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Salin, K., Villasevil, E. M., Anderson, G. J., Lamarre, S. G., Melanson, C. A., McCarthy, I., Selman, C. and Metcalfe, N. B. (2019) Differences in mitochondrial efficiency explain individual variation in growth performance. *Proceedings of the Royal Society of London Series B: Biological Sciences*, 286(1909), 20191466.

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1 **TITLE:** Differences in mitochondrial efficiency explain individual variation in growth performance

2

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13

14 **RUNNING TITLE:** ATP/O ratio explains growth performance

15

16 **KEYWORDS:** ATP/O ratio, brown trout, energy metabolism, intraspecific, mitochondrial plasticity,
17 protein synthesis.

18 **ABSTRACT**

19 The physiological causes of intraspecific differences in fitness components such as growth rate are
20 currently a source of debate. It has been suggested that differences in energy metabolism may drive
21 variation in growth, but it remains unclear whether covariation between growth rates and energy
22 metabolism is (i) a result of certain individuals acquiring and consequently allocating more resources
23 to growth, and/or is (ii) determined by variation in the efficiency with which those resources are
24 transformed into growth. Studies of individually-housed animals under standardized nutritional
25 conditions can help shed light on this debate. Here we quantify individual variation in metabolic
26 efficiency in terms of the amount of ATP generated per molecule of oxygen consumed by liver and
27 muscle mitochondria, and examine its effects both on the rate of protein synthesis within these
28 tissues and on the rate of whole-body growth of individually-fed juvenile brown trout (*Salmo trutta*)
29 receiving either a high or low food ration. As expected, fish on the high ration on average gained
30 more in body mass and protein content than those maintained on the low ration. Yet, growth
31 performance varied more than 10-fold amongst individuals on the same ration, resulting in some fish
32 on low rations growing faster than others on the high ration. This variation in growth for a given
33 ration was related to individual differences in mitochondrial properties: a high whole-body growth
34 performance was associated with high mitochondrial efficiency of ATP production in the liver. Our
35 results show for the first time that among-individual variation in the efficiency with which substrates
36 are converted into ATP can help explain marked variation in growth performance, independent of
37 food intake. This study highlights the existence of inter-individual differences in mitochondrial
38 efficiency and its potential importance in explaining intraspecific variation in whole animal
39 performance.

40 INTRODUCTION

41 Individual animals may grow at widely differing rates despite living under the same conditions - a
42 finding that has been documented across a broad range of taxa (reviewed in [1, 2]). This
43 phenomenon is often interpreted in terms of variation in individual quality. For instance, individuals
44 that grow faster typically reach maturity more quickly and can have higher fecundity than slower
45 growing individuals, suggesting direct fitness consequences of growth rate [3, 4]. However, the
46 physiological processes underlying this among-individual variation in growth rate are currently poorly
47 understood.

48 Faster growth can obviously be achieved by increasing food intake. Individuals with high rate of food
49 intake grow faster compared to individuals that have lower rate of resource intake, because high
50 amounts of food intake can lead to increased rate of resource allocation to energetically costly
51 processes, such as biomass production and, in turn, growth. However, variation in growth rate may
52 persist even when food intake is standardised. For example, individual fish fed to satiation and
53 consuming similar amount of food exhibited three-fold differences in growth performance [5].
54 Similarly, five-fold differences in the rate of growth have been shown amongst fish consuming an
55 identical amount of food [6]. This suggests that variation in growth may be, at least partly, attributed
56 to variation in the efficiency of resource utilization and its allocation to biomass production. Yet
57 surprisingly little research has investigated the possible mechanisms that might underlie this
58 variation in metabolic efficiency and thus growth performance [7].

59 Variation in the efficiency with which food is converted to energy is thought to play an important role
60 in the association between food intake and animal growth [7-9]. Energy derived from nutrients
61 becomes usable for cellular processes only following transformation into high-energy molecules of
62 adenosine triphosphate (ATP). ATP is the principal energy source for most cellular functions, such as
63 DNA, RNA and protein synthesis (and hence biomass production). The main sites of energy
64 conversion are the mitochondria, which provide over 90% of a cell's ATP [10]. Mitochondrial ATP is
65 produced via oxidative phosphorylation, a process through which energy substrates are oxidized to
66 generate a proton gradient that drives the phosphorylation of ADP to ATP. Although ATP production
67 depends on the rate of substrate oxidation, the number of ATP molecules produced for each
68 molecule of oxygen and energy substrate (i.e. pyruvate, glutamate, acetyl-CoA, etc) consumed by the
69 mitochondria can vary [11]. A proportion of the energy that is generated from substrate oxidation is
70 dissipated through proton leakage across the inner mitochondrial membrane and this leakage might
71 decrease the energy available to produce ATP [12]. The amount of energy dissipated in the
72 mitochondrial proton leak varies amongst individuals [13, 14] and this variation is known to correlate
73 with animal performance [15, 16]. This raises the possibility that variation in growth among
74 individuals could involve differences in the efficiency through which mitochondria produce ATP.

