), pectoral  
153 fin length index (PFLI) and pectoral fin width index (PFWI).

154

155 1. Condition factor (K) =  $100 * BW BL^{-3}$

156 BW: body weight (g), BL: body length (cm)

157

158 2. Eye index (EI) =  $100 * (((EDh + EDv) * 0.25)^2 p * (10 * BL)^{-1})$

159 EDh: eye diameter horizontal (mm), EDv: eye diameter vertical (mm)

160

161 3. Pectoral fin length index (PFLI) =  $100 * PFL BL^{-1}$

162 PFL: pectoral fin length (cm)

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164 4. Pectoral fin width index (PFWI) =  $100 * PFW BL^{-1}$

165 PFW: pectoral fin width (cm)

166

167 The silver index (SI) was calculated according to the procedure described by Durif et al  
168 (2005).

169 Blood samples were taken from the caudal vein with heparin flushed (10.000 IU  
170  $ml^{-1}$ ) 1 ml syringes, which were immediately replaced on ice. The blood was centrifuged  
171 for 5 min at 14,000 rpm and bloodplasm was stored at -80°C.

172 Liver, digestive tract and gonads were dissected and weighed. The following  
173 indices were calculated according to the formulae below: the gonadosomatic index (GSI),

174 the digestive tract somatic index (DTSI) and the hepatosomatic index (HSI; see for  
175 abbreviations also Table 1).

176

177 5. Gonadosomatic index (GSI) =  $(GW BW^{-1}) * 100\%$

178 GW: gonad weight (g), BW: body weight (g)

179

180 6. Digestive tract somatic index (DTSI) =  $(DTW BW^{-1}) * 100\%$

181 DTW: weight digestive tract (g)

182

183 7. Hepatosomatic index (HSI) =  $(LW BW^{-1}) * 100\%$

184 LW: liver weight (g)

185

186 Gonads of all eels solely contained oocytes so all eels were females. From the gonad  
187 at a standardised posterior location a piece of about 1 g was placed in Bouin solution at  
188 room temperature. From all swimming eels, the otoliths (sagitta) were dissected for age  
189 determination.

190

#### 191 *Otolithometry*

192 Age estimation was carried out in the laboratory of Cemagref, Bordeaux, France  
193 by otolithometry according to the method described by Daverat (2005). After extraction,  
194 otoliths were cleaned of all organic matter in distilled water, dried with ethanol, and then  
195 stored in eppendorf tubes. The otoliths were later embedded in synthetic resin (Synolith,  
196 Euroresins Benelux BV, the Netherlands), then polished to the nucleus with a polishing

197 wheel (Streuers Rotopol-35, West Lake, Ohio, USA) using 2 different grits of sandpaper  
198 (1200 and 2400). Fine polishing was done by hand with  $\text{Al}_2\text{O}_3$  (1  $\mu\text{m}$  grain) on a  
199 polishing cloth. Etching was done using 5% EDTA. A drop of this solution was applied  
200 on the mold for 3 min. The otoliths were then rinsed with distilled water and stored in dry  
201 conditions. Year rings were visualised by staining with a drop of 5% Toluidine blue and  
202 counted under a microscope. The age of each eel was determined by the number of  
203 increments starting from the nucleus which was considered as year one of the eel's life.

204

### 205 *Histology*

206 To remove the Bouin fixative, the gonads were washed in 0.1 M phosphate buffer  
207 and 70% ethanol until the solution became transparent. After dehydrating through an  
208 accumulating alcohol series the samples were embedded in air-free Technovit 7100  
209 (Kulzer Histo-Technik) in Peel-A-way molds (Polysciences Inc.) covered with a layer of  
210 air-free paraffin oil during polymerization to exclude oxygen (de Jonge et al., 2005).  
211 Sections of 10  $\mu\text{m}$  thick were cut using a sledge microtome (Jung Polycut E). Three  
212 sections were put on a slide and five slides per sample were made and stained with  
213 Mayers Haematoxylin-Eosin. The oocytes were studied visually under the microscope  
214 (Nikon Eclipse E400) and overview pictures were taken (Nikon Coolpix 4500). Per  
215 section, ten (protocol 1) or twenty (protocol 2) oocytes were randomly selected that were  
216 cut through the nucleus. For each oocyte the developmental stage (OS; see for  
217 abbreviations also Table 1) was determined. The diameter of the oocytes (OD) was  
218 determined using UTHSCSA Image Tool 2.0. The number and diameter of lipid droplets  
219 occurring in stage 3 oocytes were measured with the same method. In fact, what we call

220 lipid droplets are the empty lipid vesicles from which the lipids have been extracted by  
221 washing in ethanol.

222

### 223 *Statistics*

224 Normality of the data and homogeneity of variances were checked by  
225 Kolmogorov-Smirnov tests. Paired observations before and after swimming of  
226 morphometric parameters were tested with student t-tests with one-tailed probabilities.  
227 For SI, a Wilcoxon test with one-tailed probabilities was used.

228 With an univariate general linear model (GLM), analysis of covariance  
229 (ANCOVA) with one-tailed probabilities was performed on log transformed unpaired  
230 observations in search for group effects in swim parameters with either BL (for PFL,  
231 PFW, ED) or BW (for GW, DTW, LW, OD, number lipid droplets, diameter lipid  
232 droplets; for abbreviations see list) as cofactors. ANCOVA was similarly performed for  
233 comparison between the two weeks and the six weeks swim groups. In case of occurrence  
234 of significant group effects, ANOVA with a post-hoc Bonferroni test was performed to  
235 specify the effects between particular groups. ANCOVA was especially required for the  
236 scale difference between swim and rest groups in experiment 2 (Quinn and Keough,  
237 2002).

238 Kruskal-Wallis tests with one-tailed probabilities were performed for comparison  
239 of SI and oocyte stage. Spearman correlation tests with one-tailed probabilities were  
240 performed between start parameters (BL\*, BW\*) vs. silvering parameters (EI\*, SI\*, HSI,  
241 DTSI) vs. maturation parameters (GSI, OS, OD) for control groups, but also for pre-swim  
242 groups for parameters marked with asterisks.

