# The relationship between membrane fatty acid content and mitochondrial efficiency differs within- and betweenomega-3 dietary treatments

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#### Abstract :

An important, but underappreciated, consequence of climate change is the reduction in crucial nutrient production at the base of the marine food chain: the long-chain omega-3 highly unsaturated fatty acids (n-3 HUFA). This can have dramatic consequences on consumers, such as fish as they have limited capacity to synthesise n-3 HUFA de novo. The n-3 HUFA, such as docosahexaenoic acid (DHA, 22:6n-3) and eicosapentaenoic acid (EPA, 20:5n-3), are critical for the structure and function of all biological membranes. There is increasing evidence that fish will be badly affected by reductions in n-3 HUFA dietary availability, however the underlying mechanisms remain obscure. Hypotheses for how mitochondrial function should change with dietary n-3 HUFA availability have generally ignored ATP production, despite its importance to a cell's total energetics capacity, and in turn, whole-animal performance. Here we (i) quantified individual variation in mitochondrial efficiency (ATP/O ratio) of muscle and (ii) examined its relationship with content in EPA and DHA in muscle membrane of a primary consumer fish, the golden grey mullet Chelon auratus, receiving either a high or low n-3 HUFA diet. Mitochondria of fish fed on the low n-3 HUFA diet had higher ATP/O ratio than those of fish maintained on the high n-3 HUFA diet. Yet, mitochondrial efficiency varied up about 2-fold among individuals on the same dietary treatment, resulting in some fish consuming half the oxygen and energy substrate to produce the similar amount of ATP than conspecific on similar diet. This variation in mitochondrial efficiency among individuals from the same diet treatment was related to individual differences in fatty acid composition of the membranes: a high ATP/O ratio was associated with a high content in EPA and DHA in biological membranes. Our results highlight the existence of interindividual differences in mitochondrial efficiency and its potential importance in explaining intraspecific variation in response to food chain changes.

# Highlights

► Marine fish performance rely on fed omega-3 that is predicted to decline in a near future. ► Surprisingly, we still know little about the consequence of omega-3 deficient diet on fish performance. ► Mitochondrial ability to make ATP increased in fish fed on deficient omega-3 diet compared with those fed high levels of omega-3. ► Mitochondrial metabolism may provide new insights into the mechanisms underlying fish performance under omega-3 deficiency.

**Keywords** : ATP/O ratio, docosahexaenoic acid (DHA), eicosapentaenoic acid (EPA), food quality, Chelon auratus, global change.

# 43 1. Introduction

44 Fish provide critical sustenance for millions of people worldwide and have far reaching impacts on 45 the productivity of ecosystems (McIntyre et al., 2016). Yet, ongoing and future climate change 46 threatens the persistence of fish populations globally (Pörtner and Knust, 2007). An important, but 47 underappreciated, consequence of climate change is the reduction in production at the base of the 48 food chain of essential nutrient: the long-chain omega-3 highly unsaturated fatty acids (n-3 HUFA) 49 (da Motta Pacheco et al., 2014; Galloway and Winder, 2015; Hixson and Arts, 2016). Water warming 50 (Hixson and Arts, 2016), acidification (Bermudez et al., 2015), or UV irradiation (Kang, 2011), all affect 51 n-3 HUFA primary producers physiology and community assemblages, leading to a dominance of n-3 52 HUFA-impoverished taxa (Galloway and Winder, 2015). The n-3 HUFA such as eicosapentaenoic acid 53 (EPA, 20:5n-3) and docosahexaenoic acid (DHA, 22:6n-3) are essential to the structure and function 54 of all biological membranes and are thus considered to be important drivers of organism performance (Hulbert et al., 2005; Ishizaki et al., 2001; Mazorra et al., 2003; Vagner et al., 2015). 55 56 Endogenous biosynthesis of n-3 HUFA from precursors is limited in most vertebrates, including 57 marine fish (Alimuddin et al., 2005; Arts and Kohler, 2009; Oboh et al., 2017). Changes in the n-3 58 HUFA availability in the fish diet causes strongly correlated changes in the fatty acid composition of 59 their biological membranes (Guderley et al., 2008; Nogueira et al., 2001; Ramsey et al., 2005). Fish on 60 n-3 HUFA-deficient diets can perform badly: reductions in EPA and DHA dietary content reduced 61 growth (Norambuena et al., 2015; Vagner et al., 2015) and thermal tolerance (Vagner et al., 2014), 62 and altered whole-organism metabolic traits (Vagner et al., 2015; Vagner et al., 2014). Determining 63 what causes animal performance to vary with n-3 HUFA diet constitutes a fundamental step in 64 predicting the impacts of projected oceanic changes on fish resilience (Kang, 2011). 65 Consideration of mitochondrial capacity to make ATP may improve our understanding of the links between dietary n-3 HUFA availability and whole-animal performance. Mitochondrial ATP is 66 67 produced via oxidative phosphorylation, a process through which energy substrates are oxidized to 68 generate a protonmotive force that drives the phosphorylation of ADP to ATP. Hypotheses for how 69 mitochondrial function should change with dietary n-3 HUFA availability have focused on variation in 70 respiratory capacities and have generally ignored variation in ATP production (Kraffe et al., 2007; 71 Ramsey et al., 2005; but see Herbst, 2014; Vagner et al., 2015; Vagner et al., 2014). Although ATP 72 production depends on the rate of substrate oxidation, the number of ATP molecules produced for 73 each atom of oxygen consumed by the mitochondria during substrate oxidation (ATP/O ratio) can 74 vary (Brand, 2005; Salin et al., 2015). A fraction of the protonmotive force that is generated from

rs substrate oxidation is dissipated through proton leak across mitochondrial inner membranes and this

76 leakage can decrease the protonmotive force available to produce ATP (Brand, 2005; Kadenbach,

2003). Thus, the greater the mitochondria leak, the less efficiently an animal converts its metabolic

78 substrates into ATP, and the lower the ATP/O ratio. The proportion of energy dissipated in the 79 proton leak and the efficiency to make ATP vary among conspecific (Bottje and Carstens, 2009; 80 Robert and Bronikowski, 2010; Salin et al., 2016b) and can be influenced by environmental factors including diet (Fontaine et al., 1996; Salin et al., 2018). A number of studies have found positive links 81 82 between intraspecific heterogeneity in efficiency to produce mitochondrial ATP and the whole-83 organism performance, such as locomotory performance (Coen et al., 2012; Distefano et al., 2018; 84 but see Jahn and Seebacher, 2019), developmental rate (Salin et al., 2012), growth efficiency (Bottje 85 and Carstens, 2009; Salin et al., 2019) and reproductive output (Robert and Bronikowski, 2010), 86 suggesting that it might be a trait of ecological relevance.