75 Mitochondrial efficiency can be quantified through measurement of the ATP/O ratio; that is the ratio
76 in the amount of ATP generated per unit of oxygen consumed [17]. Thus, the higher this ratio, the
77 more efficiently an animal converts its metabolic substrates into ATP, with the ATP then available for
78 energy-demanding cellular processes such as protein synthesis and biomass production [18]. A
79 number of studies have found positive links between mean growth rate and mean mitochondrial
80 efficiency when comparing among treatment groups, populations or selection lines [9, 19-23], but
81 until now there has been no assessment of whether mitochondrial efficiency could explain variation
82 in growth rate amongst individual animals maintained with the same food intake.

83 In this study, we tested, for the first time, whether individual variation in growth performance –
84 measured both as the rate of whole-body gain in mass and as the rate of protein synthesis - was
85 related to among-individual variation in mitochondrial efficiency. To test this hypothesis, we
86 assessed the relationships between ATP/O ratio, fractional rate of protein synthesis and growth
87 performance (growth rate, growth efficiency and protein gain) among individually housed brown
88 trout (*Salmo trutta*) of the same age and maintained under standardized conditions. In order to
89 standardize their food intake, fish were fed on individual limited rations to ensure that differences in
90 growth performance could be attributed to mitochondrial efficiency differences. We chose juvenile
91 brown trout as our study organism because larger body size in brown trout is a major determinant of
92 fitness, with fast growth resulting in increased survival [24] and larger body size being linked to
93 higher fecundity [25]. We analysed mitochondrial properties and protein synthesis in the liver and
94 the white muscle, since the physiological properties of these tissues are known to influence growth
95 performance [16, 26]. We predicted positive inter-individual correlations among mitochondrial
96 efficiency, protein synthesis and growth performance.

97

98

99 **MATERIALS AND METHODS**

100 **Experimental animals**

101 Brown trout fry were moved from the hatchery (Howietoun, UK) to the University of Glasgow in June
102 2015. The fish were then kept in a communal tank and maintained under a 12 h light: 12 h dark
103 photoperiod at 12°C and fed daily in excess with trout pellet food (EWOS, West Lothian, UK). In
104 September 2016, fish (n = 60) were transferred to individual compartments within a stream tank
105 system that allowed individual daily feeding while maintaining fish under the same water quality
106 conditions. Each individual compartment contained a small shelter (a section of opaque plastic pipe).

107 The fish were first acclimated for two weeks in their individual compartments, during which they
108 were hand-fed daily to excess on the same trout pellets. Fish were then fasted for 22h and briefly

109 anesthetized (50 ml l⁻¹ benzocaine in water) for measurement of body mass (± 0.001 g) to allow
110 calculation of caloric intake and thereby food rations (as number of pellets). For the next 5-10 weeks
111 (see below) the fish were fed once daily on an intermediate ration of pellets (presumed sufficient for
112 growth but less than a maximal rate of intake) using an equation from Elliott [27]; this allowed
113 calculation of individual-specific rations in calories as a function of the fish's body mass (W) in grams
114 and water temperature (T) of 12°C as follows:

$$115 \quad \text{Intermediate ration} = 24.062 \times W^{0.737} \times \exp(0.105 \times T)$$

116 Fish were fed their ration in the early morning; all fish consumed their entire daily ration within 2 h.
117 Body mass was measured every two weeks, and food rations were recalculated to adjust for gains in
118 mass. Fish were fasted for 22h before each body mass measurement, and on return to their
119 compartment were fed 2 h later than usual to allow time to recover from the anaesthetic and to
120 ensure they ate the ration. All fish consumed their entire daily ration and gained mass during this
121 acclimation period.

122

123 **Diet treatment and growth measurements**

124 Following this period of acclimation to an intermediate diet, fish were switched to the final diet
125 treatment for 14 days. This duration was chosen because it limited the extent of mitochondrial turn-
126 over that would occur over the growth period but was sufficient to detect differences in the rate of
127 growth between individuals [28]. Since only two individuals per day could be analysed for their
128 mitochondrial function at the end of the experiment, the start of the diet treatment was staggered
129 over a 5-week period (so that the preceding acclimation period varied between 5 to 10 weeks). Two
130 fish per day (which would subsequently be processed together 14 days later) were thus randomly
131 allocated to the treatments: one fish had its ration increased to 150% of the intermediate ration
132 (high ration, n = 30) and the other had its ration decreased to 50% of the intermediate ration (low
133 ration, n = 30). The low ration was estimated to provide sufficient energy to cover maintenance
134 requirements and relatively slow growth [27], while the high ration approximated the maximal rate
135 of food intake of juvenile brown trout [27]. Body mass ranged from 3.61 to 15.48 g across individuals
136 at the start of the experiment but did not differ between fish subsequently assigned to the two food
137 treatments (High ration: 8.15 ± 0.49 g, Low ration: 8.18 ± 0.48 g, T test: $t = -0.041$, $df = 58$, $P = 0.967$).
138 Body mass was re-measured (as above) at day 7 of the diet treatment, and rations were recalculated
139 to adjust for growth. All but one fish consumed their entire daily ration within 2 h during the
140 experimental period; this fish was removed from all analyses so giving a final sample size of 59 fish
141 (High food: n = 29; Low food: n = 30).