243 Pearson correlation tests with one-tailed probabilities were performed for  
244 comparing between OD and lipid droplet number and size. OD between stages was tested  
245 with student t-tests with one-tailed probabilities. All statistical tests were performed in  
246 SPSS 10.0 for Windows. P-values < 0.05 were considered to indicate statistically  
247 significant differences. Results were calculated and plotted as mean  $\pm$  standard deviation.

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## Results

251

### *Protocol 1*

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In Table 1 the external measurements are given of the control and swim group. The pre-swim and post-swim data are from the same eels. Paired observations showed that after one week of swimming, experimental eels showed a significant weight loss of 15 g (Table 1). Also the K was significantly lower. There was a significant decline of the DTSI (compared to the controls). No increase was observed of silvering indicators.

258 Based on an eye index threshold of 6.5 (Pankhurst, 1982), 70% of all the eels was  
259 scored as silver eels. No change in EI and silver index occurred as a result of one week  
260 swimming. Still, histology data revealed significant increases of oocyte stage and oocyte  
261 diameter. The GSI of eels that had swum was on average 39% higher than control and  
262 rest groups but individual variation remained high and this difference was not significant  
263 (Table 1). Percentages of eels with GSIs >1 were 11% in the rest group and 33% in the  
264 swim group.

265 Eels of the control group in protocol 1 showed oocytes in stage 1, 2 and 3 with the  
266 latter oocytes just having entered this stage containing only a few lipid droplets. The  
267 average diameter of these oocytes was  $103 \pm 36 \mu\text{m}$ . After one week, eels from the swim  
268 group showed a significant change in the average oocyte stage (Table 1). 85% of the eels  
269 in the swim group had ovaries with exclusively stage 3 oocytes with a larger diameter of  
270  $155 \pm 47 \mu\text{m}$  and containing much more lipid droplets (20 - 100).

271

#### 272 *Protocol 2*

273 At the start of the two week protocol the eels were  $62 \pm 4 \text{ cm}$ ,  $347 \pm 76 \text{ g}$  (Table  
274 2) and 83% of them were silver (EI > 6.5). At the start of the six week protocol the eels  
275 were  $63 \pm 5 \text{ cm}$  long, weighted  $429 \pm 137 \text{ g}$  (Table 2) and 33% was silver. This  
276 difference occurred despite the random procedure for assigning eels to the different trials.  
277 Experimental eels were assigned to SI stages 2, 3 and 5, while not any fish was assigned  
278 to stage 4. The experimental eels that had swum aged between 13 and 21 years (mean 16  
279 years; Table 2). Paired observations showed that after two weeks of swimming,  
280 experimental eels showed a significant weight loss of 14 g (Table 2). Also the K was

281 significantly lower. PFLI and PFWI did not show changes. Paired observations showed  
282 that after six weeks of swimming, experimental eels showed a significant weight loss of  
283 63 g (Table 2). Again the K was significantly lower. PFLI and PFWI did not show  
284 changes in the swim group. The PFWI was however smaller in the rest group.

285 The EI increased significantly after 2 week swimming from  $8.32 \pm 2.20$  to  $10.31 \pm$   
286  $2.84$ , also a significant increase in comparison with the rest group (Table 2). After two  
287 weeks, all eels showed increases of the EI between 11 to 41% (Fig. 1), corresponding to a  
288 rise in SI. The EI increased further after six weeks of swimming between 10 up to 66%  
289 (Fig. 1) resulting in 100% silver eels. This change in eye diameter caused a significant  
290 change in the SI.

291 The GSI was found on average 3 - 5 times higher in the two week swim group  
292 than in the controls though not significantly (Table 2). The number of eels with a GSI > 1  
293 was 50% in the swim group, which was significantly different from the control and rest  
294 group. There were no eels at all in the control and rest group with GSI > 1. The GSI in  
295 the six week swim group was significantly higher than in the control and rest group. Also  
296 in comparison with two weeks of swimming, the GSI was significantly higher both in the  
297 swim group and rest group.

298 DTSI and HSI values of the 2 and 6 week swim groups were lower than the  
299 controls but not significantly. HSI was found significantly lower in the rest group both vs.  
300 control and two week swim group. DTSI and HSI were lower in the six week swim group  
301 but only significantly in the rest group. Also in comparison with two weeks of swimming,  
302 the HSI was significantly lower (Table 2).

303

304 *Oocyte developmental status*

305 Eels of the control group contained oocytes representing stages 1, 2 and 3  
306 according to Wallace and Selman (1981), Tyler and Sumpter (1996) and Adachi et al.  
307 (2003).

308 Stage 1 oocytes represented small oocytes (30 - 90  $\mu\text{m}$ ), still in nest structure or  
309 individually organised. An acellular zona radiata (zona pellucida, chorion vitelline  
310 envelop) could not be identified yet. Oocytes were oval shaped and the cytoplasm was  
311 darkly coloured (Fig. 2A). The nucleus was centred and round, contained 1 to 3 large  
312 nucleoli and about 20 smaller ones dispersed throughout the nucleus. The stage 1 oocytes  
313 that were found should be considered late stage 1 oocytes since size was already  
314 considerable in comparison with Tyler and Sumpter (1996) stating that oogonia in  
315 teleosts generally measure less than 10  $\mu\text{m}$  in diameter.

316 Stage 2 oocytes represented larger oocytes (40 - 150  $\mu\text{m}$ ) with less darkly  
317 coloured cytoplasm. In this phase a zona radiata could be identified. The number of  
318 nucleoli was higher. Islands of lightly stained cytoplasm could be distinguished in the  
319 oocyte. Lipid and connective tissue were surrounding the oocytes (Fig. 2A).

320 Stage 3 oocytes were characterised by the presence of lipid droplets. In eels from  
321 control groups, stage 3 oocytes contained very few lipid droplets, generally dispersed in  
322 the periphery of the ooplasm near the zona radiata (Fig. 2B). Mostly the nucleus was  
323 roundly shaped and nucleoli were found in the periphery of the nucleus. Eels from swim  
324 groups contained stage 3 oocytes with a large number of lipid droplets (Fig. 2C).