87 Recent research has recognized the importance of accounting for individual heterogeneity in 88 predicting responses to global changes (Hamel et al., 2018). Because some individuals perform much 89 better than others within the same environment, individual heterogeneity is likely to directly 90 influence the potential for species to evolve adaptations for a reduced n-3 HUFA availability. A 91 number of studies have found positive links between n-3 HUFA content in membranes and 92 mitochondrial proton leak when comparing among species (Brand et al., 1994; Brookes et al., 1998), 93 but contradictory results were found when comparing treatment groups with studies reporting 94 positive (Martin et al., 2013), negative (Fontaine et al., 1996; Guderley et al., 2008) or no 95 relationships (Guderley et al., 2008) between content in n-3 HUFA in mitochondrial membranes and 96 leak respiration. N-3 HUFA are thought to have important effects on the mitochondrial capacity to 97 make ATP; but until now, there has, to our knowledge, been no assessment of whether membrane n-98 3 HUFA content could explain variation in mitochondrial metabolism among individuals.

99 The present experiment integrates measurements of mitochondrial efficiency and 100 mitochondrial proton leak to determine whether reductions in EPA and DHA availability in food led to 101 changes in energy metabolism of a primary consumer fish. We first examined the effect of dietary n-102 3 HUFA content on mitochondrial function, in particular on the efficiency to produce ATP (ATP/O 103 ratio) and the respiratory capacities to offset the proton leak (LEAK respiration). Secondly, we tested 104 whether differences between individuals in mitochondrial efficiency and mitochondrial LEAK 105 respiration vary with membrane n-3 HUFA content. To address this, we experimentally manipulated 106 the quantity of n-3 HUFA in food for wild-caught juvenile golden grey mullet (Chelon auratus). Fish 107 were fed either a high n-3 HUFA or low n-3 HUFA diet, and their membrane fatty acid composition 108 and mitochondrial functioning were determined in skeletal muscle. We choose juvenile golden grey 109 mullet as our study organism because they are likely to be the first levels of the food chain to face a 110 decline in availability of dietary n-3 HUFA, as this fish fed mainly on primary producers (Lebreton et al., 2011; Mourente and Tocher, 1993). We analysed mitochondrial properties in the skeletal muscle 111 112 because the mitochondrial function of this tissue is known to influence whole-animal performance

113 (Coen et al., 2012; Salin et al., 2016a), and that fatty acid dietary content influences muscle

114 membrane fatty acid composition (McKenzie et al., 1998; Vagner et al., 2015).

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#### 116 2. Materiel and methods

117 2.1. Fish origin and care

118 Wild juvenile golden grey mullets (n = 23) were netted from the marshes of L'Houmeau France 119 (46°12'14.4"N 1°11'43.7"W) in November 2017 and transported to the laboratory Littoral 120 Environment Society (LIENSs), France, where all the experiments were conducted. Fish were kept in a 121 common thermoregulated 300 L-tank supplied with aerated recirculated sand-filtered natural 122 seawater and equipped with an external biological filter (Eheim, Deizisau, Germany). Fish were 123 maintained under a 12 L : 12 D photoperiod, and fed daily with commercial pellets (Le Gouessant<sup>®</sup>, 124 Lamballe, France). Temperature (13.8 ± 0.2°C) and salinity (28.7 ± 0.1) were monitored daily 125 (TetraCon<sup>®</sup> 325, Laboratoires Humeau, La Chapelle-sur-Erdre, France) and were kept similar to that 126 of the sampling site. Oxygen  $(87.9 \pm 2.6\% \text{ air saturation})$  was monitored once a week. Fish were 127 acclimated for seven weeks to these conditions. The collection and handling of the animals were 128 carried out under the jurisdiction of the Departmental Service of Fisheries and the Animal Care 129 Committee of France (# 12886), respectively. In January 2018, fish were anesthetized in 0.1 g L<sup>-1</sup> MS-222 (Ethyl 3-aminobenzoate 130 methanesulfonate) in seawater, measured for body mass (34.3 ± 6.5 g), individually pit-tagged 131 132 (Biolog-id, Bernay, France), and randomly assigned to one of the four replicate 300-L holding tanks (n=5-6 fish per tank). All fish were fasted for 24h beforehand to ensure their guts were empty. Fish 133 134 were acclimated for 4 weeks in their new tanks, during which time they were fed commercial pellets 135 twice daily to a total of 2% of their biomass (Le Gouessant<sup>®</sup>, Lamballe, France). In February 2018, fish were gradually acclimated to 20°C by means of a sequence of 3 step 136

increases of 2°C over a two-week period. This temperature was chosen because it is the temperature at which whole-organism performance declined in mullet fed on a low n-3 HUFA diet, but is within their natural thermal range as 20°C reflects the mean summer temperature in mullet natural environment (Vagner et al., 2015; Vagner et al., 2014). Fish were left another 2 weeks period of acclimation to initiate physiological adjustment to the change in temperature (Bouchard and Guderley, 2003). Temperature (19.9 ± 1°C) and salinity (29.6 ± 1.1) were daily measured in the four

143 experimental tanks.

144

145 2.2. Diet treatment and tissue sampling

146 Differences in dietary n-3 HUFA content were achieved by replacing fish oil (rich in n-3 HUFA) of the

147 High n-3 HUFA diet with rapeseed oil (poor in n-3 HUFA) in the Low n-3 HUFA diet (See ingredients in

148 Table S1 (a)). Experimental diets were isocaloric and isolipidic. High and Low n-3 HUFA diets 149 contained 17.5 % and 1.2 % EPA+DHA, respectively, per total fatty acid mass of the diets, which 150 represents about 15-fold difference in EPA and DHA content between diets (See Table S1 (a) for fatty acid composition of the diets). The EPA and DHA content in High n-3 HUFA diet cover the needs of 151 152 several fish species (Robin and Skalli, 2007), although the needs of the golden grey mullet are not 153 known. The Low n-3 HUFA diet was estimated to significantly reduce n-3 HUFA content in biological 154 membranes and has been shown to impair aerobic performance of golden grey mullet at 20°C 155 (Vagner et al., 2015).