142 Growth rate and growth efficiency were simultaneously estimated over a 7-day period starting at day
143 7 of the experimental treatment (termed the initial fish mass in the following equation) and ending
144 at day 14 (final fish mass). Specific growth rate (% day⁻¹) was defined as:

$$145 \quad \text{Specific growth rate} = \frac{\ln(\text{final body mass}) - \ln(\text{initial body mass})}{\text{days elapsed}} \times 100$$

146 Daily food intake was calculated from the daily food ration, and was expressed in terms of pellet
147 mass. Growth efficiency (mg gain in body mass mg⁻¹ food eaten) was measured for each fish as:

$$148 \quad \text{Growth efficiency} = \frac{\text{gain in body mass day}^{-1}}{\text{mass of pellets eaten day}^{-1}}$$

149 At the end of the food treatment period, fractional rates of protein synthesis and mitochondrial
150 properties were measured in the fish following protocols described below.

151

152 **Estimate of gain in whole-body protein**

153 The relationship between whole-body protein content and body mass of fish reared under
154 Intermediate, Low and High rations was used to estimate the protein content of each fish at the start
155 and at the end of the diet treatment and thereby estimate the gain in protein content over the
156 treatment period. Specifically, we first determined the relationship between the body mass of a fish
157 and its whole-body protein content (Figure S1), using a separate group of brown trout of the same
158 age and size (See electronic supplementary material – ESM - for full details in section “Whole-body
159 protein content”).

160 The initial whole-body protein content of each experimental fish was therefore estimated from its
161 body mass at the start of the food treatment, using the calibration regression for fish on the
162 intermediate ration. The final whole body protein content of each experimental fish was likewise
163 estimated from its body mass at the end of the food treatment, using the appropriate equation for
164 its diet treatment. Specific protein gain rate (% day⁻¹) was then defined as:

$$165 \quad \text{Specific protein gain} = \frac{\ln(\text{final whole-body protein content}) - \ln(\text{initial whole-body protein content})}{\text{days elapsed}} * 100$$

166

167 **Measurement of the fractional rate of protein synthesis**

168 The percentage of the protein mass synthesized per day – the fractional rate of protein synthesis -
169 was measured using the flooding dose assay [29], modified for using stable isotope tracer, the ring-
170 D₅-phenylalanine (D₅-Phe) [30]. In short, the ratios of the amount of D₅-Phe relative to the amount of
171 total phenylalanine (D₅-Phe plus its natural version) in both the protein pool and the free pool of

172 amino acids allow calculation of the fractional rate of protein synthesis. The assay was first validated
173 for brown trout of this age and size by conducting a preliminary time-course experiment (see ESM).
174 From this validation experiment, we determined that a D₅-Phe incubation period of approximately 60
175 min was an appropriate incorporation duration.

176 For the main experiment, the fish were fasted for 21h before being injected into the peritoneum with
177 the D₅-Phe solution. Each fish was then immediately placed in an individual tank containing 2 L of
178 aerated water for a period of approximately 1h (mean ± SE: 1h05min ± 0h00min) without food and in
179 darkness. The fish were then culled and their livers were immediately dissected, weighed and rinsed
180 with distilled water. A subsample of liver was weighed and kept in ice-cold respirometry buffer (0.1
181 mM EGTA, 15 μM EDTA, 1mM MgCl₂, 20mM Taurine, 10mM KH₂PO₄, 20mM HEPES, 110 mM D-
182 sucrose, 60 mM lactobionic acid, 1g L⁻¹ bovine serum albumin essentially fatty acid-free, pH 7.2 with
183 KOH) for subsequent measurement of mitochondrial properties (see below). A second aliquot of liver
184 for measurement of protein synthesis was weighed and immediately flash-frozen in liquid nitrogen
185 and stored at -70°C until further analysis. Likewise, two samples of white muscle were taken dorsally
186 to the lateral line (to avoid contamination with red fibres) and just behind the dorsal fin. One aliquot
187 was collected from one side of the fish and kept in respirometry buffer while the other aliquot was
188 collected from the other side and immediately flash-frozen. After extraction and quantification of the
189 the phenylalanine isotopes in both the free amino acid pool and in the protein pool (Details in ESM),
190 the fractional rate of protein synthesis (K_s in % day⁻¹) was calculated as:

$$191 \quad K_s = \frac{24}{t} * \frac{(D5Phe / Total Phe) \text{ in protein amino acid}}{(D5Phe / Total Phe) \text{ in free amino acid}} * 100$$

192 where t is the actual duration of D₅-Phe exposure in hours.