325 Yolk globuli, such as is characteristic for vitellogenesis, were not found. For  
326 comparison, Fig. 2D shows an oocyte during final maturation in an artificially matured

327 silver eel (Palstra et al., 2005). Numerous yolk globuli could be observed. The number of  
328 lipid droplets is decreased and the size increased indicating extensive fusion of the lipid  
329 droplets. The oocyte showed considerable increase in size indicating a hydration  
330 response. The nucleus (germinal vesicle) was migrating to the periphery (Germinal  
331 Vesicle Migration – GVM).

332 Eels of the control group in experiment 2 contained oocytes in stage 1 and 2,  
333 except for one eel that showed only stage 3 oocytes. The average diameter of the oocytes  
334 was  $83 \pm 39 \mu\text{m}$ . After two weeks of swimming, eels showed oocytes in a more advanced  
335 stage of development. Change in average stage was just not significant, but oocyte  
336 diameter had significantly increased ( $P < 0.01$ ; Table 2). Fifty percent of the eels had  
337 exclusively stage 3 oocytes in the ovaries with large numbers of lipid droplets in contrast  
338 to the eels in control and rest groups which only had stage 1 and 2 oocytes (Fig. 3).

339 After six weeks, eels from the swim group showed further oocyte development.  
340 The oocyte stage in the swim group was significantly different vs. control and rest group  
341 (Table 2). In the swim group, the oocytes were on average larger (not significantly). In  
342 the rest group, eels had only oocytes representing stage 1 and 2. In the swim group, only  
343 three eels showed stage 1 - 2 oocytes. The other six eels showed stage 2 - 3 or 3 oocytes  
344 with large numbers of lipid droplets (Fig. 3). The change in the rest group was also  
345 significantly different from the control group. Average stage and diameter were  
346 significantly higher after six weeks of resting than after two weeks of resting (Table 2).

347

348 *Variation of number and size of lipid droplets in stage 3 oocytes*

349 In total 770 oocytes of 39 experimental eels in protocol 2 were histologically  
350 analysed. 32 eels contained stage 1 and 2 oocytes (without any lipid droplets) with  
351 diameters between 30 and 147  $\mu\text{m}$  ( $n = 615$ ). Stage 1 oocytes were on average  $56 \pm 14$   
352  $\mu\text{m}$  ( $n = 220$ ). Stage 2 oocytes were significantly larger ( $P < 0.001$ ) and on average  $87 \pm$   
353  $23 \mu\text{m}$  ( $n = 395$ ).

354 Ten of the 39 experimental eels also had stage 3 oocytes: nine swimmers and one  
355 control eel. Stage 3 oocytes were on average  $159 \pm 36 \mu\text{m}$  ( $n = 165$ ) and again  
356 significantly larger ( $P < 0.001$ ). A large variation in the number and size of lipid droplets  
357 in stage 3 oocytes was observed (Fig. 4). They contained on average  $45 \pm 30$  lipid  
358 droplets in a range between 6 and 102 which measured on average  $11 \pm 2 \mu\text{m}$  in a range  
359 between 5 and 17  $\mu\text{m}$ . Significant positive correlations were found for these averages per  
360 eel between oocyte diameter and the number of lipid droplets ( $P < 0.001$ ) and between  
361 oocyte diameter and the diameter of the lipid droplets ( $P < 0.001$ ). Also when oocyte data  
362 were pooled for all eels (Fig. 4), we observed a positive correlation between oocyte  
363 diameter and the number of lipid droplets ( $P < 0.01$ ) and between oocyte diameter and the  
364 diameter of the lipid droplets ( $P < 0.01$ ).

365

#### 366 *Correlations between size, silvering and oocyte developmental indicators*

367 Data of experimental eels at the start (control and pre-swim measurements) were  
368 used for correlation analyses. Correlations between size (BL, BW), silvering (externally  
369 EI, SI and internally DTISI, HSI) and oocyte developmental indicators (GSI, OS, OD)  
370 were analysed (Table 3). Significant positive correlations were found between size and  
371 silvering indicators and between size and oocyte developmental indicators suggesting that

372 the silvering and maturation status of larger eels was more advanced. Between silvering  
373 and oocyte developmental indicators, significant positive correlations were found only  
374 between EI, SI and OS (Table 3). Positive correlation with GSI was just not significant.  
375 No significant correlation was found between external indicators of the level of silvering  
376 EI and SI vs. DTSI and HSI, which are often considered to change in relation to the  
377 degree of silvering.

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379

380

### Discussion

381

#### *Swimming triggers silvering*

382  
383 Continuous swimming at  $0.5 \text{ BL s}^{-1}$  resulted in an increase of the eye index (EI),  
384 which was significant after after two and 6 weeks swimming. The observed changes were  
385 even stronger after six weeks of swimming. In contrast no change could be observed in  
386 the EI of the rest groups, indicating that the effect was not caused by time and/or  
387 starvation. As increase of EI has been described with an external parameter for  
388 progression of sexual maturation (Pankhurst, 1982; Tesch, 2003; Durif et al., 2005), these  
389 observations suggest that swimming induces early maturation.

390 The mean eye index of all the eels that just arrived from Lake Balaton in  
391 September was  $8.2 \pm 3.0$ . As an EI of 6.5 is used as the threshold for silvering (Pankhurst,  
392 1982), 59% of the group can be classified as silver eels. The degree of silvering and  
393 oocyte development was positively correlated with body size (Table 3). This observation  
394 of silver eels in Lake Balaton contradicts Bíró (1992), who stated that in Lake Balaton

395 eels never metamorphose into silver eels. However, we found even in the control group  
396 some silver eels in migrant stage 5. These eels had a GSI > 1 and gonads with some stage  
397 3 oocytes, although with few lipid droplets. Surprisingly, the migrant SI stage 4 (Durif et  
398 al., 2005) was not represented at all, not before nor after the swim experiments. This  
399 stage is characterised with elongated pectoral fins, however no changes of the pectoral  
400 fins were observed due to swimming. In field studies it was found that downstream silver  
401 eels have longer pectoral fins, from which it was concluded that swimming likely causes  
402 the fins to grow (Tesch, 2003, Durif et al., 2005). As in our swim trials no change in fin  
403 length occurred, we must conclude that the increase in fin size during downstream  
404 migration must be due to other factors than swimming.