156 Following the period of acclimation to water temperature of 20°C, fish were switched to the 157 experimental diets for about two months (mean ± SEM = 61 ± 2 days, range: 45 – 73 days). This 158 duration was chosen because it is sufficient to detect differences in the membrane fatty acid composition between diet treatments (Robin and Skalli, 2007). Because only two fish per day could 159 160 be analysed for their mitochondrial function at the end of the experiment, the duration of the diet 161 treatment differed between processing batches. Fish were randomly allocated to the treatment: fish had their food progressively switched to either the High n-3 HUFA diet (n = 11) or the Low n-3 HUFA 162 diet (n = 12). The experimental diet to commercial diet ratio was increased every second day from 163 164 25%:75% to 50%:50%, 75%:25% and finally to 100%:0%. Body mass did not differ between fish 165 groups subsequently assigned to the two food treatments (High n-3 HUFA diet: 31.3 ± 1.8 g; Low n-3 HUFA diet:  $34.9 \pm 1.5$  g, t-test:  $t_{21} = -1.510$ , p = 0.146). Body mass was re-measured (as above) every 166 167 2-3 weeks and rations were recalculated to adjust for growth.

168At the end of the food treatment period, fish were fasted for 24 h before being anesthetized169and culled. Fish were weighed, measured and two samples of skeletal muscle were immediately170dissected, taken dorsally from the lateral line (to have both red and white muscle) and just behind171the head. One aliquot was collected from one side of the fish and kept in ice-cold isolation buffer172(sucrose 250 mM, EGTA 1mM, Tris-HCl 20mM, pH 7.4 at 4°C) for immediate mitochondrial assay,173while the other aliquot (≈1 g) was collected from the other side and immediately flash-frozen for174subsequent fatty acid analysis.

175

#### 176 2.3. Determination of fatty acid composition

177 2.3.1. Lipid extraction

178 Lipids from muscle of fish were extracted following the method used in Mathieu-Resuge et al. (2019).

179 Muscle aliquots were ground into a fine homogeneous powder in liquid nitrogen. Lipids from 200-

180 250 mg of muscle powder were then extracted in 6 ml of mixture chloroform/methanol (2:1; v/v). To

181 ensure complete lipid extraction, samples were sonicated at 4 °C during 2 x 5 min. Lipids from

182 experimental diets were extracted following the same method. Pellets were ground in a mortar and

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- powder was transferred in chloroform/methanol (2:1; v/v). All lipid extracts were then stored at-20
   °C under nitrogen atmosphere until further analysis.
- 185

#### 186 2.3.2. Purification of polar lipids

Fatty acids of biological membranes are connected to a polar head group, altogether named polar 187 188 lipid (PL), unlike fatty acids of energy stores that constitute triglyceride, a neutral lipid. To analyse the 189 FA composition of biological membrane, PL were separated from neutral FA and FA from only PL 190 were considered. PL were separated from neutral one by solid-phase extraction following the 191 method described in (Martin et al., 2013). Briefly, an aliquot of muscle total lipid extract (1/6) was 192 evaporated to dryness under nitrogen, recovered with 3 washings of 0.5 ml of chloroform/methanol 193 (98:2; v/v) and deposited at the top of a silica gel column (40 mm × 4 mm, silica gel 60A 63–200  $\mu$ m) 194 previously heated at 450°C and deactivated with 6 weight % H<sub>2</sub>O. After elution of neutral lipids with 195 10 mL of chloroform/methanol (98:2; v/v), PL were eluted with 20 mL of methanol, transferred into a 196 vial containing 2.3 µg of internal standard (tricosanoic acid C23:0) and stored at -20°C under nitrogen 197 atmosphere. 198

# 199 2.3.3. Polar lipid transesterification

200 Polar lipid transesterification was realized as described in Mathieu-Resuge et al. (2019). PL fractions 201 were evaporated to dryness under nitrogen. Fatty acid methyl esters (FAME) were obtained by 202 addition of 800  $\mu$ L of H<sub>2</sub>SO<sub>4</sub> / methanol (3.4 %; v/v) and heating at 100°C for 10 min. FAME were then 203 extracted in 800  $\mu$ L of hexane and washed three times with 1.5 ml of distilled water saturated in 204 hexane.

205

#### 206 2.3.4. Gas chromatography analysis of FAME

207 FA composition was analysed by GC coupled with Flame-Ionization Detector (GC-FID) as described by 208 Mathieu-Resuge et al. (2019) with a Varian CP8400 gas chromatograph. After splitless-mode 209 injection, FAME were separated in parallel on two different columns (DBWAX 30 m × 0.25 mm ID x 210 0.2  $\mu$ m and DB5 30 m × 0.25 mm ID x 0.2  $\mu$ m, Agilent). FAME were identified by comparisons of their 211 retention time with those of commercial standards (Supelco 37 Component FAME Mix, PUFA No.1 212 and No.3, and Bacterial Acid Methyl Ester Mix, Sigma) and lab-made standard mixtures. The internal standard (C23:0) allowed to calculate the FA content (µg mg<sup>-1</sup>muscle wet mass). Fatty acid content 213 was then determined as the percentage of fatty acid mass per total mass of fatty acids in membrane 214 215 lipids from the polar lipids fraction. The sum of EPA and DHA content in membrane lipids was

216 calculated as:

# $\Sigma$ EPA + DHA content = $\frac{\text{mass of EPA} + \text{mass of DHA}}{\text{total mass of fatty acids}} \times 100$

217

# 218 2.4. Mitochondrial function

219 2.4.1. Isolation of mitochondria

220 Isolation of muscle mitochondria and measurement of mitochondrial efficiency were adjusted from 221 published protocols (Ghanizadeh Kazerouni et al., 2016; Salin et al., 2016c). Briefly, mitochondria 222 were isolated from 1 g (mean ± SEM: 1.07 ± 0.03 g) of muscle. Tissue was finely chopped on ice and 223 homogenized in 10 mL of isolation buffer by four gentle passes in a Potter-Elvehjem homogenizer and centrifuged at 500 g for 5 min at 4°C. The supernatant was transferred in a clean tube and 224 225 centrifuged at 1000 g for 5 min at 4°C and then centrifuged at 9000 g for 10 min at 4°C. The pellet 226 containing mitochondria was gently resuspended in 500 μL assay buffer at 4°C (20mM Taurine, 10mM KH<sub>2</sub>PO<sub>4</sub>, 20 mM HEPES, 110 mM D-sucrose, 60 mM K-lactobionate, 1g L<sup>-1</sup> BSA fatty acid free, 227 228 pH 7.2 at 20°C).