193

194 **Measurement of mitochondrial properties**

195 Since only two samples could be run simultaneously to measure mitochondrial properties, liver
196 samples of the two individuals in a processing batch were first homogenized as in [15, 16] and
197 assessed for mitochondrial function, while the subsample of white muscle was preserved in
198 respirometry buffer on ice for the subsequent run.

199 Oxygen and magnesium green fluorescence signals were detected simultaneously using two
200 respirometry chambers equipped with fluorescent sensors and recorded using DatLab software
201 (Oroboros Instruments, Innsbruck Austria). Tissue homogenate from each fish was added to one of
202 the two measurement chambers immediately following preparation. Mitochondrial efficiency was
203 measured as in Salin, Villasevil [31]. Briefly, we used a protocol for estimating the ATP/O ratio that
204 simultaneously measures both oxygen consumption and ATP production on the same sample.

205 Cytochrome c oxidase (COX) respiration was then measured to allow standardization of the
206 mitochondrial density of the tissues [32]. The rate of oxygen consumption simultaneously to ATP
207 production was assessed by adding saturating ADP to the chamber containing complex I and II
208 substrates. COX activity was measured after addition of ascorbate and N,N,N',N'-Tetramethyl-p-
209 phenylenediamine dihydrochloride. The muscle trial was identical to the liver trial using the
210 subsample of muscle that was kept on ice (see ESM for full details of the protocol).

211 Rates of mass-specific oxygen consumption and ATP production at each step of the protocol were
212 averaged over 30 to 60 seconds of stabilisation. Fluxes of O₂ and ATP were expressed in pmoles s⁻¹
213 mg⁻¹ wet weight of tissue. The ATP/O ratio was calculated as the ratio of corrected ATP production to
214 double the rate of O₂ consumption at the time that the ATP was being produced.

215

216 ***Statistical analysis:***

217 We first used correlation analysis to test whether physiological parameters (mitochondrial efficiency
218 [ATP/O ratio], mitochondrial density [COX activity] and fractional rate of protein synthesis [Ks]) were
219 correlated between the liver and white muscle within the same fish. We then used linear mixed
220 models to determine the links between mitochondrial efficiency of the liver and/or muscle and the
221 fractional rate of protein synthesis for different rates of food intake. The models included Ks of liver
222 or muscle as the dependent variable, ATP/O ratio of liver and muscle as continuous predictors, and
223 the food intake (high or low) as a fixed factor, and two-way interactions between food intake and
224 covariates. To control for effects of mitochondrial density on the fractional rate of protein synthesis,
225 the models included COX activity of the liver and muscle as a covariate and in two-way interactions
226 with food intake, with Ks as the dependent variable. Processing batch was included as a random
227 effect to control for the order in which fish were processed. Preliminary analyses showed that the
228 fractional rate of protein synthesis was not affected by the duration of D₅-Phe exposure or the mass
229 of sample used for the extraction of the phenylalanine isotopes, so exposure duration and mass of
230 sample were not included as covariates in the final models. We finally tested whether the degree of
231 mitochondrial efficiency and the fractional rate of protein synthesis of the liver and/or the muscle
232 explained individual variation in growth performance using a linear mixed model approach. The
233 models included the growth performance (Specific growth rate, Growth efficiency and Specific
234 protein gain) as dependent variables, and ATP/O ratio and Ks of liver and muscle as continuous
235 predictors, the food intake as a fixed factor, with processing batch as a random factor. To control for
236 effects of mitochondrial density on growth performance, COX activity of the liver or muscle were
237 included as a covariate in the models with specific growth rate, growth efficiency and specific protein
238 gain as the dependent variable. These models also included two-way interactions between covariates

239 and food regime. To control for effects of initial body size on growth performance, initial body mass
240 was included as a covariate in the models with specific growth rate or growth efficiency as the
241 dependent variable, while the initial estimate for whole-body protein content was included as a
242 covariate in the model for specific protein gain. All models were simplified by removing non-
243 significant terms in a backward deletion procedure, starting with two-way interactions; significance
244 was tested when terms were dropped from the model. All statistical analyses were performed in IBM
245 SPSS Statistics 21 (Chicago, IL). Data are presented as means \pm standard error, and the significance
246 level was set to $P < 0.05$.

247

248

249 RESULTS

250 The mitochondrial efficiency (ATP/O ratio) showed significant inter-individual variation, varying at
251 least twofold for each tissue across individuals having the same food intake (table S1). The fractional
252 rate of protein synthesis K_s differed up to two- or five-fold in liver and muscle, respectively, among
253 individuals with the same food intake (table S1). There was no correlation between the physiological
254 traits (ATP/O ratio and K_s) of the liver and muscle from the same fish (table S2).