405

#### 406 *Swimming triggers oocyte development*

407 After six weeks of swimming, changes were much more pronounced than after  
408 two weeks of swimming, both GSI and oocyte diameter was significantly higher. More  
409 than 50% of the eels that had swum for two and six weeks had oocytes predominantly in  
410 stage 3, while in contrast resting eels had no stage 3 oocytes at all (Fig. 3). Oocytes in  
411 stage 3 showed a high variation in numbers and diameter of lipid droplets, in other words  
412 in total lipid content. The increase of lipid content is typical for stage 3. As resting eels  
413 showed a slight oocyte development but not any lipid deposition in the oocytes, we  
414 conclude that the deposition of lipids in oocytes occurs under conditions of increased  
415 lipid mobilisation.

416 Recently, van Ginneken et al. (2007) simulated a complete migration of 5,500-km  
417 using 3 year old silver eels from the farm. Those eels were  $71 \pm 4$  cm long and weighed

418 792 ± 104 g, bigger than the ones used in this study. Significant higher levels of pituitary  
419 LH and plasma estradiol were found in the swim group as compared to the controls, also  
420 the oocyte diameter was increased. Those results indicate that long term swimming had  
421 an effect on maturation of younger farmed eels. However, no changes in EI and GSI were  
422 found. Thus, despite the increased hormone levels, the gonads did not develop.

423 The more explicit changes in this study, already after two weeks of swimming,  
424 might be explained by the difference in age. The hatchery eels were young (3 years)  
425 while the Lake Balaton eels in our study were much older (13 to 21 years). This finding  
426 supports the hypothesis that older eels are more responsive and sensitive for maturation.  
427 Durif et al. (2006) observed positive correlations between age vs. condition factor, liver  
428 weight and vitellogenin level, suggesting that older eels are more suited for reproduction.  
429 Indeed, we found recently that older eels require a shorter hormonal treatment to mature  
430 (Palstra et al., 2006b), indicating a higher sensitivity for maturation.

431 In this study, we observed swimming induced oocyte development up to stage 3,  
432 the lipid droplet stage. Stage 3 was found to be highly variable with respect to the  
433 arrangement, number and size of the lipid droplets (Fig. 4). Oocytes of eels that had  
434 swum contained more than 100 larger droplets. However, oocytes did not develop further  
435 than stage 3. Most developed oocytes had lipid droplets that covered >50% of the  
436 cytoplasm and formed a complete ring around the circumference of the oocyte (Couillard  
437 et al., 1997), which is typical for pre-vitellogenic oocytes (Colombo et al., 1984). In  
438 *Anguilla rostrata* this condition is shown to be the start of vitellogenesis (Cottril et al.  
439 2001). However, we did not observe any yolk globuli in the oocytes of eels that had  
440 swum, quite in contrast to artificially matured oocytes which are packed with yolk globuli

441 (Fig. 2D). Also the oocytes did not reach sizes that are characteristic for vitellogenesis.  
442 Vitellogenesis is the major cause for oocyte growth in teleosts in general (Tyler, 1991),  
443 including eel (Nagahama, 1994). Cottril et al. (2001) found oocytes of 200  $\mu\text{m}$  for *A.*  
444 *rostrata* and considered them already vitellogenic. Such values are, however, not typical  
445 for vitellogenesis. Adachi et al. (2003) showed for *Anguilla japonica* that vitellogenesis  
446 begins when oocytes are about 250  $\mu\text{m}$  in diameter. In this study we found maximum  
447 oocyte diameters of 236  $\mu\text{m}$ , which are quite close to the onset of vitellogenesis.

448

449 *Perspectives*

450 The eels in this experiment swam for max 6 weeks, still all individuals showed  
451 progression in sexual maturation. Earlier experiments with 3 year old farmed eels, that  
452 swam for 5,500-km, showed hormonal changes but no increase in gonadal mass.  
453 Therefore it is possible that maturation sensitivity depends on age, and that older eels  
454 may develop further during continued swimming.

455

455 **List of abbreviations**

---

BL	bodylength
BW	bodyweight
DTSI	digestive tract somatic index
DTW	digestive tract weight
ED	eye diameter
EDh	eye diameter horizontal
EDv	eye diameter vertical
EI	eye index
GSI	gonadosomatic index
GW	gonad weight
HSI	hepatosomatic index
K	condition factor
LW	liver weight
OD	oocyte diameter
OS	oocyte stage
PFL	pectoral fin length
PFLI	pectoral fin length index
PFW	pectoral fin width
PFWI	pectoral fin width index
SI	silver index

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456

457 *Acknowledgements*

458 This research was subsidised by the EU contract EELREP no Q5RS-2001-01836. The  
459 authors like to thank Dr. F. Daverat (CEMAGREF) for advice and M. Casteleijn for  
460 assistance with otolithometry. We like to thank R. van der Linden and R. Heymans for  
461 technical support, P. Niemantsverdriet, S. van Schie and L. Wagenaar for animal care,  
462 and E. Clavero and W. Spoor for oocyte histology. We would like to thank I. Báthory, G.  
463 Dobos, G. Ostoros and A. Specziár of the VMRI (Hungarian National Research Fund  
464 OTKA no T45891) for providing Lake Balaton eels. Prof. dr. S. Dufour, Prof. dr. F.  
465 Volckaert and Prof. dr. M. Richardson are greatly acknowledged for critically reading  
466 this manuscript. Experiments comply with the current laws of the Netherlands and were  
467 approved by the animal experimental commission.