229

# 230 2.4.2 Mitochondrial ATP/O ratio and LEAK respiration measurements

Mitochondrial efficiency has been quantified through the measurement of the ATP/O ratio, which is
the amount of ATP generated per unit of oxygen consumed. Mitochondrial LEAK respiration was
determined as the rate of oxygen consumption when ATP synthesis was inhibited. We used a
protocol that simultaneously measures both mitochondrial ATP production and the oxygen
consumption related to that ATP production, as in Salin et al. (2016c). Assays were conducted at
20°C.

Oxygen and magnesium green fluorescence signals were detected simultaneously using two respirometry chambers equipped with fluorescent sensors and recorded using DatLab software (Oroboros Instruments, Innsbruck Austria). To estimate ATP production we used the magnesiumsensitive fluorescent probe, Magnesium Green, to determine changes in [Mg<sup>2+</sup>] (Szmacinski and Lakowicz, 1996). ATP production was then calculated from the rate of change in [Mg<sup>2+</sup>] and is based on the unequal affinities of ATP and ADP for Mg<sup>2+</sup> (Chinopoulos et al., 2009).

The oxygen electrodes were calibrated at two points: air-saturated assay buffer (daily) and zero oxygen after sodium dithionite addition (fortnightly). Stepwise additions of MgCl<sub>2</sub> at each run were performed for calibration of the fluorescent signal into Mg<sup>2+</sup>. The two binding affinity (K<sub>d</sub>) values of ATP and ADP for Mg<sup>2+</sup> were determined in presence of isolated mitochondria and calculated as in (Chinopoulos et al., 2014) ; the values were K<sub>d-ATP</sub> = 0.197 mM and K<sub>d-ADP</sub> = 1.609 mM. Isolated mitochondria from each fish were added to one of the two measurement chambers of the oxygraph immediately following isolation; fish from a processing pair (i.e. a Low and a High n-3

250 HUFA) were measured in parallel. The remaining part of the isolated mitochondria was preserved on

251 ice for use in a replicate trial. Assays contained 200  $\mu$ l of isolated mitochondria with 1.8 mL of assay 252 buffer in the presence of complex I substrates (5 mM pyruvate and 0.5 mM malate) and complex II 253 substrate (10 mM succinate). Magnesium green (2.2  $\mu$ M), EGTA (0.1 mM), EDTA (5  $\mu$ M) and MgCl<sub>2</sub> (1 254 mM) were added to determine changes in [Mg<sup>2+</sup>] and so to calculate the rate of ATP production as in 255 (Chinopoulos et al., 2014).

256 The rate of oxygen consumption to support ATP production was assessed by adding a 257 saturating concentration of ADP (2 mM, Mg<sup>2+</sup> free) to the chamber. The raw rate of ATP production 258 was also measured in this condition. The rate of oxygen consumption to offset the leakage of proton 259 - LEAK respiration - was then measured after inhibition of mitochondrial ATP synthesis (with 4  $\mu$ M carboxyatractyloside). The rate of ATP hydrolysis was also measured in this condition. The rate of 260 ATP hydrolysis was then added to the raw rate of ATP production to obtain the corrected rate of ATP 261 262 production. Addition of complex I inhibitor (0.5 μM rotenone) and complex III inhibitor (2.5 μM 263 antimycin A) allowed determination of residual oxygen consumption, which was then subtracted 264 from all other oxygen consumption values. The second replicated trial was used to control for 265 repeatability of the assay. It was identical to the first one, but started an hour and half later, using 266 the remaining isolated mitochondrial and the same measurement chamber. Every second day, the 267 measurement chamber associated with a treatment group was reversed to control for any inter-268 respirometry chamber difference in readings. No effect of the choice of measurement chamber on 269 mitochondrial function was detected.

270

# 271 2.4.3. Analysis of Mitochondrial ATP/O ratio and LEAK respiration

We expressed respiration rate as pmoles of  $O_2 \sec^{-1} ml^{-1}$  of assay buffer and ATP production 272 as pmoles of ATP sec<sup>-1</sup> ml<sup>-1</sup> of assay buffer for each replicate. Finally, the ATP/O ratio was calculated 273 274 as the ratio of corrected ATP production to two-fold respiration that supported ATP production; the 275 rate of respiration is doubled since each molecule of oxygen is comprised of two oxygen atoms. 276 Replicated ATP/O ratios were highly correlated (Pearson's r = 0.749, P < 0.001). However, we found a 277 consistent shift in the values of the measurements between the first and the second trials of ATP/O 278 ratio (drift between trials: Paired t = -6.194, P < 0.001). The data from the second trial of muscle 279 assay were excluded because the mitochondrial integrity may have been impaired with time post-280 isolation, as previously found in liver mitochondria (Salin et al., 2016c). Only data of the first LEAK 281 respiration and ATP/O ratio trial were kept for the main analyses. Protein concentration was 282 measured at 595 nm using the Bradford method with bovine serum albumin as a standard. We expressed LEAK respiration as pmoles of O<sub>2</sub> sec<sup>-1</sup> mg<sup>-1</sup> of mitochondrial protein. See descriptive 283 statistics of the mitochondrial function in Table S2. 284

286 2.5. Statistical analysis

287 We first examined the effect of the treatments (High and Low dietary n-3 HUFA content) on 288 the membrane fatty acid composition ( $\Sigma$  EPA + DHA content). Normality (Shapiro-Wilk test) and homoscedasticity (Bartlett-test) were tested. When normality and homoscedasticity were not 289 290 satisfied, non-parametric analyses were carried out. We used Wilcoxon tests to determine whether 291 muscle membrane content of EPA and DHA, and their sum ( $\Sigma$  EPA + DHA content) differed between 292 dietary treatment groups. To examine potential differences in final body mass between treatments 293 that could explain differences in mitochondrial function, a t-test was employed. The effect of the 294 dietary treatment on mitochondrial function (ATP/O ratio and LEAK respiration) were also analysed 295 using t-tests.