255 The fractional rate of muscle protein synthesis K_s in a fish depended on the ATP/O ratio of its liver
256 mitochondria, although this effect depended on food intake (liver ATP/O by food intake interaction,
257 table 1). While muscle K_s was positively related to the ATP/O ratio in the liver mitochondria of fish
258 with the high food ration ($t = 2.80$; $df = 36$; $P = 0.008$), there was no such relationship in fish receiving
259 a low food ration ($t = -0.92$; $df = 36$; $P = 0.362$; figure 1). Amongst-individual variation in the fractional
260 rate of protein synthesis K_s in the liver was not explained by the mitochondrial efficiency in either
261 liver or muscle (LMM, $P > 0.05$).

262 Not surprisingly, food intake had a positive effect on specific growth rates, with fish on average
263 having a specific growth rate threefold higher at the high compared to the low ration (table S1).
264 However, individuals from the same food treatment varied considerably in their specific growth rate,
265 with the fastest growing fish in the low ration exceeding the growth of some fish on the high ration
266 (figure 2a; low food intake: -6.00 to 110.57 mg day⁻¹; high food intake: 68.86 to 394.43 mg day⁻¹). This
267 individual variation in growth rate was partially explained by differences in liver mitochondrial
268 efficiency, although the effect depended on food intake (liver ATP/O by food treatment interaction;
269 table 2). The specific growth rate of fish receiving high rations was strongly and positively linked to
270 the ATP/O ratio in their liver mitochondria ($t = 4.46$, $df = 41$, $P < 0.001$, figure 2a), whereas the trend
271 was not significant when food intake was low ($t = 0.33$, $df = 41$, $P = 0.745$). Regardless of the food
272 intake, the specific growth rate of a fish was strongly but negatively linked to the K_s in its muscle

273 after controlling for liver ATP/O (table 2). Specific growth rates under either ration were unrelated to
274 the ATP/O ratio in muscle mitochondria, or to the Ks in the liver (table 2).

275 Growth efficiency varied among individuals from -0.13 to 2.23 gain in body mass per mass of food
276 eaten but did not differ between low and high food fish (table S1). Regardless of their food intake,
277 individuals that had the higher ATP/O ratio in the liver had the highest growth efficiency (table 2,
278 figure 2b).

279 The rate of protein gain of the trout also differed considerably amongst individuals, ranging from -
280 1.98 to 17.74 mg day⁻¹ for fish eating the low ration and from -0.21 to 60.79 mg day⁻¹ for fish on the
281 high ration. Individuals that had a higher ATP/O ratio in their liver mitochondria, and a lower Ks in
282 their muscle had a faster specific gain in protein mass (table 2). The specific rate of protein gain was
283 not related to ATP/O ratio in the muscle mitochondria nor to Ks in the liver (table 2).

284

285

286 **DISCUSSION**

287 While the general trend was for growth performance to increase when food intake was higher,
288 individuals exhibited markedly differing growth performance even when having identical food intake.
289 This variation in growth was related to mitochondrial function: individuals that were more efficient at
290 producing ATP within their liver mitochondria grew faster, more efficiently and accumulated more
291 protein than those individuals with less efficient mitochondria. Individuals that had a higher liver
292 mitochondrial efficiency under high food levels had a faster rate of protein synthesis in their muscle.
293 However, these differences in protein synthesis had an effect on growth performance in the
294 completely opposition direction to our initial prediction that “protein synthesis promotes growth”. In
295 summary, our study shows for the first time that under conditions of a fixed food intake, the
296 mitochondrial efficiency of an individual animal can determine whether it grows fast or slow.

297 Individual variation in growth performance is likely to be a complex, integrative characteristic
298 influenced by several physiological and behavioural traits. Because individual differences in growth
299 rate covary with behaviours that increase feeding rates [33], only studies of animals with controlled
300 food intakes can shed light on the physiological drivers of growth differences. Food intake in our
301 experiment was standardized, revealing that growth of fish under the same ration could vary more
302 than 3-fold amongst individuals. Consequently, some fish on the low ration treatment were actually
303 faster growing than others on the high ration treatment that were consuming three times as much
304 food. While it has previously been shown that increased mitochondrial efficiency promotes fitness-
305 related traits (physical performance [34], growth performance [9, 21-23, 35], reproductive output
306 [36] and ageing [9, 14, 36, 37]), here we demonstrate that this relationship can even occur when