468

469

470

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- 567

567 **Table 1** Effect of swimming (protocol 1) on morphometric and gonad parameters (mean  
 568  $\pm$  SD). Control eels (n = 10) were sampled at the start. Swimming eels (n = 40) were  
 569 subjected to a swim fitness test (*see text for explanation*), corresponding to about a week  
 570 of swimming, or kept resting (n = 9). Swimming eels were measured before (pre-swim)  
 571 and after swimming (post-swim). Significant differences ( $P < 0.05$ ) are indicated in bold.  
 572 Symbols mark differences vs. the controls (\*) or pre-swim values (&). (*For abbreviations*  
 573 *see List*)

parameters	control	1 week		
		pre-swim	post-swim	rest
n (eels)	10	40	40	9
<u>external</u>				
BL (cm)	67 $\pm$ 4	69 $\pm$ 6	69 $\pm$ 6	66 $\pm$ 5
BW (g)	475 $\pm$ 77	525 $\pm$ 142	<b>510<math>\pm</math>142</b> &	437 $\pm$ 123
K	0.16 $\pm$ 0.01	0.16 $\pm$ 0.02	<b>0.15<math>\pm</math>0.02</b> &	0.15 $\pm$ 0.02
PFLI	4.68 $\pm$ 0.39	4.96 $\pm$ 0.43	4.88 $\pm$ 0.42	4.68 $\pm$ 0.49
EI	7.38 $\pm$ 3.30	9.14 $\pm$ 2.99	8.64 $\pm$ 3.12	7.31 $\pm$ 2.49
SI	3 $\pm$ 1	4 $\pm$ 1	4 $\pm$ 1	3 $\pm$ 1
<u>internal</u>				
GSI	0.59 $\pm$ 0.34		0.82 $\pm$ 0.43	0.59 $\pm$ 0.32
DTSI	2.42 $\pm$ 0.65		<b>1.66<math>\pm</math>0.67</b> *	2.01 $\pm$ 0.52
HSI	0.92 $\pm$ 0.09		0.84 $\pm$ 0.13	0.86 $\pm$ 0.19
<u>oocytes</u>				
OS	2.7 $\pm$ 0.3		<b>2.8<math>\pm</math>0.4</b> *	<b>2.9<math>\pm</math>0.2</b> *
OD ( $\mu$ m)	103 $\pm$ 36		<b>155<math>\pm</math>47</b> *	128 $\pm$ 48

574

575

575 **Table 2** Effect of swimming (protocol 2) for two weeks and six weeks on morphometric  
 576 and gonad parameters (mean  $\pm$  SD). Control eels (n = 10) were sampled at the start. Eels  
 577 swam two (n = 6) or six weeks (n = 9) at 0.5 BL s<sup>-1</sup> or were kept resting for two (n = 10)  
 578 or six weeks (n = 6). Swimming eels were measured before (pre-swim) and after  
 579 swimming (post-swim). Significant differences (P < 0.05) are indicated in bold. Symbols  
 580 mark differences vs. the controls (\*), pre-swim values (&) or the rest group (#). The 6  
 581 weeks rest group had an average OS value of 1.73 and was found significantly different  
 582 by a Kruskal-Wallis test from the control group. The control group consisted of 9 fish  
 583 that together had an average value of 1.58 with one fish that solely had stage 3 oocytes.  
 584 (*For abbreviations see List*)

parameters	control	2 weeks			6 weeks		
		pre-swim	post-swim	rest	pre-swim	post-swim	rest
n (eels)	10	6	6	10	9	9	6
age (years)			16 $\pm$ 1			16 $\pm$ 3	
<u>external</u>							
BL (cm)	59 $\pm$ 5	62 $\pm$ 4	62 $\pm$ 4	55 $\pm$ 2	63 $\pm$ 5	63 $\pm$ 5	53 $\pm$ 2
BW (g)	267 $\pm$ 68	347 $\pm$ 76	<b>333<math>\pm</math>63</b> <sup>&amp;</sup>	212 $\pm$ 26	429 $\pm$ 137	<b>366<math>\pm</math>114</b> <sup>&amp;</sup>	174 $\pm$ 23
K	0.13 $\pm$ 0.01	0.14 $\pm$ 0.02	<b>0.14<math>\pm</math>0.01</b> <sup>&amp;</sup>	0.13 $\pm$ 0.02	0.16 $\pm$ 0.03	<b>0.14<math>\pm</math>0.02</b> <sup>&amp;</sup>	0.12 $\pm$ 0.02
PFLI	4.55 $\pm$ 0.19	4.89 $\pm$ 0.25	4.89 $\pm$ 0.21	4.39 $\pm$ 0.25	4.78 $\pm$ 0.55	4.78 $\pm$ 0.57	4.33 $\pm$ 0.25
PFWI	2.32 $\pm$ 0.26	2.64 $\pm$ 0.43	2.45 $\pm$ 0.46	2.30 $\pm$ 0.17	2.52 $\pm$ 0.18	2.52 $\pm$ 0.16	<b>2.03<math>\pm</math>0.21</b> <sup>*</sup>
EI	6.20 $\pm$ 1.81	8.32 $\pm$ 2.20	<b>10.31<math>\pm</math>2.84</b> <sup>&amp;#</sup>	5.55 $\pm$ 0.43	6.89 $\pm$ 3.16	<b>9.09<math>\pm</math>2.91</b> <sup>&amp;#</sup>	5.53 $\pm$ 0.60
SI	3 $\pm$ 1	3 $\pm$ 2	4 $\pm$ 1	2 $\pm$ 0	2 $\pm$ 2	<b>3<math>\pm</math>1</b> <sup>&amp;</sup>	2 $\pm$ 0
<u>internal</u>							
GSI	0.26 $\pm$ 0.31		0.74 $\pm$ 0.48	0.13 $\pm$ 0.12		<b>0.80<math>\pm</math>0.35</b> <sup>*#</sup>	0.38 $\pm$ 0.18
DTSI	2.58 $\pm$ 0.81		1.95 $\pm$ 0.77	2.40 $\pm$ 0.52		2.14 $\pm$ 0.54	<b>2.18<math>\pm</math>0.20</b> <sup>*</sup>
HSI	1.13 $\pm$ 0.20		1.02 $\pm$ 0.15	<b>0.87<math>\pm</math>0.09</b> <sup>*</sup>		0.77 $\pm$ 0.13	<b>0.72<math>\pm</math>0.12</b> <sup>*</sup>
<u>oocytes</u>							
OS	1.7 $\pm$ 0.5		2.4 $\pm$ 0.7	1.5 $\pm$ 0.2		<b>2.4<math>\pm</math>0.6</b> <sup>*#</sup>	<b>1.7<math>\pm</math>0.1</b> <sup>*</sup>
OD ( $\mu$ m)	83 $\pm$ 39		<b>136<math>\pm</math>46</b> <sup>#</sup>	61 $\pm$ 14		109 $\pm$ 41	94 $\pm$ 17