We then used linear model analyses to test whether ATP/O ratio and LEAK respiration were correlated with membrane lipid content in  $\Sigma$  EPA + DHA. The models included mitochondrial function (ATP/O or LEAK respiration) as the dependant variable, dietary treatments as a categorical effect, and  $\Sigma$  EPA + DHA content as continuous predictor. The normality of the linear model residuals was validated for all models using Shapiro–Wilk tests. We also tested the interactions between dietary treatment and membrane lipid content in  $\Sigma$  EPA + DHA. All interactions were non-significant, so they were removed from the models.

The duration of the dietary treatment was included as potential covariate, the processing batch and fish tanks were included as potential random factor in initial analysis. Each of these variable were not significant, so were removed from all the final models. All statistical analyses were performed with the free software R (R Core Team, 2017), with R Version 3.6.1, with significance level set to p < 0.05. Data are presented as means ± standard error of the mean (SEM).

308

309 3. Results

310 3.1. Effect of the dietary treatment on membrane fatty acid composition

311 Muscle of mullets fed on Low n-3 HUFA diet showed significant differences in their membrane 312 fatty acid composition compared to those of fish fed on High n-3 HUFA diet (Table S3). EPA and DHA 313 content in muscle membranes were significantly lower in Low n-3 HUFA fish compared to High n-3 314 HUFA fish (EPA: W = 119, p < 0.001; DHA: W = 132, p < 0.001; Table 1). Not surprisingly, fish on 315 average had a lower content of  $\Sigma$  EPA+ DHA in their muscle when fed the Low n-3 HUFA diet (W = 316 130, p < 0.001, Table 1). Final body mass did not differ between High and Low n-3 HUFA fish (t = -317 1.668; p = 0.112), excluding body mass as confounding sources of differences between dietary 318 treatments.

319

320 3.2. Effect of the dietary treatment on mitochondrial function

ATP/O ratio was significantly higher in the mitochondria of Low compared to High n-3 HUFA mullet (t = -3.107, p = 0.005; Fig. 1A). However, there was no effect of dietary n-3 HUFA content on LEAK respiration in muscle mitochondria (t = -0.423, p = 0.677; Fig. 1B).

324

325 3.3. Relationships across individuals between mitochondrial function and membrane fatty acid
326 composition

327 The ATP/O ratio ranged from 1.46 to 3.33 for mitochondria of mullet eating the High n-3 328 HUFA diet and from 2.05 to 3.67 for mitochondria of fish on Low n-3 HUFA diet (Fig. 2A). Variation in 329 mitochondrial function between individuals was mainly explained by differences in n-3 HUFA content 330 in muscle (Figure S1). Regardless of the food treatment, the muscle ATP/O ratio of a fish was strongly and positively related to its  $\Sigma$  EPA + DHA content: individuals that had the higher mitochondrial 331 332 efficiency under either diet had the highest content in EPA and DHA in their membranes (t = 2.513, p 333 = 0.021; Fig. 2A and Table 2). There was no relationship between a fish's mitochondrial LEAK 334 respiration and its  $\Sigma$  EPA + DHA content among individuals with the same dietary treatment (t = -0.104, p = 0.919, Fig. 2B and Table 2). We found no correlation between ATP/O ratio and LEAK 335 336 respiration from the same mitochondria (High n-3 HUFA diet: Pearson's r = 0.229, p = 0.499; Low n-3 337 HUFA diet: Pearson's r = -0.157, p = 0.627).

338

#### 339 **4. Discussion**

340 We asked whether a decline in dietary n-3 HUFA content leads to changes in mitochondrial 341 metabolic phenotype for a model of primary consumer fish, the golden grey mullet. We manipulated 342 dietary content in n-3 HUFA and assessed membrane FA composition and mitochondrial function in 343 mullet. We found that diet strongly influenced membrane fatty acid composition: mullets on a Low 344 n-3 HUFA diet had lower levels of n-3 HUFA in muscle biological membranes, which suggested that 345 the mitochondrial membranes contained less n-3 HUFA. Previous studies have indeed demonstrated 346 that modification of the fatty acid composition of the diet causes strongly correlated changes in the 347 membrane fatty acid composition of mitochondria (Herbst et al., 2014; Jeromson and Hunter, 2014; 348 Ramsey et al., 2005), including those of fish species (Guderley et al., 2008; Morash et al., 2009). Our 349 findings reveal that ATP/O ratio increased significantly in mitochondria of fish fed a low n-3 HUFA 350 diet, so that a decline in dietary content in n-3 HUFA elicits greater mitochondrial efficiency to 351 produce ATP. Surprisingly, the differences in membrane n-3 HUFA content between individuals had 352 an opposite effect on mitochondrial efficiency: under the same dietary treatment, individuals that 353 displayed higher EPA and DHA membrane content had mitochondria with the highest ATP/O ratio.

Aerobic performance of marine fish will not only be challenged in a near future by increased oxygen needs and low oxygen availability at high temperatures but probably also by a shift in ocean

356 productivity (Behrenfeld et al., 2006), where EPA and DHA availability might decline considerably 357 (Hixson and Arts, 2016). Mitochondrial aerobic capacity has traditionally been determined in terms 358 of oxygen consumption, enzymatic activities or density of mitochondria (Brookes et al., 1998; Guderley et al., 2008; Martin et al., 2013; Morash et al., 2009; Ramsey et al., 2005; Yu et al., 2014). 359 360 However, oxygen consumption provides only an indirect measure of ATP production, and the 361 relationship between oxygen consumption and ATP production can vary both among and within 362 individuals. Our results clearly show that when evaluating mitochondrial adjustment in response to a 363 reduction in n-3 HUFA availability in diet, variation in the amount of ATP generated per molecule of 364 oxygen consumed by mitochondria can have major importance in explaining associated changes in 365 whole-animal performance. Past work on mullets found that fish fed on a low n-3 HUFA diet consumed less oxygen compared to the fish fed a high n-3 diet to reach a similar swimming speed 366 367 (Vagner et al., 2015; Vagner et al., 2014). Our results suggest that higher mitochondrial efficiency 368 may result in lower oxygen requirement for fish to produce ATP and sustain muscle contraction for 369 locomotory performance, as previously demonstrated in human (Coen et al., 2012). Mitochondria 370 with high efficiency can be beneficial for energy-demanding cellular processes (Salin et al., 2015), yet 371 there could also be a cost. Mitochondria are a major producer of reactive oxygen species (ROS) and 372 mitochondrial efficiency can be positively related to ROS production (Brand, 2000; Salin et al., 2015), 373 an area which would require further study.