307 animals are experiencing similar rates of food intake. As well as varying amongst individuals,
308 mitochondrial efficiency is a flexible trait that can change in response to environmental conditions
309 [38, 39] and stage of life [34, 40]. A higher mitochondrial efficiency may also have a cost, since
310 mitochondria are a major producer of reactive oxygen species (ROS) and mitochondrial efficiency can
311 be positively related to ROS production [17, 37]. When the generation of ROS in an organism exceeds
312 the capacity of its antioxidant defence and repair mechanisms to combat its effects, there can be an
313 accumulation of oxidative damage [41]. ROS have been proposed as an important factor underlying
314 cellular and whole-organism senescence [41] and therefore, a potential cost linked to fast growth
315 [42, 43]. Despite this cost, in some contexts natural selection may favour phenotypes with relatively
316 high mitochondrial efficiency (since this can lead to faster growth, increased body size at maturity,
317 minimized mortality risk and higher number of eggs), whereas in other contexts a lower
318 mitochondrial efficiency and decreased ROS production might be beneficial (e.g. under conditions of
319 *ad libitum* food availability) [7, 17, 37, 44]. This hypothesis is in accordance with several recent
320 studies suggesting that variation in mitochondrial function is a key target of natural selection [45,
321 46]. Our findings that fish with high liver mitochondrial efficiency had a high rate of protein synthesis
322 in their muscles and faster growth match our predictions that a higher efficacy at converting food
323 into ATP can lead to an increased allocation to energetically-costly processes such as protein
324 synthesis and growth. Contrary to expectations, the rate of protein synthesis in white muscle was
325 negatively correlated with growth performance; individuals that grew the best displayed lower rates
326 of muscle protein synthesis for a given liver mitochondrial efficiency. An explanation for this
327 discrepancy might lie in the fact that rates of protein synthesis are tissue-specific [47] and the
328 correlation of protein synthesis rates across different tissues in the same individual can be poor (as
329 shown by this study), and so the range of tissues that have been measured in our study might not be
330 representative of the overall rate of protein synthesis in the entire animal since this would be
331 defined as the sum of the individual tissue-specific rates of protein synthesis [48]. However, positive
332 relationships between protein synthesis in white muscle and body growth have been reported in
333 other species [26, 47]. An alternative explanation is based on the fact that body proteins are
334 continually being broken down as well as synthesised, and so protein synthesis will only result in
335 growth if the rate of synthesis exceeds the rate of degradation; it has previously been shown that
336 growth variation among individual fish is more explained by variation in rates of protein degradation
337 than rates of protein synthesis [26]. While measurements of protein degradation rates were beyond
338 the scope of the present study, it may only be possible to explain observed patterns of protein
339 growth if all aspects of protein metabolism (synthesis and degradation) are considered [49].

340 In conclusion, our study has demonstrated a clear positive relationship between the efficiency with
341 which liver mitochondria convert energy substrates into ATP and whole animal growth performance.
342 Future research should focus on quantifying the presumed costs of highly efficient mitochondria.

343 Information on the causes and consequences of variation in mitochondrial efficiency would allow
344 prediction of the consequences for whole animal performance of variation in mitochondrial function,
345 so linking cellular processes to organismal fitness.

346

347 **Acknowledgements.** We thank Graham Law, Ross Phillips and Alastair Kirk for help with fish
348 husbandry. All procedures were carried out under the jurisdiction of a UK Home Office project
349 license (PPL 60/4292).

350

351 **Author contributions.** KS, SGL, IMcC, CC and NBM conceived the ideas and designed methodology.
352 KS, EMV, GJA, SGL and CAM collected the data. KS, EMV, IMcC, SGL and CAM analysed the data. KS
353 led the writing of the manuscript; SGL, IMcC, CS and NBM revised the manuscript and added
354 comments. All authors gave final approval for publication.

355

356 **Competing interests.** The authors declare they have no competing interests.

357

358 Data accessibility. The dataset supporting this article are available from the Dryad Digital Repository:
359 DOI: <https://datadryad.org/review?doi=doi:10.5061/dryad.5c5372c>

360

361 **Funding.** This research was supported by a European Research Council Advanced Grant (number
362 322784) to NBM

363

364

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- 507

508 **Table 1.** Results from linear mixed model analysis of the fractional rate of protein synthesis (Ks) in
 509 the muscle of a brown trout as a function of its food intake and the properties (ATP/O ratio and
 510 cytochrome c oxidase [COX] activity) of mitochondria in its muscle and liver. Processing batch was
 511 included as a random effect to control for the order in which fish were processed. Non-significant
 512 terms were excluded from the final analysis. Bold denotes significant results.

Dependant variable	Source of variation	Parameter estimate \pm SE	F	d.f.	P
Muscle Ks*	Intercept	-0.00 \pm 0.41			
	Food Intake#	0.88 \pm 0.42	4.38	1, 39.71	0.043
	Liver COX activity	0.00 \pm 0.01	0.04	1, 46.95	0.837
	Muscle COX activity	0.03 \pm 0.01	5.25	1, 36.56	0.028
	Liver ATP/O ratio	0.66 \pm 0.23	1.30	1, 30.99	0.262
	Muscle ATP/O ratio	-0.03 \pm 0.03	1.17	1, 26.98	0.289
	Food Intake# x Liver ATP/O ratio	-0.92 \pm 0.39	5.58	1, 40.41	0.023

513 #Food intake: Two-level fixed factor (Low and High food intake).

514 *Full model: Muscle Ks = Food intake + Liver COX activity + Muscle COX activity + Liver ATP/O ratio + Muscle ATP/O ratio + Food intake x

515 Liver ATP/O ratio + Food intake x Liver COX activity + Food intake x Muscle COX activity + Food intake x Muscle ATP/O ratio .