585

585 **Table 3** Correlations between parameters of control eels (n = 19 - 20) and eels before  
 586 swimming (n = 75). Significant correlations (P < 0.05) are given in bold. Significant  
 587 correlations were found between size (BL, BW) with silvering (EI, SI) and oocyte  
 588 developmental indicators (GSI, OS, OD). DTSI and HSI were correlated with oocyte  
 589 developmental indicators rather than silvering parameters (*For abbreviations see List*)

		BW	EI	SI	DTSI	HSI	GSI	OS	OD
<b>BL</b>	corr.	<b>0.944</b>	<b>0.740</b>	<b>0.680</b>	-0.218	<b>0.478</b>	<b>0.703</b>	<b>0.900</b>	<b>0.612</b>
	P	<b>0.000</b>	<b>0.000</b>	<b>0.000</b>	0.178	<b>0.017</b>	<b>0.000</b>	<b>0.000</b>	<b>0.003</b>
	n	<b>75</b>	<b>75</b>	<b>75</b>	20	<b>20</b>	<b>20</b>	<b>19</b>	<b>19</b>
<b>BW</b>	corr.		<b>0.729</b>	<b>0.658</b>	-0.147	<b>-0.522</b>	<b>0.734</b>	<b>0.884</b>	<b>0.582</b>
	P		<b>0.000</b>	<b>0.000</b>	0.268	<b>0.009</b>	<b>0.000</b>	<b>0.000</b>	<b>0.004</b>
	n		<b>75</b>	<b>75</b>	20	<b>20</b>	<b>20</b>	<b>19</b>	<b>19</b>
<b>EI</b>	corr.			<b>0.884</b>	-0.211	-0.062	0.311	<b>0.510</b>	0.212
	P			<b>0.000</b>	0.186	0.398	0.091	<b>0.013</b>	0.191
	n			<b>75</b>	20	20	20	<b>19</b>	19
<b>SI</b>	corr.				-0.090	-0.020	0.351	<b>0.603</b>	0.269
	P				0.353	0.467	0.065	<b>0.003</b>	0.133
	n				20	20	20	<b>19</b>	19
<b>DTSI</b>	corr.					-0.013	<b>-0.432</b>	-0.323	-0.203
	P					0.479	<b>0.028</b>	0.088	0.203
	n					20	<b>20</b>	19	19
<b>HSI</b>	corr.						-0.342	<b>-0.390</b>	-0.229
	P						0.070	<b>0.049</b>	0.173
	n						20	<b>19</b>	19
<b>GSI</b>	corr.							<b>0.721</b>	<b>0.730</b>
	P							<b>0.000</b>	<b>0.000</b>
	n							<b>19</b>	<b>19</b>
<b>OS</b>	corr.								<b>0.744</b>
	P								<b>0.000</b>
	n								<b>19</b>

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600 **Captions to figures**

601

602 **Figure 1** Swimming induced changes of eye diameter. Eels were sampled after  
603 swimming continuously at  $0.5 \text{ BL s}^{-1}$  for two and six weeks (also Table 2). The EI had  
604 increased significantly after two and six weeks swimming. This bar graph shows that the  
605 EI of each individual had increased. After two weeks swimming, the EI had increased by  
606  $23.9 \pm 9.6\%$  (range 11 - 41%). After six swimming, the EI had increased by  $36.6 \pm 16.0\%$   
607 (range 10 - 66%). Individual increase was more pronounced after six weeks swimming.

608

609 **Figure 2** Gonad histology. Technovit 7100 sections of  $10 \mu\text{m}$  after staining with Mayers  
610 Haematoxilin-Eosin. A) Oocytes in stage 1 and 2 (small cells with dense cytoplasm and  
611 nucleus), B) Oocytes early stage 3 (larger cells with light cytoplasm, few lipid droplets  
612 swollen nucleus), C) Oocytes late stage 3 (larger cells with a large number of lipid  
613 droplets, swollen nucleus with many nucleoli), D) Oocytes in late stage 5 displaying:  
614 large cells with packed lipid droplets and yolk globuli, and very transparent nucleus in  
615 stage of Germinal Vesicle Migration (GVM). This picture is from an artificially matured  
616 eel, and included for comparison (Palstra et al., 2005). Scale bars are  $100 \mu\text{m}$ . Note that  
617 the mature stage 5 oocyte (D) is much larger than the oocytes in stage 1, 2 and 3 (A, B,  
618 C). Stages were assigned according to Wallace and Selman (1981), Tyler and Sumpter  
619 (1996) and Adachi et al. (2003).

620

621 **Figure 3** Gonad developmental in relation to swimming. The % of silver eels with  
622 oocytes in stage 1 + 2 (white bars), stage 2 + 3 (grey bars), and stage 3 (black bars) is

623 presented in control eels (initial situation), resting eels, and in eels after continuous  
624 swimming at  $0.5 \text{ BL s}^{-1}$  for two and six weeks. In each eel the developmental stage of the  
625 oocytes was found rather homogeneous. The difference in stage for two weeks swimming  
626 was just not significant vs. the controls but significant vs. resting eels ( $P < 0.01$ ). The  
627 difference for 6 weeks swimming was significant vs. the controls ( $P < 0.01$ ) vs. resting  
628 eels ( $P = 0.05$ ).