374 Here, we found no significant differences in LEAK respiration of mullet muscle across dietary 375 treatments. In contrast, LEAK respiration in trout muscle were higher for mitochondria from fish fed 376 a low n-3 HUFA diet than for mitochondria from fish fed a high n-3 HUFA diet (Guderley et al., 2008). 377 Another experiment in trout has shown the opposite effect: animals fed a lower n-3 HUFA diet 378 displayed lower LEAK respiration in muscle mitochondria (Martin et al., 2013). We stress that it is 379 entirely possible for the mitochondrial LEAK response to dietary content in n-3 HUFA to vary among 380 individuals, species and other environmental factors, but also that the consequence on the proton 381 leakage might depend on the magnitude and duration of the dietary treatment. Equivocal support 382 may indicate that the responses of LEAK to changes in membrane fatty acid composition does not fall 383 along a linear response curve, as typically assumed. Instead, the effect of dietary content in n-3 HUFA 384 on membrane composition, and in turn proton permeability might be biphasic (Abbott et al., 2010) 385 and might promote non-linear response of LEAK. While testing a large range of n-3 HUFA dietary 386 content was beyond the scope of the present study, it may only be possible to explain contrasted 387 pattern of LEAK response if many dietary levels of n-3 HUFA are considered.

In our study, mullets from the same food treatment displayed important individual variation in mitochondrial efficiency that covary with membrane fatty acid composition: individual that had higher EPA and DHA content in their membranes had mitochondria that synthesized ATP more

391 efficiently than those of individuals with lower EPA and DHA content. Previous studies suggested that 392 variation in FA composition of the mitochondrial inner membrane can influence the mitochondrial 393 membrane conductance of protons, and in turn the proportion of oxygen used to offset the proton leakage (Hulbert et al., 2007), leading to variation in the efficiency of ATP production (Brand, 2005). 394 395 However, this does not appear to be the case in the present study, as LEAK did not covary with 396 ATP/O ratio and membrane fatty acid composition. An explanation for this discrepancy might lie in 397 the fact that mitochondrial efficiency also depends on protein abundance and activity of the 398 mitochondrial inner membrane-bound complexes involved in oxidative phosphorylation (i.e. electron 399 transport complexes and ATP synthase). This alternative explanation is based on the fact that fatty 400 acid composition of membrane lipids may affect lipid-protein interactions and therefore the function 401 of embedded proteins (Brenner, 1984). As well as varying with membrane FA composition and 402 protein function, mitochondrial ATP/O ratio can shift in response to energy substrate use (Brand, 403 2005). Previous work in Atlantic salmon has demonstrated that individual variation in DHA and EPA in 404 muscle content was associated with differences in the expression of genes involved in lipid 405 catabolism and carbohydrate metabolism (Horn et al., 2019). However, further research is needed to 406 determine whether differences in energy substrates utilization determine individual variation in 407 mitochondrial efficiency in mullet.

408 Some mullets on the low n-3 HUFA dietary treatment had actually higher EPA and DHA 409 contents than others on the high n-3 HUFA dietary treatment that were eating almost 15 times as 410 much EPA and DHA. Individual variation in membrane fatty acid composition is likely to be a 411 complex, integrative characteristics influenced by several metabolic pathways. If individual 412 differences in fatty acid membrane composition covary with the rates of lipid assimilation, 413 biosynthesis and degradation, it is important to recognize that fatty acid composition may vary with pathways that are generally neglected. For example, ability to biosynthesis EPA and DHA in marine 414 415 fish is very limited, as it is generally insufficient to compensate dietary deficiency (Tocher, 2003), but 416 the link between individual variation in n-3 HUFA content and their rate of biosynthesis might only 417 appear across individuals eating the same amount of n-3 HUFA. Another explanation might be an 418 individual heterogeneity in the rate of degradation of EPA and DHA, while also considered minor in 419 fish (McKenzie, 2001). Individual variation in the rate of assimilation may also be significant for n-3 420 HUFA content in membrane. Regardless of their food intake, individual fish that have higher levels of 421 EPA and DHA in membrane lipids may have preferentially retained these fatty acids from their diet. 422 Our observations illustrate that an understanding of individual variability in membrane fatty acid 423 composition can be gained only through consideration of differential metabolic pathways of fatty 424 acids.

425 The individual covariation between mitochondrial efficiency and membrane fatty acid 426 composition we observed might be because higher EPA and DHA contents in the membrane promote 427 higher efficiency of the mitochondria, as explained above. However, given that our study is correlative, it is perhaps high mitochondrial efficiency that may promote membranes with relatively 428 429 high levels of EPA and DHA, because more ATP is available to fuel metabolic pathways that retain 430 dietary n-3 HUFA. Assimilation of fatty acids in the intestine can be energetically costly (Mansbach 431 and Gorelick, 2007), and for example, fish that have higher efficiency to make ATP may actually have 432 higher assimilation rate of EPA and DHA compared with fish at lower mitochondrial efficiency. The 433 variation in mitochondrial efficiency of muscle tissue studied here might be representative of the 434 mitochondrial efficiency in other tissues, including intestine, although previous studies looking at 435 correlation of mitochondrial efficiency across tissues in the same individual have shown equivocal 436 results (Salin et al., 2019). Further study looking at mitochondrial metabolism in different tissues will 437 be necessary to determine if golden grey mullets are able to improve their mitochondrial efficiency 438 across multiple tissues.

439 Flexibility in mitochondrial efficiency may be particularly important since the capacity of the 440 mitochondria to produce ATP can set limits on the capacity of an organism to respond to 441 environmental changes (Blier et al., 2013; Sokolova et al., 2012). Our data imply that the greater 442 mitochondrial efficiency induced by the Low n-3 HUFA diet might benefit for the ability of the 443 mitochondrial to make ATP for energy-demanding cellular processes when reduction in EPA and DHA 444 availability in food web. Information on the consequences of individual heterogeneity in 445 mitochondrial efficiency on fish performance would allow a better understanding of the effect of 446 decline in EPA and DHA availability in marine food web. Further research should focus on identifying 447 the individual variation in metabolic pathway of EPA and DHA. This type of variation may be very 448 important in an evolutionary context as well, not only generating phenotypic variation among 449 individuals, but also allowing animals to reduce the energy costs of making ATP by increasing the 450 mitochondrial efficiency.