516 **Table 2.** Results from linear mixed model analyses of indices of growth performance in individual
517 brown trout as a function of their initial mass, their liver and muscle mitochondrial density
518 (cytochrome *c* oxidase [COX] activity), food intake, liver and muscle mitochondrial efficiency (ATP/O
519 ratio) and fractional rates of protein synthesis (Ks). Processing batch was included as a random effect
520 to control for the order in which fish were processed. Non-significant terms were excluded from the
521 final analysis. Bold denotes significant results.

Dependant variable	Source of variation	Parameter estimate ± SE	F	d.f.	P
Specific Growth Rate*	Intercept	-0.38 ± 0.59			
	Initial Body Mass	0.05 ± 0.02	9.69	1, 41	0.003
	Liver COX activity	-0.01 ± 0.01	0.71	1, 41	0.403
	Muscle COX activity	0.06 ± 0.02	8.27	1, 41	0.006
	Food Intake [#]	0.55 ± 0.59	0.87	1, 41	0.355
	Liver ATP/O ratio	1.61 ± 0.36	11.7	1, 41	0.001
	Muscle ATP/O ratio	-0.02 ± 0.04	0.18	1, 41	0.671
	Liver Ks	-0.01 ± 0.02	0.30	1, 41	0.586
	Muscle Ks	-0.54 ± 0.20	7.58	1, 41	0.009
	Food Intake[#] x Liver ATP/O ratio	-1.49 ± 0.54	7.56	1, 41	0.009
Growth Efficiency[‡]	Intercept	0.13 ± 0.41			
	Initial Body Mass	0.06 ± 0.02	10.8	1, 48	0.002
	Liver ATP/O ratio	0.72 ± 0.33	4.87	1, 48	0.032
Specific Protein Gain[‡]	Intercept	-3.03 ± 0.74			
	Initial Protein Mass	0.00 ± 0.00	81.3	1, 31.25	< 0.001
	Liver COX activity	0.02 ± 0.01	2.80	1, 39.94	0.102
	Muscle COX activity	0.09 ± 0.03	0.11	1, 33.93	0.299
	Food Intake [#]	2.15 ± 0.85	6.34	1, 33.81	0.017
	Liver ATP/O ratio	1.04 ± 0.29	13.0	1, 30.84	< 0.001
	Muscle ATP/O ratio	0.02 ± 0.05	0.16	1, 18.86	0.690
	Liver Ks	0.01 ± 0.02	0.28	1, 35.40	0.601
	Muscle Ks	-0.51 ± 0.24	4.44	1, 37.94	0.042
	Food Intake[#] x Initial Protein Mass	-0.00 ± 0.00	29.4	1, 19.30	< 0.001
	Food Intake[#] x Muscle COX activity	-0.13 ± 0.05	6.84	1, 32.27	0.013

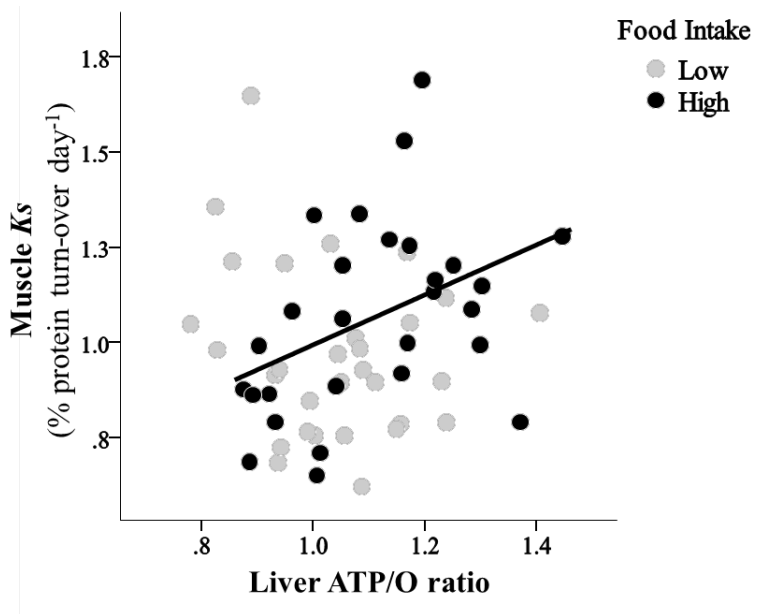
522 [#]Food intake: Two-level fixed factor (Low and High food intake).

523 [‡]Full model: Specific Growth Rate = Liver COX activity + Muscle COX activity + Initial Body Mass + Food intake + Liver ATP/O ratio + Muscle
524 ATP/O ratio + Liver Ks + Muscle Ks + Food intake x Liver COX activity + Food intake x Muscle COX activity + Food intake x Initial Body Mass +
525 Food intake x Liver ATP/O ratio + Food intake x Muscle ATP/O ratio + Food intake x Liver Ks + Food intake x Muscle Ks.