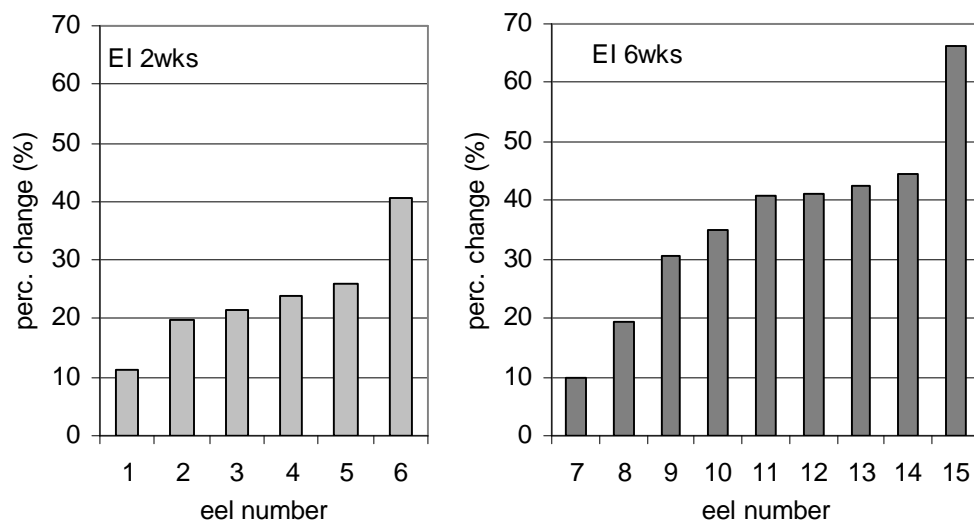
629

630 **Figure 4** Relationship between oocyte diameter and lipid deposition in stage 3 oocytes.  
631 Depicted are the relation between the oocyte diameter and the number of lipid droplets  
632 (upper panel) and the diameter of the lipid droplets (lower panel). All samples originate  
633 from eels from experiment 2. Only eels that had swum had stage 3 oocytes. These eels  
634 had stage 3 oocytes with an average diameter of  $154 \pm 34 \mu\text{m}$  containing  $38.4 \pm 26.8$  lipid  
635 droplets with a diameter of  $10.4 \pm 2.5 \mu\text{m}$ . Significant positive correlations were found  
636 for these averages per eel between oocyte diameter and the number of lipid droplets ( $P <$   
637  $0.001$ ) and between oocyte diameter and the diameter of the lipid droplets ( $P < 0.001$ ).  
638 Representative pictures showing the relation between number and size of lipid droplets  
639 and oocyte diameter are given on the right side. Scale bars are  $50 \mu\text{m}$ .

640

640 **Figure 1**

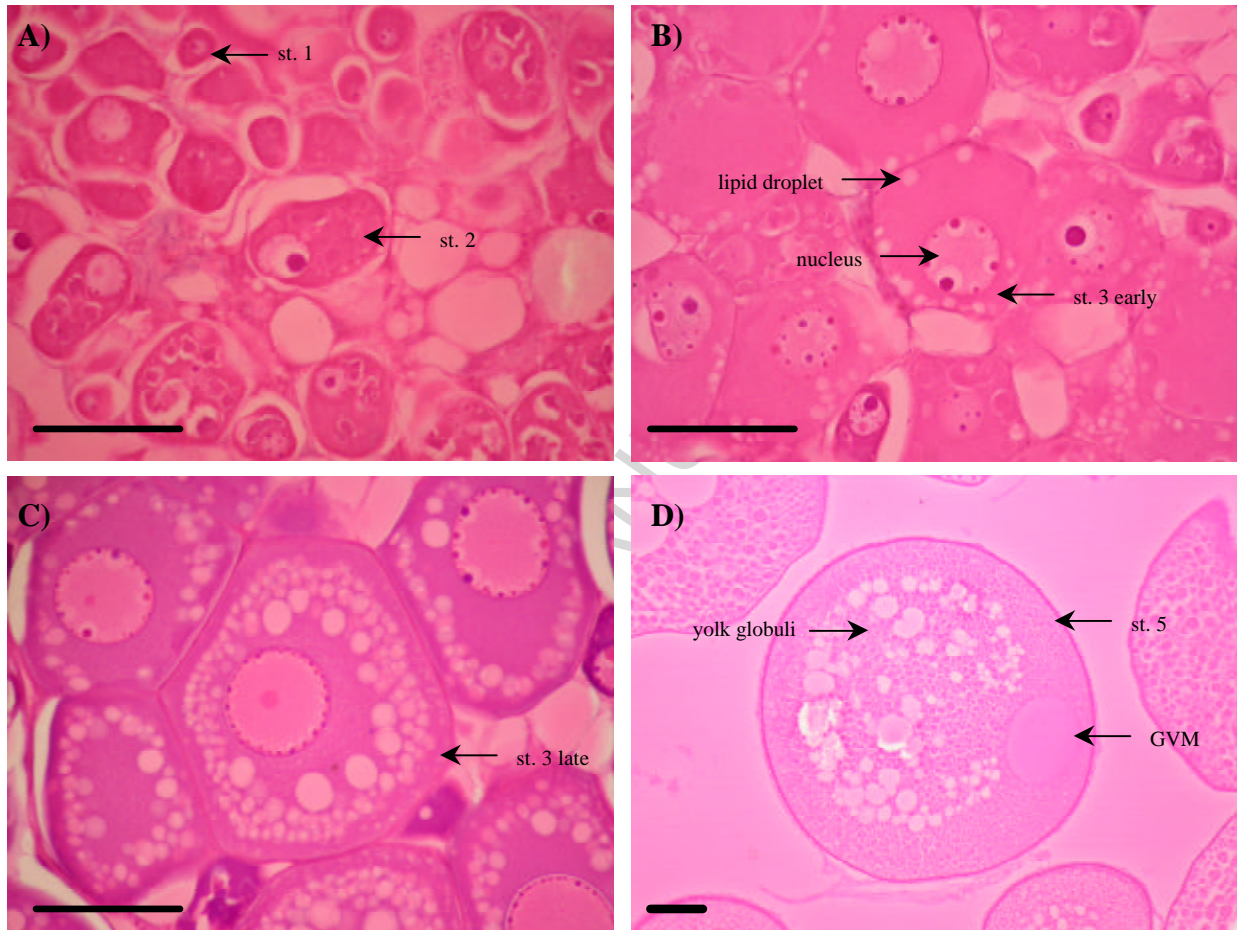
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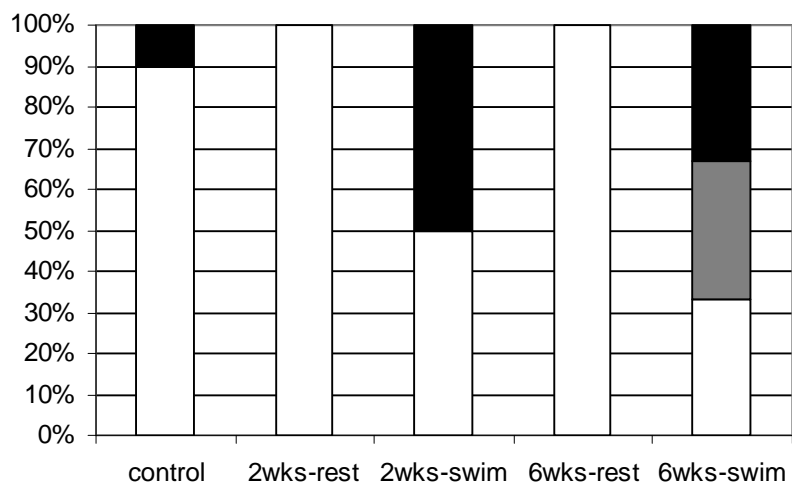
641 **Figure 2**