451

ACKNOWLEDGEMENTS: We thank Salomé Ducos for help with fish handling and José-Luis Zambonino
Infante for designing the diets. We thank the three referees for helpful comments that improved the
manuscript.

455

456 <u>CRediT author statement</u>: KS, MMR, FLG, PS and MV conceived the ideas and designed methodology.

457 KS, MMR, NG, ED, FLG and MV collected the data. KS, MMR, NG, FLG, PS and MV analysed the data.

458 KS led the writing of the manuscript except for section 2.3 written by FLG; MMR, FLG, PS and MV

459 revised the manuscript and added comments. All authors gave final approval for publication.

460	
461	DECLARATION OF INTEREST: The authors declare that they have no known competing financial
462	interests or personal relationships that could have appeared to influence the work reported in this
463	paper.
464	
465	DATA ACCESSIBILITY: The dataset supporting this article will be made available by archiving it in the
466	Dryad Digital Repository.
467	
468	FUNDING: This work is a contribution to the project ECONAT Axe 1 - Ressources Marines Littorales:
469	qualité et éco-valorisation, funded by the Contrat de Plan Etat-Région and the CNRS and the
470	European Regional Development Fund.
471	
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- 659

660	Table 1: Mean ± SEM eicosapentaenoic acid (EPA), docosahexaenoic acid (DHA) content and sum of
661	EPA and DHA content ( $\Sigma$ EPA + DHA) in muscle membrane fatty acids (percentage of the EPA and DHA
662	mass per total fatty acids mass in membrane, %), of juvenile golden grey mullet given High (n=11) or
663	Low (n=12) omega-3 highly unsaturated fatty acids (n-3 HUFA) diet. Different letters within a row
664	indicate significant differences between diet treatments (Wilcoxon tests, at the significant level p <
665	0.05). More details on FA composition of muscle membranes are provided Table S3.
666	

Muscle membrane FA composition (%) High n-3 HUFA fish Low n-3 HUFA fish

EPA	$9.8 \pm 0.3^{a}$	5.7 ± 1.0 <sup>b</sup>
DHA	$20.8 \pm 0.7$ <sup>a</sup>	$10.1 \pm 2.0$ <sup>b</sup>
Σ ΕΡΑ + DHA	$30.6 \pm 0.8^{a}$	15.8 ± 3.0 <sup>b</sup>

667	
668	
669	Table 2: Results from linear model analyses of muscle mitochondrial efficiency (ATP/O ratio) and
670	respiratory capacities to offset the proton leak (LEAK respiration) in juvenile golden grey mullet as a
671	function of the sum of eicosapentaenoic acid (EPA), docosahexaenoic acid (DHA) content - $\Sigma$ EPA +
672	DHA in muscle (percentage of the EPA and DHA mass per total fatty acid mass in membrane). Bold
673	denotes significant results.

Dependant variable	Source of variation	Parameter estimate ± SEM	t-value	p-value
ATP/O ratio	Intercept	2.21 ± 0.23	7.978	< 0.001
	Σ ΕΡΑ + DHA	$0.04 \pm 0.01$	2.513	0.021
	diet	$1.30 \pm 0.31$	4.232	< 0.001
LEAK respiration	Intercept	56.59 ± 6.45	8.769	<0.001
	Σ EPA + DHA	$0.04 \pm 0.34$	-0.104	0.919
	diet	2.61 ± 7.17	0.364	0.720

# 676 FIGURES

- 677 **Figure 1**. Effect of omega-3 highly unsaturated fatty acids (n-3 HUFA) dietary content on (*a*)
- 678 mitochondrial efficiency estimated as ATP/O ratio ( $t_{21} = -3.107$ , p = 0.005) and (b) mitochondrial LEAK
- 679 respiration ( $t_{21}$  = -0.423, p = 0.677) of muscle of golden grey mullet that were kept either on High
- 680 (n=11) or Low (n=12) n-3 HUFA diet. Data are plotted as mean ± SEM. \* denotes significant effect.

681 *(a)* 







# 697 ELECTRONIC SUPPLEMENTARY MATERIAL

- **Table S1**: (a) Ingredient (in g 100 g<sup>-1</sup>) and (b) fatty acid (FA) composition expressed as a percentage
- of FA mass per total FA mass (% mean  $\pm$  standard error of the mean; n = 3) in the High and Low
- 700 omega-3 highly unsaturated fatty acid (n-3 HUFA) diets (made at INRA, Donzag, France). Only FA that
- 701 occur above 1 % in at least a treatment group are represented. Bold denotes significant results of
- one-way ANOVA. Assumptions of normal distribution and homoscedasticity of residuals were met.
- 703 *(a)*

Ingredients <sup>a</sup>	High n-3 HUFA diet	Low n-3 HUFA diet	
Fish meal LT 94	17	17	X
Casein	30	30	
Rapeseed oil	2	10	
Fish oil	8	0	
Precooked starch	30	30	
Vitamin mixture <sup>b</sup>	8	8	
Mineral mixture <sup>c</sup>	4	4	
Betaine	1	1	

704

705 <sup>a</sup> Sources: fish meal LT 94: Norse (Fyllingsdalen, Norway); casein: Sigma-Aldrich (Germany); rapeseed oil:

- Système U (Créteil, France); fish oil: pure cod oil Cooper (Melun, France); precooked starch: Prégéflo Roquette
   Frères (Lestrem, France); vitamin mixture (INRA Jouy-en-Josas, France)
- 708 <sup>b</sup> Vitamin mixture (g kg<sup>-1</sup> vitamin mix): retinyl acetate 1; cholecalciferol 2.5; Dl- $\alpha$  tocopheryl acetate 5;

709 menadione 1; thiamine-HCl 0.1; riboflavin 0.4; D-calcium pantothenate 2; pyridoxine–HCl 0.3; cyanocobalamin

710 1; niacin 1; choline 200; ascorbic acid (ascorbyl polyphosphate) 5; folic acid 0.1; D-biotin 1; mesoinositol 30.