526 [‡]Full model: Growth Efficiency = Liver COX activity + Muscle COX activity + Initial Body Mass + Food intake + Liver ATP/O ratio + Muscle
527 ATP/O ratio + Liver Ks + Muscle Ks + Food intake x Liver COX activity + Food intake x Muscle COX activity + Food intake x Initial Body Mass +
528 Food intake x Liver ATP/O ratio + Food intake x Muscle ATP/O ratio + Food intake x Liver Ks + Food intake x Muscle Ks.

529 [‡]Full model: Specific Protein Gain = Liver COX activity + Muscle COX activity + Initial Protein Mass + Food intake + Liver ATP/O ratio +
530 Muscle ATP/O ratio + Liver Ks + Muscle Ks + Food intake x Liver COX activity + Food intake x Muscle COX activity + Food intake x Initial
531 Protein Mass + Food intake x Liver ATP/O ratio + Food intake x Muscle ATP/O ratio + Food intake x Liver Ks + Food intake x Muscle Ks.

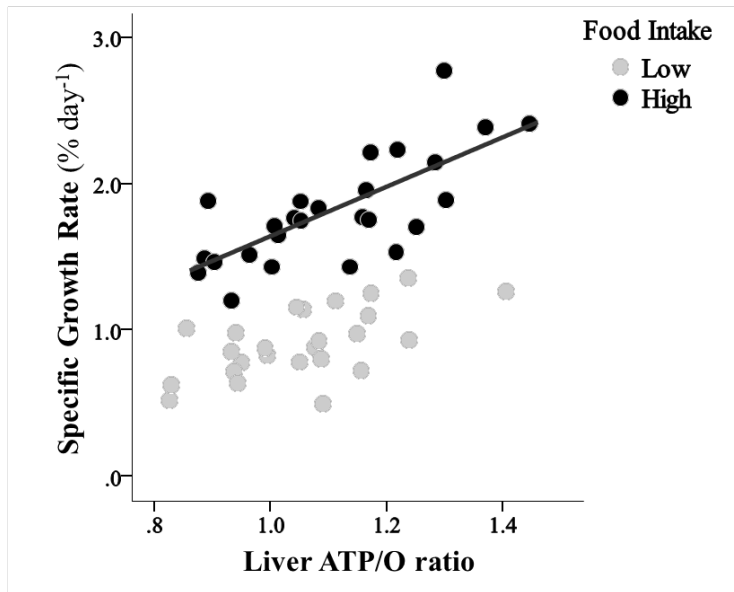
532 **Figure 1.** Relationship between the fractional rate of protein synthesis (K_s) in the muscle and
533 mitochondrial efficiency (ATP/O ratio) in the liver of juvenile brown trout at low vs high food intake.
534 Continuous lines show significant effect. N = 28-30 fish per food level. See Table 1 for statistical
535 analyses.



536

537 **Figure 2.** Relationships between indices of growth performance and mitochondrial efficiency in
538 juvenile brown trout at low vs high food levels. **(a)** Specific Growth Rate in relation to liver
539 mitochondrial efficiency (ATP/O ratio), and **(b)** Growth Efficiency in relation to liver ATP/O ratio.
540 Continuous lines show significant effects. N = 29-30 fish per food level. See Table 2 for statistical
541 analyses.

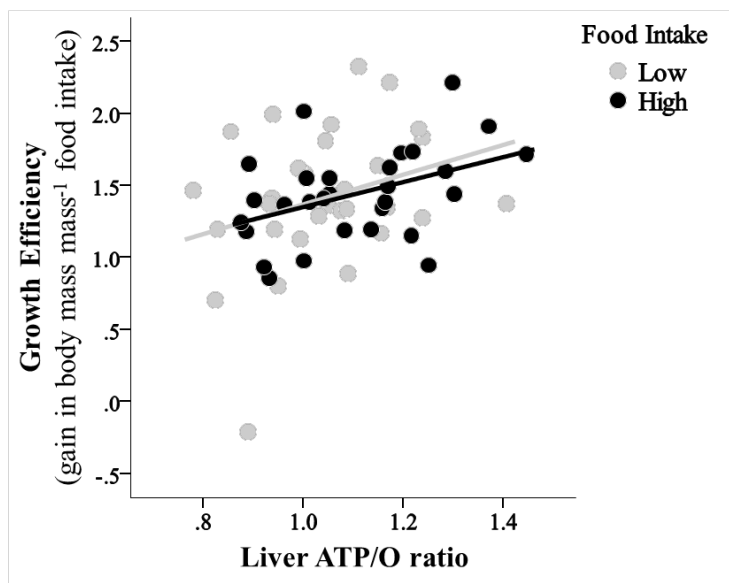
542 **(a)**



543

544 Plotted are partial residuals of specific growth rate for fish at high food ration evaluated at mean
545 initial body mass = 9.59 g.

546 **(b)**



547

548 Plotted are partial residuals of growth efficiency evaluated at mean initial body mass = 9.02 mg.