642

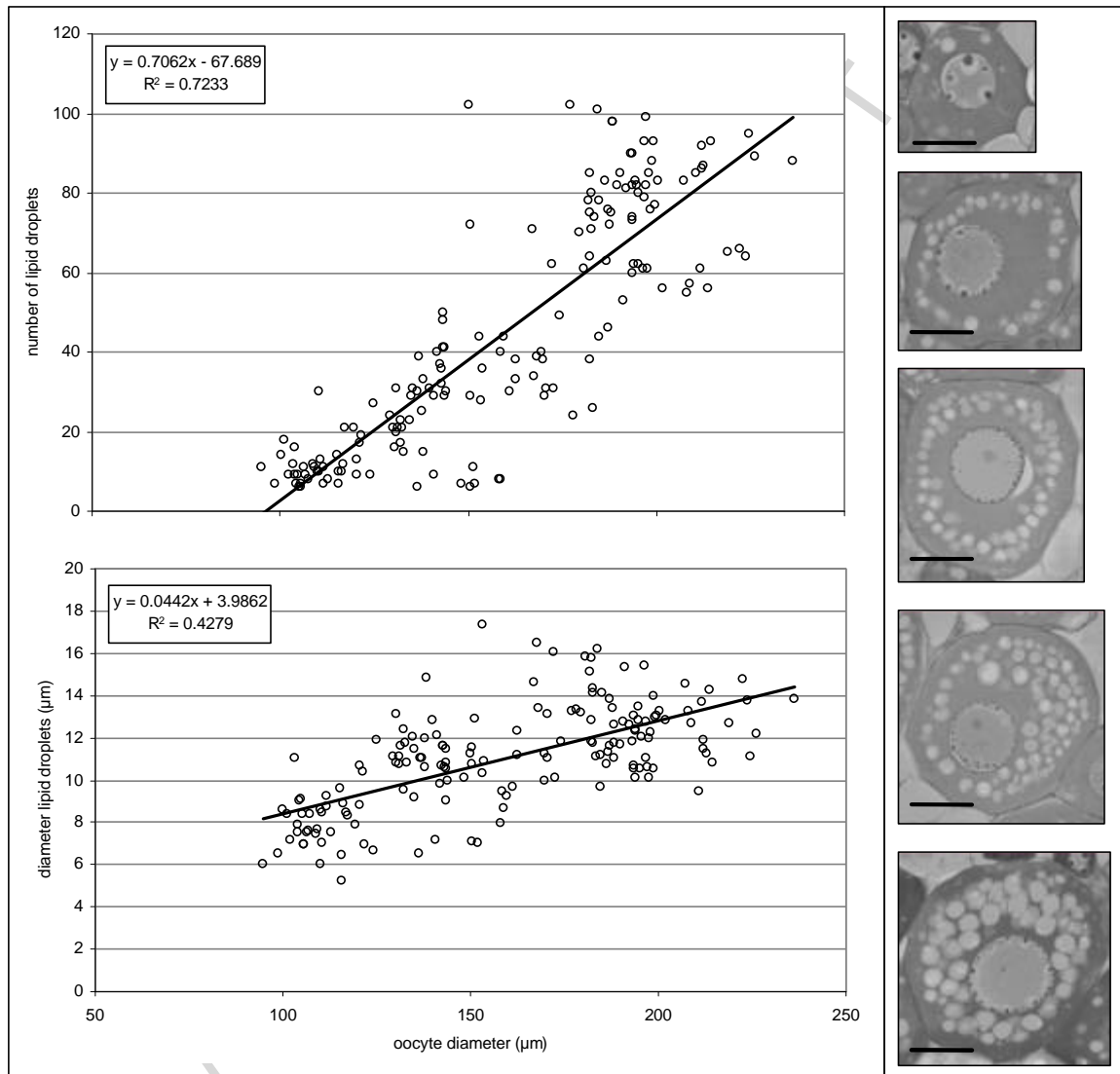


642 **Figure 3**

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643 **Figure 4**

644

645