711 <sup>C</sup> Mineral mixture (g kg<sup>-1</sup> mineral mix): KCl 90; Kl 0,04; CaHPO<sub>4</sub>·2H<sub>2</sub>O 500; NaCl 40; CuSO<sub>4</sub>·5H<sub>2</sub>O 3; ZnSO<sub>4</sub>·7H<sub>2</sub>O 4;

712 CoSO<sub>4</sub> 0.02; FeSO<sub>4</sub>·7H<sub>2</sub>O 20; MnSO<sub>4</sub>·H<sub>2</sub>O 3; CaCo<sub>3</sub> 215; MgOH 124; Na<sub>2</sub>SeO<sub>3</sub> 0 03; NaF 1.

<sup>714</sup> *(b)* 

Diet FA composition (%)	High n-3 HUFA	Low n-3 HUFA	F	Df	p-value
14 :0	4.30 ± 0.20	$0.41 \pm 0.02$	362.2	1	< 0.001
16 :0	18.67 ± 0.80	6.10 ± 0.26	223.6	1	< 0.001
18 :0	4.59 ± 0.21	1.91 ± 0.09	142.7	1	< 0.001
16 :1n-7	4.83 ± 0.11	$0.44 \pm 0.01$	1630	1	< 0.001
18 :1n-9	25.04 ± 0.23	56.74 ± 0.30	7004	1	< 0.001
18 :1n-7	2.56 ± 0.36	0.05 ± 0.05	46.2	1	< 0.01
20 :1n-9	$1.82 \pm 0.08$	4.94 ± 2.09	2.2	1	0.21
18 :2n-6	5.06 ± 0.36	15.13 ± 1.82	29.6	1	< 0.01
18 :3n-3	2.42 ± 0.25	6.49 ± 1.03	14.7	1	0.02
20 :4n-6	$1.38 \pm 0.07$	0.27 ± 0.04	205.1	1	< 0.001
20 :5n-3	6.63 ± 0.60	0.39 ± 0.07	106.3	1	< 0.001
22 :5n-3	$1.64 \pm 0.09$	$0.88 \pm 0.31$	5.5	1	0.07
22 :6n-3	10.95 ± 0.63	0.78 ± 0.14	246.3	1	< 0.001

716	Table S2: Mean ± SEM mitochondrial function of muscle of golden grey mullet fed with High (n=11)
717	and Low omega-3 highly unsaturated fatty acid (n-3 HUFA) diet. Rate of ATP production, rate of
718	oxygen consumption to support ATP production (OXPHOS respiration), and to offset the proton
719	leakage (LEAK respiration). Fluxes are expressed in pmol ATP produced and oxygen consumed s <sup>-1</sup> mg <sup>-1</sup>
720	of mitochondrial protein. The respiratory control ratio (RCR), an index of mitochondrial coupling
721	(Brand and Nicholls, 2011) was calculated as the ratio of OXPHOS respiration to the LEAK respiration.

Muscle mitochondrial function High n-3 HUFA fish Low n-3 HUFA fish

ATP production	3267.6 ± 445.8	3869.7 ± 237.4
OXPHOS respiration	836.8 ± 112.0	733.6 ± 60.8
LEAK respiration	55.1 ± 4.0	57.1 ± 2.9
RCR	14.6 ± 1.5	13.0 ± 1.0

722

Table S3: Results from Kruskal-Wallis tests comparing fatty acid contents (percent of FA mass per total FA mass) in membrane lipids of the muscle of golden grey mullet (*Chelon auratus*) that were either fed a High or Low omega-3 highly unsaturated fatty acid (n-3 HUFA) diet. Only FA that occur above 1 % in at least a treatment group are represented. Means are expressed ± standard error of the mean. Bold denotes significant results. N = 11-12 per treatment group.

728

Muscle membrane FA composition (%)	High n-3 HUFA	Low n-3 HUFA	Chi <sup>2</sup>	p-value
15 :0	0.97 ± 0.08	1.68 ± 0.29	4.125	0.04
16 :0	18.49 ± 0.75	21.15 ± 2.20	0.015	0.902
18 :0	9.78 ± 0.39	$9.91 \pm 1.44$	0.379	0.54
17 :1n-7	$1.29 \pm 0.09$	$1.66 \pm 0.17$	1.83	0.18
18 :1n-9	11.42 ± 0.46	19.32 ± 1.98	15.51	< 0.001
18 :1n-7	2.95 ± 0.07	$2.81 \pm 0.08$	1.67	0.20
20 :1n-9	$1.03 \pm 0.06$	1.41 ± 0.09	8.72	0.003
22 :1n-7	$1.42 \pm 0.10$	$1.40 \pm 0.19$	0.034	0.85
18 :2n-6	1.70 ± 0.24	4.95 ± 0.72	11.05	< 0.001
20 :4n-6	4.16 ± 0.08	2.31 ± 0.42	16.5	< 0.001
20 :5n-3	9.82 ± 0.32	5.70 ± 1.03	10.64	< 0.001
22 :5n-6	1.35 ± 0.08	0.89 ± 0.14	8.37	< 0.01
22 :5n-3	$5.58 \pm 0.15$	3.97 ± 0.62	2.76	0.09
22 :6n-3	20.78 ± 0.66	10.08 ± 1.96	16.5	< 0.001

729

- 731 Figure S1: Spearman's correlation analyses between the membrane lipid fatty acids and the
- 732 mitochondrial function of muscle of golden grey mullet fed with (a) High (n=11) and (b) Low omega-3
- highly unsaturated fatty acid (n-3 HUFA) diet. FA composition were calculated as percent of FA mass
- per total FA mass. Only FA that occur above 1 % in at least a treatment group are represented.
- 735 Mitochondrial function were expressed as the mitochondrial efficiency (ATP/O) and the LEAK
- 736 respiration. N = 11-12 per treatment group. The coloured circles represent significant correlations
- 737 between two variables (p < 0.05). The numbers inside cells are the associated Spearman's correlation
- 738 (*r*<sub>S</sub>).



# Highlights MERE\_2020\_399

Marine fish performance rely on fed omega-3 that is predicted to decline in a near future.

Surprisingly, we still know little about the consequence of omega-3 deficient diet on fish performance.

Mitochondrial ability to make ATP increased in fish fed on deficient omega-3 diet compared with those fed high levels of omega-3.

Mitochondrial metabolism may provide new insights into the mechanisms underlying fish performance under omega-3 deficiency.

Journal Prevention

DECLARATION OF INTEREST: The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.